
Copyright © 2013 American Society of Tropical Medicine and Hygiene

A copy can be downloaded for personal non-commercial research or study, without prior permission or charge

The content must not be changed in any way or reproduced in any format or medium without the formal permission of the copyright holder(s)

When referring to this work, full bibliographic details must be given

http://eprints.gla.ac.uk/78630/

Deposited on: 25 April 2013
Coxiella burnetii in Humans, Domestic Ruminants, and Ticks in Rural Western Kenya


Boyd Orr Centre for Population and Ecosystem Health, Institute of Biodiversity, Animal Health and Comparative Medicine, University of Glasgow, Glasgow, United Kingdom; Department of Veterinary Tropical Diseases, University of Pretoria, Onderstepoort, South Africa; Jomo Kenyatta University of Agriculture and Technology, Nairobi, Kenya; Viral and Rickettsial Diseases Department, Naval Medical Research Center, Silver Spring, Maryland; School of Health, Sports and Biosciences, University of East London, London, United Kingdom; Kenya Medical Research Institute/Centers for Disease Control and Prevention-Kenya, Kisumu, Kenya; Global Disease Detection Division, CDC-Kenya, Nairobi, Kenya

Abstract. We conducted serological surveys for *Coxiella burnetii* in archived sera from patients that visited a rural clinic in western Kenya from 2007 to 2008 and in cattle, sheep, and goats from the same area in 2009. We also conducted serological and polymerase chain reaction-based surveillance for the pathogen in 2009–2010, in human patients with acute lower respiratory illness, in ruminants following parturition, and in ticks collected from ruminants and domestic dogs. Antibodies against *C. burnetii* were detected in 30.9% (*N* = 246) of archived patient sera and in 28.3% (*N* = 463) of cattle, 32.0% (*N* = 378) of goats, and 18.2% (*N* = 159) of sheep surveyed. Four of 135 (3%) patients with acute lower respiratory illness showed seroconversion to *C. burnetii*. The pathogen was detected by polymerase chain reaction in specimens collected from three of six small ruminants that gave birth within the preceding 24 hours, and in five of 10 pools (50%) of *Haemaphysalis leachi* ticks collected from domestic dogs.

INTRODUCTION

Q fever is a disease of humans caused by infection with the obligate intracellular bacterial pathogen, *Coxiella burnetii*. It is a zoonotic infection, typically transmitted from animal hosts to humans through inhalation of contaminated aerosols or ingestion of infected animal products such as milk or cheese.1

Ticks have also been implicated as vectors, and over 40 species of ticks have been found naturally infected with the agent.1 *Coxiella burnetii* has a wide host range, including wild and domestic mammals, birds, reptiles, and arthropods.2 Domestic ruminants (primarily goats, cattle, and sheep) represent the most frequent source of human infection,1 although transmission from dogs and cats is also documented.4,5 The uterus and mammary glands of female animals are sites of chronic *C. burnetii* infection, and infected females may shed large amounts of bacteria into the environment during parturition or spontaneous abortion.2 Once shed, the organisms may remain infective in the environment for several months.1 Infection in humans, usually by inhalation, may be asymptomatic (up to 60% of infected individuals) or may manifest clinically after an incubation period ranging between 1 and 3 weeks.1,6 Clinical signs of acute Q fever include fever of 2–14 days’ duration, atypical pneumonia, and/or hepatitis.1 Although the disease is typically self-limiting, severe debilitating illness requiring hospitalization can occur in a small proportion (2–5%) of acutely infected cases.6 Chronic disease may develop following infection, particularly in patients with predisposing conditions such as preexisting cardiac valvulopathy, pregnancy, or immunosuppression.3 Common manifestations of chronic disease include endocarditis and vascular infection.1 *Coxiella burnetii* displays antigenic (phase) variation associated with loss of virulence and mutational variation in the lipopolysaccharide.1 High levels of antibodies to phase I antigens are detected during chronic Q fever, whereas antibodies to phase II antigens are produced in acute disease.6

