In today’s presentation we will cover information regarding the organism that causes Q Fever and its epidemiology. We will also talk briefly about the history of the disease, how it is transmitted and clinical disease in humans and animals. Finally, we will address prevention and control measures for Q fever.

Coxiella burnetii is a obligate intracellular gram-negative pathogen. It was previously identified as a rickettsial agent, but has been recently reclassified as Proteobacteria, the same group that contains bacteria such as Legionella. It replicates in host monocytes and macrophages. It has tremendous stability and can reach high concentrations in animal environments. Because it forms unusual spore-like structures it is highly resistant to environmental conditions and many disinfectants. Coxiella burnetii can survive 7 to 10 days on wool at room temperature, 1 month on fresh meat in cold storage, 120 days in dust and more than 40 months in skim milk. The organism is killed by pasteurization. Coxiella burnetii exists in two antigenic phases. This is important in the diagnosis of Q fever. Phase I is pathogenic and found in infected animals or in nature. Phase II is less pathogenic and is recovered only after multiple lab passages in eggs or cell cultures. Increased antibodies to phase II antigens indicate acute infection while a rise in phase I reflects a chronic infection of Q fever.
Q Fever

**History**

- 1935
  - Queensland, Australia (abattoir worker)
  - Montana, USA (ticks)
- Outbreaks
  - Military troops
  - When present in areas with infected animals
  - Cities and towns
  - Downwind from farms
  - By roads traveled by animals

Q “Query” fever was first reported in Brisbane, Queensland, Australia, in 1935 by Derrick, who described outbreaks of febrile illness in abattoir workers. Burnet and his associate Freeman subsequently isolated the organism and investigated the epidemiology of the disease. Concurrently, a similar agent (initially called the “Nine Mile agent”) was isolated from ticks in Montana by Davis and Cox and was subsequently found to be the same organism as that found in Queensland. In 1938 the organism was named *Coxiella burnetii* in honor of Cox and Burnet. In 1944 there were outbreaks among British and American troops stationed in the Mediterranean (Italy); outbreaks also occurred during World War II and during the Persian Gulf. Outbreaks have similarly been reported among persons residing in cities and towns downwind from sites where infected animals are kept.

**Transmission**

- Aerosol
  - Parturient fluids
  - 10⁹ bacteria released per gram of placenta
  - Urine, feces, milk
- Direct contact
- Fomites
- Ingestion
- Arthropods (ticks)

Aerosolization is the primary mode of transmission in humans and domestic ruminants represent the most frequent source of infection. Organisms can be found in airborne droplets or dust contaminated by placental tissues, birth fluids, or excreta of infected animals. Shedding of *C. burnetii* into the environment occurs mainly during parturition; over 10⁹ bacteria per gram of placenta are released at the time of delivery. Aerosol or direct transmission can occur when infected animals are processed as meat, during necropsies, or while assisting deliveries.

Due to persistence of the organism in the environment, dried infective material can contaminate water, dust, and soil; *C. burnetii* has been isolated downwind up to ½ mile or more from a known source. Fomites (i.e., newborn animals, wool, bedding, clothing) can also be contaminated and serve as a source of infection. Shedding in the milk occurs due to infected mammary glands, but pasteurization kills this organism.

*C. burnetii* has been naturally and experimentally isolated from a variety of arthropods, (mainly ticks but also cockroaches, beetles, flies, fleas, lice, mites). Over 40 tick species are naturally infected with *C. burnetii*. 
and transovarial (mother to offspring) and transstadial (between developmental life stages) transmission has been documented. Feces of infected arthropods can serve as a source of \textit{C. burnetii} infection and can remain infective for at least 19 months.

Animals typically acquire Q fever through exposure to other infected animals, either through direct contact with contaminated material or aerosol exposure.

Person-to-person transmission is extremely rare. Transplacental transmission may occur resulting in congenital infection. Transmission from blood transfusions, bone marrow transplants, and intradermal inoculations have also been reported. Transmission via sexual intercourse has been hypothesized. Sexual transmission of \textit{Coxiella burnetii} has been documented in mice and guinea pigs and hypothesized for a rare number of human cases.

Q fever is a zoonosis with worldwide distribution. It has been reported on all continents, except New Zealand, and is endemic in areas where reservoir animals are found. The animal reservoir is large and includes many wild and domestic mammals, birds, and arthropods. However, the primary reservoirs are considered to be cattle, sheep, goats, and ticks. Wildlife species reported as reservoirs include snowshoe hares, moose and white-tailed deer in Nova Scotia, wild Dall sheep in Alaska, and black bears in Idaho and California.

