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Oroya Fever and Verruga Peruana: Bartonelloses Unique to South America

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Abstract: Bartonella bacilliformis is the bacterial agent of Carrión’s disease and is presumed to be transmitted between humans by phlebotomine sand flies. Carrión’s disease is endemic to high-altitude valleys of the South American Andes, and the first reported outbreak (1871) resulted in over 4,000 casualties. Since then, numerous outbreaks have been documented in endemic regions, and over the last two decades, outbreaks have occurred at atypical elevations, strongly suggesting that the area of endemicity is expanding. Approximately 1.7 million South Americans are estimated to be at risk in an area covering roughly 145,000 km² of Ecuador, Colombia, and Peru. Although disease manifestations vary, two disparate syndromes can occur independently or sequentially. The first, Oroya fever, occurs approximately 60 days following the bite of an infected sand fly, in which infection of nearly all erythrocytes results in an acute hemolytic anemia with attendant symptoms of fever, jaundice, and myalgia. This phase of Carrión’s disease often includes secondary infections and is fatal in up to 88% of patients without antimicrobial intervention. The second syndrome, referred to as verruga peruana, describes the endothelial-cell-derived, blood-filled tumors that develop on the surface of the skin. Verrugae are rarely fatal, but can bleed and scar the patient. Moreover, these persistently infected humans provide a reservoir for infecting sand flies and thus maintaining Bartonella bacilliformis in nature. Here, we discuss the current state of knowledge regarding this life-threatening, neglected bacterial pathogen and review its host-cell parasitism, molecular pathogenesis, phylogeny, sand fly vectors, diagnostics, and prospects for control.

Methods

Authors were included based upon their relevant expertise in the field. Available databases were searched with key words for applicable articles in both English and Spanish. All cited work was previously peer-reviewed and published elsewhere.

Janus: The Two Faces of Carrión’s Disease

A brief history

Bartonellosis, or Carrión’s disease, is a remarkable bi-phasic illness that manifests as either angiogenic lesions that occur primarily on the skin or as a severe hemolytic anemia. It is caused by Bartonella bacilliformis, a bacterium presumed to be transmitted between humans through bites of infected female phlebotomine sand flies, most notably Lutzomyia verrucarum [1–4]. Humans are the only known reservoir. Carrión’s disease is endemic to arid, high-altitude (approximately 600–3200 m) valleys in the Andes Mountains of Peru, Ecuador, and Colombia [5–10], coincident with the habitat of the sand fly vector [5]. Evidence for bartonellosis in the region predates Columbus. Skin lesions akin to those typical of bartonellosis are depicted on 2,000-year-old pre-Inca ceramics, and Bartonella-like bacteria have been observed in skin lesions taken from the mummified remains of a human sacrifice victim who died over 1,000 years ago [11]. Descriptions of these lesions, referred to as verruga peruana (VP), appear in the journals of Spanish conquistadors who arrived in South America during the 16th century, and in 1630, Gago de Vadillo published the first medical treatise on the disease [12].

Two events inexorably linked with the early history of bartonellosis occurred within 15 years of one another at the end of the 19th century. The first was a dramatic outbreak of hemolytic anemia among workers based in the upper Rimac valley, above Lima, during the construction of a railway line to the mining town of La Oroya in 1871. During the course of a few weeks at least 4,000 men died of what became known as Oroya Fever (OF), although the agent of the syndrome remained elusive. Second, the common etiology of OF and VP was revealed when, in 1985, a medical student named Daniel Carrión injected himself with exudate from a verruga lesion, then developed OF and died [12]. Carrión sacrificed his life to establish the common etiology of VP and OF and is one of the most heroic figures in Peruvian medical history. In honor of his contribution, South American bartonellosis is commonly known as Carrión’s disease. Finally, in 1905, Alberto Barton solved the riddle of the etiological agent of Carrión’s disease when he observed intracellular bacteria in blood smears from Oroya fever patients. These bacteria were subsequently named Bartonella bacilliformis in his honor.

Carrión’s disease is most common in Peru, where nearly all cases have occurred over the last 70 years in the Departments of Ancash, Cajamarca, Cuzco, Lima, and Amazonas [13]. Sporadic cases may also have occurred in Chile, Bolivia, and Guatemala.

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During the last two decades, several outbreaks have occurred at abnormal altitudes and in historically nonendemic regions of Ecuador, Colombia, and Peru (e.g., near sea level in Huaral and Ica, Peru [13,15]; the Utcubamba River Valley in Amazonas [16]; Urubamba Valley and Sacred Valley of the Incas near Cuzco [17,18]; and the high Amazonas jungle [19]). In addition, multiple outbreaks have occurred concurrently in endemic regions (e.g., Ancash Region, Peru; Zamora-Chinchipe Province, Ecuador) [5–8,20,21]. Children have been hardest hit by these outbreaks, with fatality rates of approximately 10% [13,15,22]. A 10-fold increase in cases (948 to 10,390) was reported in Peru from 1997–2005 [23], with 52 cases per 100,000 Peruvians in 1999 [13]. Reported rates are considerably higher in endemic regions of Peru, at 242 cases per 100,000 inhabitants (http://www.dge.gob.pe/). An estimate of the at-risk population for Carrión’s disease is 1.7 million people in an area of 145,000 km² [15]. In one prospective cohort study of a Peruvian community in an endemic area, an incidence rate of 12.7 per 100 person-years was estimated, reflecting a clustering of 70% of cases within 18% of homes [24].