*Coxiella burnetii* is found worldwide, with the exception of New Zealand.7 A recent large outbreak in the Netherlands involved at least 3,523 human cases from 2007 through 2009,8 and was characterized by a high rate of hospitalization, with 20% of notified cases admitted to hospital in 2008–2009.9 Pneumonia was the predominant clinical presentation.9 The Netherlands outbreak has been linked to the increase in the country’s dairy goat population, which more than doubled in size between 2000 and 2009, and highlights the public health risks of Q fever epidemics posed by domestic ruminants. Despite the high-profile nature of some Q fever outbreaks, and the attention that *Coxiella burnetii* has received as a potential bioterrorism agent,10 information on the prevalence of infection in sub-Saharan Africa is scant.11,12 In Kenya, serological evidence of Q fever in patients with acute febrile and respiratory illness was shown in the 1950s.13 Other studies showed the prevalence of antibodies to *C. burnetii* among Kenyans to range between 10% and 20%.14,15 A more recent investigation found that four people (8%) of a group of 50 travelers to Kenya contracted Q fever,16 and in another recent study investigators diagnosed acute Q fever in 5% of febrile patients admitted to two hospitals from September 2007 to August 2008, in neighboring northern Tanzania.17 Among domestic ruminants in Kenya, the prevalence of antibodies was reported as 7–57% in cattle and 33–34% in goats.14,15,18,19 There are no other recent reports of investigations into disease prevalence in resident human or livestock populations, or any information on the relationship between prevalence in human and animal populations.

The establishment of a population-based infectious disease surveillance program in western Kenya20 provided a valuable opportunity for generating domestic animal prevalence data that could be linked with human health outcomes. To assess the current status of Q fever among humans and infection prevalence in domestic ruminants, we conducted surveys for *C. burnetii* in cattle, goats, sheep, and ticks in this rural agropastoral community, and tested specimens from human patients presenting to a clinic in the same area for Q fever.
MATERIALS AND METHODS

**Study site.** All data were collected in Asembo within Rarieda (formerly Bondo) District in western Kenya in 2007–2010. This rural site on the eastern shore of Lake Victoria falls within a health and demographic surveillance system (HDSS) that has been run by the Kenya Medical Research Institute (KEMRI) and U.S. Centers for Disease Prevention and Control (CDC) since 2001. The HDSS collects household demographic and socioeconomic data three times per year, and includes information on the number of livestock owned. Households are clustered into compounds composed of related family units, with most compounds having between one and five family units. The primary economic activity is subsistence smallholder agro-pastoralism and fishing. In this area, 44% of households own cattle (mean number owned: 1.84) and 43% own at least one sheep or goat (mean number owned: 2.12; HDSS data for 2008, unpublished).

The Kenyan International Emerging Infections Program of KEMRI/CDC has conducted population-based infectious disease surveillance (PBIDS) of people in Asembo since late 2005, with between 23,500 and 25,000 people under surveillance in 33 villages. Participants enrolled in PBIDS have household visits to determine health status every 2 weeks, and receive free medical care for all acute illnesses at a centrally located clinic, St. Elizabeth Lwak Mission Hospital (henceforth Lwak Hospital). Specimen collection and diagnostic testing are focused on determining etiologies for four infectious disease syndromes, namely acute respiratory infection, diarrhea, jaundice, and febrile illness. Sera are collected from patients participating in PBIDS presenting to Lwak Hospital and meeting case definitions for acute respiratory illness, jaundice or acute febrile illness, or who are hospitalized for any non-traumatic illness. Between January 2007 and January 2010, 3,948 serum specimens were collected and archived at −80°C at the KEMRI/CDC laboratory in Kisumu.

**Ethical review.** The collection of specimens from humans was approved by the KEMRI Ethical Review Committee (protocol no. 932) and CDC Institutional Review Board (protocol no. 4566), and from animals by the KEMRI and CDC Animal Care and Use Committees (protocol no. 1191 and 1562BRETBDX, respectively). Written informed consent was obtained from all patients or animal owners before specimen collection.