Q fever is an occupational hazard for persons in contact with domestic animals, such as cattle, sheep and goats. Persons at risk include farmers, livestock producers, veterinarians, abattoir workers, those in contact with dairy products, and laboratory personnel performing culture and diagnostics. There has been an increase in reports of sporadic cases in people living in urban areas after occasional contact with farm animals or after contact with infected pets, such as dogs and cats.
Q Fever

There were a total of 120 human cases of Q fever reported to the CDC in 2008; the above map shows the geographic distribution. [N=Report of disease is not required in this jurisdiction]. Although relatively few human cases are reported annually, the disease is believed to be substantially underreported because of its nonspecific presentation and the subsequent failure to suspect infection and request appropriate diagnostic tests.

[Source: http://www.cdc.gov/mmwr/preview/mmwrhtml/mm5754a1.htm]

In humans, the incubation period varies from 2 to 40 days (mean around 20 days). As few as one organism is capable of causing disease. Humans are considered to be dead end hosts and are the only species known to develop illness regularly as a result of infection. Most cases of Q fever are asymptomatic; only about 50% of all people infected with C. burnetii show signs of clinical illness. The two clinical forms of the disease are acute (less than 6 months duration) and chronic (greater than 6 months duration).

Symptoms of acute disease can vary in severity and duration; a self-limited febrile or flu-like illness often occurs. Signs include fever, chills, “sweats”, retrobulbar headache, fatigue, anorexia, malaise, myalgia, and chest pain. Illness typically lasts from one to three weeks. 30 to 50% of patients with symptomatic illness will develop pneumonia. In more severe cases, a nonproductive cough with pneumonitis may develop. Radiographs of patients with pneumonia resemble those of patients with viral pneumonia etiologies. Multiple rounded opacities of both lungs on x-ray may be noted, and pleural effusion may also be seen. Additionally, many clinically ill patients will have abnormal liver enzymes, and some will develop hepatitis although jaundice is rare. Exanthema (rash) occurs in about 10% of cases. Rarely meningoencephalitis or pericarditis may occur with acute infection. Only 2% of acute infections require hospitalization and a similar percentage result in death.
### Chronic Disease

<table>
<thead>
<tr>
<th>1</th>
<th>to 5% of those infected</th>
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<tbody>
<tr>
<td>-</td>
<td>Prior heart disease, pregnant women, immunocompromised</td>
</tr>
<tr>
<td>2</td>
<td>Endocarditis</td>
</tr>
<tr>
<td>3</td>
<td>Other</td>
</tr>
<tr>
<td>4</td>
<td>- Granulomatous hepatitis</td>
</tr>
<tr>
<td>5</td>
<td>- Cirrhosis</td>
</tr>
<tr>
<td>6</td>
<td>- Osteomyelitis</td>
</tr>
<tr>
<td>7</td>
<td>50% relapse rate after antibiotic therapy</td>
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</tbody>
</table>

Chronic Q fever (infection greater than six months in duration) occurs in 1 to 5% of those infected and is relatively uncommon. It typically develops in persons with pre-existing cardiac valvular disease. Immunocompromised persons and pregnant women are also at great risk for the chronic form. Endocarditis is the major clinical presentation and accounts for 60 to 70% of all chronic Q fever cases. Infection can also affect the liver causing granulomatous hepatitis or cirrhosis. Kupffer cells are considered to be target cells for *C. burnetii*. Involvement in bone and arteries has also been reported. Patients who have had acute Q fever may also develop the chronic form as soon as 1 year or as long as 20 years after initial infection.

### Risk to Pregnant Women

<table>
<thead>
<tr>
<th>1</th>
<th>Most asymptomatic</th>
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<tbody>
<tr>
<td>2</td>
<td>Transplacental transmission</td>
</tr>
<tr>
<td>3</td>
<td>Reported complications</td>
</tr>
<tr>
<td>4</td>
<td>- In-utero death</td>
</tr>
<tr>
<td>5</td>
<td>- Premature birth</td>
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<tr>
<td>6</td>
<td>- Low birth weight</td>
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<tr>
<td>7</td>
<td>- Placentitis</td>
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</tbody>
</table>

Pregnant women who become infected by *C. burnetii* are typically asymptomatic. However, the organism can be transplacentally transmitted. Depending on the timing of infection abortion, neonatal death, premature birth, low birth weight, or placentitis may occur. The greatest risk is during the first trimester. Pregnant women are also at a greater risk of developing chronic Q fever infection. Pregnant women with Q fever may pose a degree of risk to medical staff.