Factors contributing to the emergence and geographical expansion of *B. bacilliformis* are unclear but include climatic events that favor sand fly propagation, such as increased temperature, humidity, and rainfall during El Niño cycles [21,25,26], and possibly additional sand fly species serving as competent vectors (see “Sand fly vectors,” below) [5,27–29]. In short, Carrión’s disease is a reemerging public health threat in several South American nations and to travelers who visit these countries [10,30].

**Oroya fever (OF)**

Carrión’s disease manifests as two distinct syndromes that occur independently or sequentially. The first phase, OF (also known as the acute, primary or hematic phase), is more common in children (>60% of cases [13,22,31]) in both endemic and nonendemic areas, and it is characterized by an acute bacteremia at about 60 days (range of 10–210 days; [32]) following the bite of an infected sand fly. From the inoculation site, *B. bacilliformis* colonizes the entire circulatory system, and nearly every erythrocyte is infected (Figure 1A). OF involves a severe reduction in hematocrit (>80% destruction) and acute hemolytic anemia from splenic culling of infected erythrocytes [33–35]. Patients with OF present for 1–4 weeks with an array of symptoms, including pallor, fever, anorexia, malaise, cardiac murmur, myalgia, headache, jaundice, tachycardia, and hepatomegaly. Complications are common during OF, and include high mortality in pregnant women and their unborn children, cardiovascular and neurological problems, respiratory infections, and arthralgia [22,36]. This phase of disease can be fatal in 40%–88% of infected patients without antimicrobial intervention [2,5,12,14,32,36,37] and is by far the highest reported case-fatality rate for any *Bartonella* species. Secondary infections are also common and potentially life-threatening, and they may arise due to an immune-compromised state that accompanies the disease. These include salmonellosis, toxoplasmosis, malaria, shigellosis, histoplasmosis, and pneumocystosis [5,14,37]. Even with appropriate antimicrobial therapy, fatality rates of 9%–11% are typical [13,15,38]. A milder form of OF has also been reported with a significantly lower case-fatality rate (0.7%), suggesting that virulence potential and associated disease severity may be strain-dependent [16]. *Bartonella rochalimae*, a distantly related *Bartonella* species isolated from a tourist who visited South America has been reported to cause a relatively mild febrile illness that could be confused with OF. Although associated symptoms are similar, no intraerythrocytic bacteria are observed in blood smears and there is only a slight drop in hematocrit and mild anemia [39].

![Figure 1. Clinical manifestations of Carrión’s disease.](https://www.plosntds.org/article/f1d97e647915f02f0d397d0b571f44c5)
Candidatus for some chronic bartonellosis cases in endemic areas [42]. Whether indicates that species other than \textit{B. bacilliformis} as \textit{Candidatus} [2,22,32]. The recent discovery of a new Phlebotomus verrucarum phlebotomine sand flies to humans living in the high mountain and joint pains, headache, and a chronic accompanied by fever and malaise, lymphadenopathy, acute bone lesions can bleed or scar the patient, and eruptions can be sand fly is a vector of human disease. Applied to the transmission results from bacterial invasion of capillary endothelium and generates skin, termed VP or Peruvian warts (Figure 1B). Tissue involvement certainty that a sand fly is a vector of a given pathogen. \textit{Candidatus} Bartonella ancashi can cause OF is unknown.

Sand fly vectors

\textit{B. bacilliformis} is believed to be transmitted by bites of phlebotomine sand flies to humans living in the high mountain valleys of Peru and neighboring countries of Andean South America. Townsend was the first to advance this view, and he identified \textit{Phlebotomus verrucarum} (now \textit{L. verrucarum}) as the vector of “Verruga” [43–45]. Experiments by Townsend and other workers provided strong circumstantial evidence of sand fly vectorship but no conclusive proof.

Having the suspicion based on circumstantial evidence that a suspected vector insect transmits a specific pathogen to man or other animals does not constitute incrimination. One must determine the role the insect plays in the natural history of the pathogen. Killick-Kendrick and Ward [46] outlined five criteria that must be fulfilled to declare with reasonable certainty that a sand fly is a vector of human disease. Applied to the transmission of \textit{B. bacilliformis}, these are:

1. The suspected vector sand fly must bite humans and be present in a place where humans become infected with \textit{B. bacilliformis}.
2. The distribution of the suspected vector should encompass the distribution of the disease in humans, and the sand fly must be sufficiently abundant to maintain pathogen transmission.
3. \textit{B. bacilliformis} should be isolated from wild-caught sand flies and be shown to be indistinguishable from the pathogen causing disease in humans in the same area.
4. It should be demonstrated that naturally or experimentally infected flies can maintain the infection through the complete extrinsic life cycle of the pathogen.
5. Experimental transmission of \textit{B. bacilliformis} by bite or some other means would generally be considered conclusive proof that a sand fly is a vector of a given pathogen.