**Human serosurvey.** To assess previous exposure to *C. burnetii* in humans, we retrospectively identified paired serum specimens collected from ALRI cases occurring from July 2009 through January 2010. Convalescent-phase serum specimens were screened for IgG antibodies against *C. burnetii* phase II antigen by indirect enzyme-linked immunosorbent assay (ELISA) (Panbio, Brisbane, Australia) using the manufacturer’s recommended cutoff. End-point titers of IgG antibodies in acute and convalescent sera were determined by IFA assay for all pairs in which the convalescent-phase tested positive or equivocal on ELISA, with serial dilutions starting from 1:16. The operator who performed the IFA was blind to the status (acute or convalescent) of the specimen. A ≥ 4-fold increase in IFA titer to *C. burnetii* phase II antigen defined acute Q fever, and a titer ≥ 1:1,024 to *C. burnetii* phase I antigen in either acute- or convalescent-phase sera defined possible chronic Q fever.

**Domestic ruminants cross-sectional study.** Data were collected from January through May 2009. The study population comprised all domestic ruminants (cattle, goats, and sheep) living within compounds in the 33 villages of the PBIDS. The sampling frame was compiled from data on the number of each species of domestic animal within the compounds, as reported at the most recent HDSS visit for which data were available. Three hundred livestock-owning compounds (LOCs) were randomly selected from the sampling frame of 4,528 LOCs, with a LOC defined by ownership of one or more animals of any of the following species: cattle, sheep, goats, chickens. The heads of the selected compounds were approached and offered enrollment. If livestock were no longer owned, replacement LOCs was then randomly selected from the same village.

Specimens were collected from a maximum of three randomly selected animals of any age from each of the species of domestic ruminants present at enrolled compounds. Blood was collected by venipuncture of the jugular vein into plain vacutainers (BD, Franklin Lakes, NJ). Vaginal swabs were collected from adult females only, using nylon flocked swabs (eSwabs, Copan Innovation, Brescia, Italy) and placed in universal transport medium (Copan Innovation). Blood collection tubes were kept at ambient temperature for 15–30 minutes, and then transported to the field laboratory on ice with the vaginal swabs. At the field laboratory, blood tubes were centrifuged at 2,000–3,000 revolutions per minute for

**Human respiratory patients.** Under the PBIDS clinic surveillance protocol, blood specimens are collected from patients who meet the case definition for acute lower respiratory infection (ALRI). This is defined in persons 5 years of age or older as cough, difficulty breathing or chest pain and either documented axillary temperature ≥ 38.0°C or oxygen saturation < 90%, and in children under 5 years of age as cough or difficulty breathing with one of the following: elevated respiratory rate for age (non-severe pneumonia), or lower chest wall indrawing, stridor, oxygen saturation < 90%, maternal report of convulsions, inability to drink or breastfeed, or vomiting everything, or on exam lethargy or unconsciousness (severe and very severe pneumonia). All patients who meet the case definition have acute-phase sera collected and archived at the time of presentation, and patients are requested to return to the clinic after 4–6 weeks for the collection of convalescent-phase serum specimens. Paired sera are tested serologically for a range of respiratory pathogens.

To determine the prevalence of *C. burnetii* infection in these patients with respiratory illness, we retrospectively identified paired serum specimens collected from ALRI cases occurring from July 2009 through January 2010. Convalescent-phase serum specimens were screened for IgG antibodies against *C. burnetii* phase II antigen by indirect enzyme-linked immunosorbent assay (ELISA) (Panbio, Brisbane, Australia) using the manufacturer’s recommended cutoff. End-point titers of IgG antibodies in acute and convalescent sera were determined by IFA assay for all pairs in which the convalescent-phase tested positive or equivocal on ELISA, with serial dilutions starting from 1:16. The operator who performed the IFA was blind to the status (acute or convalescent) of the specimen. A ≥ 4-fold increase in IFA titer to *C. burnetii* phase II antigen defined acute Q fever, and a titer ≥ 1:1,024 to *C. burnetii* phase I antigen in either acute- or convalescent-phase sera defined possible chronic Q fever.