### Prognosis

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<thead>
<tr>
<th>1</th>
<th>Usually self-limiting</th>
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<tbody>
<tr>
<td>2</td>
<td>Only 2% develop severe disease</td>
</tr>
<tr>
<td>3</td>
<td>Active chronic disease</td>
</tr>
<tr>
<td>4</td>
<td>- Usually fatal if left untreated</td>
</tr>
<tr>
<td>5</td>
<td>- Fatality for endocarditis: 45 to 65%</td>
</tr>
<tr>
<td>6</td>
<td>- 50 to 60% need valve replacement</td>
</tr>
<tr>
<td>7</td>
<td>Case-fatality rate: &lt;1 to 2.4%</td>
</tr>
</tbody>
</table>

Q fever is usually a self-limiting illness and most cases resolve within 2 days to 2 weeks. Approximately 50 to 60% of cases are thought to be asymptomatic, and complications from the acute form of disease are rare. Only 2% of persons infected with *Coxiella burnetii* develop severe disease and require hospitalization. In general, the mortality rate is 1% or lower if treated. Active chronic disease is usually fatal if untreated. In patients with endocarditis, the fatality rate can range from 45 to 65%; additionally, 50 to 60% need valve replacement surgery. Because severe disease is rare the overall case-fatality rate for Q fever ranges from <1 to 2.4%.

### Diagnosis

<table>
<thead>
<tr>
<th>1</th>
<th>Serology (rise in titer)</th>
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<tbody>
<tr>
<td>2</td>
<td>- IFA, CF, ELISA, microagglutination</td>
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<tr>
<td>3</td>
<td>DNA detection methods</td>
</tr>
<tr>
<td>4</td>
<td>- PCR</td>
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<tr>
<td>5</td>
<td>Isolation of organism</td>
</tr>
<tr>
<td>6</td>
<td>- Risk to laboratory personnel</td>
</tr>
<tr>
<td>7</td>
<td>- Rarely done</td>
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</table>

In humans, Q fever is usually diagnosed by serology (rise in antibody titer levels) which can be done as early as the second week of illness. There are a variety of serological tests for Q fever including IFA (immunofluorescence assay), CF (complement fixation), ELISA (enzyme-linked immunosorbent assay), and microagglutination. The indirect IFA is the most dependable and widely used method. *C. burnetii* may also be identified in infected tissue by IHC (immunohistochemistry) and DNA detection methods (PCR-polymerase chain reaction). Isolation of the organism is rarely done due to the risk *C. burnetii* poses for laboratory personnel. Clinical signs and patient history can also aid in diagnosis.
The antibiotic treatment of choice is doxycycline. Antibiotic treatment is most effective when initiated within the first three days of illness. For chronic disease, treatment may be necessary for 2 to 3 years. Doxycycline and quinolones are contraindicated in pregnant women but long term therapy with co-trimoxazole (trimethoprim/sulfamethoxazole combination) has prevented fetal death in some cases. Persons recovering from Q fever are thought to develop long-lasting (possibly lifelong) immunity.

Now let's look at a few case studies of Q fever. A 46 year old male dairy farmer from Georgia reported a sudden onset of fever, chills, cough, and weight loss. A physician initially diagnosed influenza in the patient. Two weeks later the symptoms still persisted and the patient presented to the emergency room where he was again diagnosed with influenza.

The emergency room doctor then referred the patient to an infectious disease specialist. The infectious disease specialist tested the patient for Q fever and he was positive. The patient took a five day course of a fluoroquinolone and symptoms resolved within 2 weeks. Although the patient owned several dairy cows, no recent calvings had occurred at his farm. Two beef cattle herds (approximately 35 animals per herd) were pastured across the road from the patient’s farm. Fourteen animals from the neighboring beef herds were then tested; two animals were found to be positive for Q fever.

In 1985, a cluster of Q fever cases occurred in Nova Scotia, Canada. Most of the affected persons had symptoms of fever, sweats, chills, fatigue, myalgia and headache. Seventeen of the patients developed a cough and 14 had pneumonia. Epidemiological investigation revealed that 25 patients were exposed to a cat that had given birth to stillborn kittens 2 weeks prior. The majority of human cases lived or worked in 4 buildings near the apartment where the cat lived, and the cat visited the other buildings frequently. The cat tested positive for antibodies to C. burnetii. All human patients recovered uneventfully.