Blood was collected by venipuncture of the jugular vein into plain vacutainers (BD, Franklin Lakes, NJ). Vaginal swabs were collected from adult females only, using nylon flocked swabs (eSwabs, Copan Innovation, Brescia, Italy) and placed in universal transport medium (Copan Innovation). Blood collection tubes were kept at ambient temperature for 15–30 minutes, and then transported to the field laboratory on ice with the vaginal swabs. At the field laboratory, blood tubes were centrifuged at 2,000–3,000 revolutions per minute for

**Human respiratory patients.** Under the PBIDS clinic surveillance protocol, blood specimens are collected from patients who meet the case definition for acute lower respiratory infection (ALRI). This is defined in persons 5 years of age or older as cough, difficulty breathing or chest pain and either documented axillary temperature ≥ 38.0°C or oxygen saturation < 90%, and in children under 5 years of age as cough or difficulty breathing with one of the following: elevated respiratory rate for age (non-severe pneumonia), or lower chest wall indrawing, stridor, oxygen saturation < 90%, maternal report of convulsions, inability to drink or breastfeed, or vomiting everything, or on exam lethargy or unconsciousness (severe and very severe pneumonia). All patients who meet the case definition have acute-phase sera collected and archived at the time of presentation, and patients are requested to return to the clinic after 4–6 weeks for the collection of convalescent-phase serum specimens. Paired sera are tested serologically for a range of respiratory pathogens.

To determine the prevalence of *C. burnetii* infection in these patients with respiratory illness, we retrospectively identified paired serum specimens collected from ALRI cases occurring from July 2009 through January 2010. Convalescent-phase serum specimens were screened for IgG antibodies against *C. burnetii* phase II antigen by indirect enzyme-linked immunosorbent assay (ELISA) (Panbio, Brisbane, Australia) using the manufacturer’s recommended cutoff. End-point titers of IgG antibodies in acute and convalescent sera were determined by IFA assay for all pairs in which the convalescent-phase tested positive or equivocal on ELISA, with serial dilutions starting from 1:16. The operator who performed the IFA was blind to the status (acute or convalescent) of the specimen. A ≥ 4-fold increase in IFA titer to *C. burnetii* phase II antigen defined acute Q fever, and a titer ≥ 1:1,024 to *C. burnetii* phase I antigen in either acute- or convalescent-phase sera defined possible chronic Q fever.

**Domestic ruminants cross-sectional study.** Data were collected from January through May 2009. The study population comprised all domestic ruminants (cattle, goats, and sheep) living within compounds in the 33 villages of the PBIDS. The sampling frame was compiled from data on the number of each species of domestic animal within the compounds, as reported at the most recent HDSS visit for which data were available. Three hundred livestock-owning compounds (LOCs) were randomly selected from the sampling frame of 4,528 LOCs, with a LOC defined by ownership of one or more animals of any of the following species: cattle, sheep, goats, chickens. The heads of the selected compounds were approached and offered enrollment. If livestock were no longer owned, replacement LOCs was then randomly selected from the same village.

Specimens were collected from a maximum of three randomly selected animals of any age from each of the species of domestic ruminants present at enrolled compounds. Blood was collected by venipuncture of the jugular vein into plain vacutainers (BD, Franklin Lakes, NJ). Vaginal swabs were collected from adult females only, using nylon flocked swabs (eSwabs, Copan Innovation, Brescia, Italy) and placed in universal transport medium (Copan Innovation). Blood collection tubes were kept at ambient temperature for 15–30 minutes, and then transported to the field laboratory on ice with the vaginal swabs. At the field laboratory, blood tubes were centrifuged at 2,000–3,000 revolutions per minute for
7–10 minutes, and serum pipetted into cryovials. Sera and swabs were then transported on ice to the KEMRI-CDRC laboratory and stored at −80°C until testing. Ruminant serum samples were tested for the presence of IgG antibodies to *C. burnetii* by indirect ELISA (Chekit Q fever ELISA, IDEXX), following the manufacturer’s instructions. DNA was extracted from vaginal swab specimens using QIAamp DNA Mini Kit (Qiagen, Valencia, CA) in a biosafety level 3 laboratory, and tested by quantitative real-time polymerase chain reaction (qPCR) assay using the *IS1111* gene target for *C. burnetii*, as previously described.24

**Domestic ruminant births/abortions.** To further investigate shedding of *C. burnetii* by domestic ruminants, we conducted surveillance for births and abortions in the ruminant population in the PBIDS area during 1 week in August 2009, as part of a pilot study for a larger surveillance program on animal disease syndromes. A network of animal health reporters (AHRs) was established in the 33 villages in the PBIDS (1–2 AHRs per village), through which all livestock owners could report births or abortions in their ruminant herds. The AHRs then contacted the study field veterinary team by mobile phone, who responded to the case as soon as possible. Available samples were collected by the veterinary team, including blood and vaginal swabs from dams, placenta, and liver, lungs, spleen, and brain of aborted fetuses or stillborn animals. DNA was extracted and tested as described previously.