[Source: MMWR October 18, 2002/51(41);924-927]
Sheep, cattle, and goats are the most common reservoirs of Q fever. The incubation period for animals is variable. Affected animals may be asymptomatic; when clinical disease occurs, reproductive failure is usually the only symptom seen. This may include abortions, stillbirths, retained placentas, infertility, weak newborns, and mastitis in dairy cattle. Lambs born following *Coxiella* abortions may be carried to term. However, ewes can remain chronically infected and continue to shed organisms. Organisms may be shed in milk and feces for several days after parturition.

Dogs, cats, horses, pigs, and most mammals and fowl species may carry *C. burnetii*. Animals may be infected by a tick bite, consumption of placentas or milk from infected ruminants, or by the aerosol route. Most infections are asymptomatic; however, the most symptoms are related to reproductive failure. Still births and weak offspring are commonly reported.

Placentitis is the most characteristic lesion in ruminants. The placenta is typically leathery and thickened. It may contain large amounts of creamy, white-yellow exudate at the edges of cotyledons and in the intercotyledonary area. Lesions in aborted fetuses are usually non-specific.

*C. burnetii* can be detected in vaginal discharges, the placenta or its fluids, aborted fetuses, milk, urine and feces. Organism identification can be accomplished with Modified Ziehl-Neelson or Gimenez stains, but are usually not detected by Gram stain. IHC can also confirm bacterial identity. PCR techniques are also available in some laboratories. A number of serological tests are also available (i.e., immunofluorescence (IFA), enzyme-linked immunosorbent assays (ELISA) and complement fixation (CF)). The complement fixation test is done most commonly. Although isolation of the organism can be accomplished in a variety of methods, it is dangerous to laboratory personnel, and must be completed in a Biosafety-Level 3 laboratory. It is therefore rarely used.
Little is known about the effectiveness of treating animals with antibiotics. Tetracycline has been given in water in the weeks preceding parturition in enzootic herds. This is believed to help reduce shedding in birthing materials. Antimicrobial therapy may not eradicate the “carrier” state of *C. burnetii* infection, but may suppress the number of abortion.

Information on the prevalence of Q fever in animal species is limited. In endemic areas, (i.e. areas of California), it was found that 18 to 55% of sheep and up to 82% of cows in some dairies had antibodies to *C. burnetii*. In sheep, abortion can affect 5 to 50% of the flock. Death is rare in naturally occurring infections.

Good husbandry plays an important role in the prevention and control of this disease. Tick prevention should be used to help prevent the spread of this disease. Animals that are about to give birth should be separated from the rest of the herd. Fetal membranes and aborted fetuses should be disposed of immediately either by burying or burning. New or sick animals should be separated from the rest of the herd until it can be determined that the animals are not contagious or do not pose an infectious risk. Vaccinations for humans and animals have been developed for this disease. However, they are not currently licensed for use in the United States.

Pasteurization of milk from cows, sheep, and goats is important in stopping the spread of Q fever by contaminated milk sources. The amount of *C. burnetii* in the environment can be greatly reduced by thorough cleaning. A 10% bleach solution should be used to disinfect areas after cleaning where animals give birth. Eradication is impossible because of environmental stability, infectivity for wild animals, asymptomatic and carrier state in animals and people, and arthropods.
Q Fever as a Biological Weapon

- Accessibility
- Low infectious dose
- Stable in the environment
- Aerosol transmission
- WHO estimate
  - 5 kg agent released on 5 million persons
  - 125,000 ill - 150 deaths
  - Could travel downwind for over 20 km

Because of its highly infectious nature, stability in the environment, and aerosol route of transmission, *C. burnetii* can be considered a potential agent of bioterrorism. Although overall mortality associated with the disease is low, it could be considered a debilitating agent. The World Health Organization (WHO) estimated that if Q fever was aerosolized in a city of approximately 5 million people there would be 125,000 ill and 150 deaths. They estimated that the agent could travel downwind for greater than 20 km.

Additional Resources

- World Organization for Animal Health (OIE)
  - www.oie.int
- U.S. Department of Agriculture (USDA)
  - www.aphis.usda.gov
- Center for Food Security and Public Health
  - www.cfsph.iastate.edu
- USAHA Foreign Animal Diseases ("The Gray Book")
  - www.usaha.org/pubs/fad.pdf

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