**Collection of arthropods.** All domestic ruminants from which blood was collected were also inspected for ticks. In addition and when present, up to three domestic dogs in each enrolled compound were also inspected. In each animal, we examined specific body sites (ruminants: ear, dewlap, shoulder, belly, groin, udder, and perineum; dogs: ear, neck, mape, belly, and groin) and collected 1–2 non-engorged adult ticks of different genera per site (although only *Amblyomma* ticks were tested from ruminants). Ticks were preserved in 70% ethanol at room temperature until delivery to the laboratory, and then stored in ethanol at −80°C. Ticks were pooled by individual host animal and tick species (1–6 ticks per pool). Ticks of the genus *Amblyomma* collected from ruminants and all ticks collected from dogs were identified using published entomological keys25; after identification, ticks were washed in molecular-grade water and mechanically disrupted using a bead mill (Qiagen TissueLyser LT). Genomic DNA was extracted using QIAamp blood and tissue kits (Qiagen) according to the manufacturer’s instructions, using a final elution volume of 100 μL, and tested by *IS1111* qPCR.24

**RESULTS**

**Human serosurvey.** A total of 248 and 246 archived sera were tested by IFA for IgG antibodies against *C. burnetii* phase I and phase II antigen, respectively. The IgG antibodies against *C. burnetii* phase I antigen were detected in 58 specimens (23.4%, exact binomial 95% confidence intervals [CI]: 18.3–29.2%), and IgG antibodies against phase II antigen were detected in 76 specimens (30.9%, 95% CI: 25.2–37.1%). All the specimens that reacted with phase I antigen also reacted with phase II antigen.

**Human respiratory patients.** Of 2,246 patients who met the case definition for ALRI from July 2009 through January 2010, 493 (22%) returned for collection of convalescent sera. Of these, 135 pairs were available for testing. Convalescent-phase sera were screened for IgG antibodies against *C. burnetii* phase II antigen by indirect ELISA. Nineteen patients (14.1% of ALRI patients, 95% CI: 8.7–21.1%) had positive titers on ELISA. Of these, 4 (3% of ALRI patients, 95% CI: 0.8–7.4%) were diagnosed serologically with acute Q fever by IFA as determined by a ≥ 4-fold rise between acute and convalescent sera. All four patients were males, 7–31 years of age, who presented with fever (documented axillary temperatures of 38.2–39.5°C), chills, cough, and headache (Table 1). The reported duration of illness ranged from 2 to 10 days. All four patients were from households that owned one or more domestic ruminants around the time of illness. One of the patients (#2 in Table 1) also showed seroconversion to influenza B virus. No patients were diagnosed with possible chronic Q fever.

**Domestic ruminant births/abortions.** Six births (five in goats and one in a sheep) and one abortion in a cow were detected during the week-long surveillance. In all cases, specimens were collected within 24 hours of the event. Three births in goats resulted in one stillborn kid in each, along with one to two live offspring. Samples from the placenta of one of these cases, and vaginal swabs from a sheep and a goat in the same compound that gave birth on consecutive days, were positive for *C. burnetii* on the qPCR assay for the *IS1111* gene target.

**Arthropods.** A total of 258 adult *Amblyomma variegatum* ticks were collected from 162 cattle in the study area, and pooled by individual host. *Coxiella burnetii* DNA was detected by *IS1111* PCR in vaginal swabs from 5 of 233 (2.1%) cattle, 5 of 222 (2.3%) goats, and 4 of 85 (4.7%) sheep.

**Domestic ruminant births/abortions.** Six births (five in goats and one in a sheep) and one abortion in a cow were detected during the week-long surveillance. In all cases, specimens were collected within 24 hours of the event. Three births in goats resulted in one stillborn kid in each, along with one to two live offspring. Samples from the placenta of one of these cases, and vaginal swabs from a sheep and a goat in the same compound that gave birth on consecutive days, were positive for *C. burnetii* on the qPCR assay for the *IS1111* gene target.

**Arthropods.** A total of 258 adult *Amblyomma variegatum* ticks were collected from 162 cattle in the study area, and pooled by individual host. *Coxiella burnetii* DNA was detected by *IS1111* PCR in 4 of 162 (2.5%) pools. In dogs, 102 ticks were collected from 36 animals, resulting in pools of the following species: *Rhipicephalus sanguineus* (15 pools), *Rhipicephalus appendiculatus* (9 pools), unspeciated *Rhipicephalus* (10 pools), *A. variegatum* (10 pools), *Haemaphysalis leachi* (10 pools), and

---

**Table 1.** Clinical features of four patients from rural western Kenya with acute lower respiratory illness, diagnosed serologically with acute Q fever

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Male</td>
<td>31</td>
<td>Y</td>
<td>38.3°C</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>N</td>
<td>N</td>
<td>N</td>
<td>N</td>
<td>N</td>
<td>N</td>
<td>N</td>
</tr>
<tr>
<td>2</td>
<td>Male</td>
<td>22</td>
<td>Y</td>
<td>38.7°C</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>N</td>
<td>N</td>
<td>N</td>
<td>N</td>
<td>N</td>
<td>N</td>
<td>N</td>
</tr>
<tr>
<td>3</td>
<td>Male</td>
<td>10</td>
<td>Y</td>
<td>38.2°C</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>N</td>
<td>N</td>
<td>N</td>
<td>N</td>
<td>N</td>
<td>N</td>
<td>N</td>
</tr>
<tr>
<td>4</td>
<td>Male</td>
<td>7</td>
<td>Y</td>
<td>39.5°C</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>N</td>
<td>N</td>
<td>N</td>
<td>N</td>
<td>N</td>
<td>N</td>
<td>N</td>
</tr>
</tbody>
</table>

---

Of 2,246 patients who met the case definition for ALRI from July 2009 through January 2010, 493 (22%) returned for collection of convalescent sera.
Rhizophalus (Boophilus) decoloratus (5 pools). The number (and percent) of pools positive using the IS1111 PCR assays were: Rh. sanguineus 2 (20%), Rh. appendiculatus 1 (11.1%), unspeciated Rhizophalus 2 (20%), A. variegatum 2 (20%), H. leachi 5 (50%), and Rh. (Boophilus) decoloratus 1 (20%).

**DISCUSSION**

We identified Q fever as a possible cause of acute lower respiratory illness among rural residents in western Kenya. In addition, we found a high prevalence of antibodies to C. burnetii in domestic ruminants in the same area, with high rates of shedding of the organism shortly after parturition in a small sample of animals.

The prevalence of antibodies to C. burnetii in patients attending Lwak Hospital is similar to that reported in several population-based studies elsewhere in sub-Saharan Africa (17–37%), indicating high levels of exposure to the pathogen. In addition to the recent study in northern Tanzania in which acute Q fever was diagnosed in 5.0% of febrile inpatients, another study reported recent infection with C. burnetii in 9.5% of febrile patients >5 years of age in two urban areas in Mali. A literature review did not identify any recent investigations of Q fever as a cause of respiratory illness in Africa, although in the report of Q fever in travelers returning from Kenya, the two symptomatic patients both had respiratory involvement.

In addition to ALRI, it is likely that Q fever also contributes to the burden of acute febrile illness in patients in our study area. Both ALRI and febrile illness are managed empirically in this setting, due largely to a lack of reliable rapid diagnostic tests for diseases other than malaria. Although malaria may be responsible for >50% of febrile illness episodes in this holoendemic area, knowledge of the full range of prevalent etiologies is important to improve the empiric treatment of this syndrome. Doxycycline is considered the preferred antibiotic for the treatment of acute Q fever, and there is evidence that newer macrolides and fluoroquinolones are also effective. Notably, C. burnetii is resistant in vitro to those antimicrobial agents that are typically used for the empirical treatment of ALRI in Kenya (penicillin and its derivatives, and aminoglycosides). A diagnosis of Q fever pneumonia should therefore be considered in patients not responding to first-line antimicrobial therapy, particularly in those with exposure to known risk factors.

Inhalation of fomites contaminated by the parturient fluids of infected animals is the main mode of human infection with C. burnetii. Despite a very small sample, a high proportion of small ruminants in our study were found to shed the pathogen at parturition (2 of 5 goats and 1 of 1 sheep sampled). The stratified age-seroprevalence curves in ruminants show an increase in prevalence with age until around 2–3 years, followed by a plateau. This pattern has been observed elsewhere and is consistent with cumulative exposure to the pathogen from a young age, with the highest prevalence of antibody in reproductively mature animals.

Coxiella burnetii was detected by PCR in all species of ticks studied. Ticks become infected during the transient bacteremia that occurs in the vertebrate host early after infection. In our study, the highest prevalence of infection with C. burnetii was found in H. leachi (the yellow dog tick), one of two ticks adapted to feeding on domestic dogs in tropical and subtropical areas. Detection of C. burnetii in this species has been reported in Kenya previously; these findings suggest that domestic dogs may play a role as reservoir hosts and sources of human infection with C. burnetii in Kenya, as has been reported elsewhere.

Our findings show that Q fever is a significant yet under-diagnosed cause of human respiratory illness in Kenya.
Domestic ruminants are sources of human infection, through direct contact or contamination of the environment during parturition or abortion. Although not assessed in this study, consumption of dairy products from infected ruminants is also likely to pose a risk. Although risk factor studies would be helpful to identify the principal modes of transmission to humans in this setting, our results support the likely efficacy of recognized public health and hygiene measures for mitigating the risk of transmission from domestic ruminants, such as confinement of domestic animals during parturition, avoidance of contaminated pastures and contact with placental material, and boiling or pasteurization of milk before consumption. This study also suggests that investigation of the role of domestic dogs and tick-borne transmission is warranted to identify potential sources of infection and risk factors that are currently not well recognized.

Received March 15, 2012. Accepted for publication December 15, 2012.

Acknowledgments: We thank the IEIP-Z field team (Samuel Asenso, Michael Oticeno, James Oyigo, and Pauline Oticeno) for the collection of ticks and specimens from domestic animals, and Immaculate Amadi and Catherine Sonye for assistance with laboratory testing.

Financial support: This research was supported by the Wellcome Trust, UK (grant no. 081828/B/06/Z) and U.S. Centers of Disease Control and Prevention and Global Emerging Infections Surveillance and Response System Program (work unit no. 188M.0931.001.A0074).

Disclaimer: The findings and conclusions are those of the authors. They do not necessarily represent the official policy or position of the Centers for Disease Control and Prevention-Kenya, International Emerging Infections Program, Kenya, E-mails: EOgola@kemricdc.org and sarah.cleaveland@glasgow.ac.uk. and Comparative Medicine, College of Medical, Veterinary and Life Sciences, University of Glasgow, United Kingdom, E-mails: S.Cutler@uel.ac.uk. Eric Ogola and Mutunzi Junghae, Kenya Medical Research Institute, Centers for Disease Control and Prevention-Kenya, International Emerging Infections Program, Kenya, E-mails: EGola@kemricdc.org and mjungahae@ke.cdc.gov. Daniel R. Feikin, Johns Hopkins Bloomberg School of Public Health and Centers for Disease Control and Prevention, Baltimore, MD, E-mail: dfeikin@jhsph.edu. Jo E. B. Halliday and Sarah Cleaveland, Institute of Biodiversity, Animal Health and Comparative Medicine, College of Medical, Veterinary and Life Sciences, University of Glasgow, United Kingdom, E-mails: jo.halliday@glasgow.ac.uk and sarah.cleaveland@glasgow.ac.uk. Robert F. Breiman, International Emerging Infections Program, Global Disease Detection Division, Centers for Disease Control and Prevention, Kenya, E-mail: rbreiman@ke.cdc.gov. M. Kariuki Njenga, Integrated Human-Animal Health Program, Global Disease Detection Division, Centers for Disease Control and Prevention, Kenya, E-mail: knjenga@ke.cdc.gov.

REFERENCES