Of these criteria, the first two are fairly easy to satisfy through observations of sand fly behavior in the field, and numerous workers have done so. The remaining three criteria are more difficult to satisfy and must be accomplished through painstaking analysis and experimentation to conclusively incriminate a sand fly species as a vector of \textit{B. bacilliformis}.

Beginning in 1926, Noguchi and associates conducted several experiments in which triturated, wild-caught sand flies from the “Verruga zone” in Peru were inoculated into monkeys to produce infection [47,48]. Around the same time, Battistini performed a series of experiments in which sand flies collected from the Verruga zone were released into a screened cage with a rhesus monkey. Although this was the first sand fly experiment in which the disease was transmitted by means other than inoculation, the sand flies were not observed and it was not determined whether transmission was by bite [49–51]. Between 1937 and 1939, Hettig and associates conducted several feeding experiments in which wild-caught sand flies were either released into small cages with a rhesus monkey, or placed in small feeding cages and strapped to the monkey’s shaved belly. Five monkeys became infected as determined by blood culture but no cultures were made from the sand flies [1]. Furthermore, uncertainty remained as to whether transmission was by bite or some other means such as by delection, as “…there existed opportunities for other types of contact than that of the mouthparts between sand fly and monkey, such as the deposit of fecal matter if such there should be” [1]. Later, these scientists isolated what they said was \textit{B. bacilliformis} from the proboscis of each of two “Phlebotomus verrucarum” (\textit{L. verrucarum}) collected from an epidemic focus at Huinco in the Santa Eulalia Valley [1]. Because culture methods used at that time were likely not sufficiently refined to distinguish between \textit{B. bacilliformis} and other \textit{Bartonella} species, there is a lingering question as to whether this was the same pathogen causing disease in humans. However, it is fair to say that Hettig’s team was the first to incriminate \textit{L. verrucarum} as a vector of \textit{B. bacilliformis}.

Attempts to culture \textit{B. bacilliformis} from sand fly guts were unsuccessful; hence, they were unable to demonstrate any developmental cycle in the sand fly that might explain the mechanism of transmission [1].

In preliminary experiments conducted recently by three of the authors (MFM, JMB, and PGL), \textit{L. verrucarum} from a laboratory colony (Walter Reed Army Institute of Research) were fed through a chick-skin membrane on human blood containing green fluorescent protein (GFP)-labeled \textit{B. bacilliformis}. Serial sections of putatively infected flies and microscopic examination of their guts under UV over seven days revealed massive proliferation of the bacteria within the blood meal and invasion of red blood cells during the first three days (Figure 2). The infection persisted for at least seven days in the gut of the sand fly. Contrary to what Hettig observed [1], no \textit{B. bacilliformis} were seen on the mouthparts or in any organ other than the midgut. Diuretic droplets deposited on glass cover slips following feeding contained no labeled bacteria, an indication that posterior stage transmission via diuresis is unlikely. Transmission by bite during a second blood meal was not attempted, nor were fecal droplets from the digested blood meal examined for \textit{B. bacilliformis} contamination. The mechanism of transmission of \textit{B. bacilliformis} by sand flies is still unclear and is the subject of ongoing investigations.

Other vector sand flies. Owing to discrepancies between the distribution of Carrión’s disease and \textit{L. verrucarum}, the existence of secondary vectors was suggested [27]. Whereas \textit{L. verrucarum} is limited to altitudes below approximately 3,200 m above sea level \textit{L. peruensis} has been collected in human habitations in endemic villages as high as 3,740 m ASL [27; 2005, Wilfredo Arque, unpublished data, personal communication]. Since Hettig’s landmark experiments, substantial evidence
has accumulated in Peru to also incriminate *L. peruensis*, which is highly anthropophilic and abundant or dominant in bartonellosis foci in Ancash and Cusco departments, respectively. In studies conducted during an outbreak in the Urubamba Valley, *L. peruensis* in large numbers and a few *L. pescei* were the only sand fly species found. *B. bacilliformis* was identified using PCR in 2% of *L. peruensis* analyzed [17]. In two other foci in Peru, 2%–8% of pools of *L. verrucarum* collected from patient houses in the Caraz area, Ancash Department, and 2% of pools of *L. peruensis* collected from patient houses in villages 70 km south of Cusco, Cusco Department were PCR positive for *B. bacilliformis* (2005, Sofia Romero, unpublished data, personal communication).

Besides *L. verrucarum* and *L. peruensis*, several other sand fly species are suspected vectors of *B. bacilliformis*. Caceres et al., working in the Peruvian provinces of Jaen, San Ignacio and Utcubamba, found *L. robusta* and *L. maranonensis* in great abundance in intradomiciliary habitats and, because of the absence of *L. verrucarum* and *L. peruensis*, considered these as likely vectors of *B. bacilliformis* in associated foci [28]. Elsewhere, on the eastern slope of the Andes in what is known as “Yungas,” or high jungle, Tejada et al. found that *L. serrana*, an avid man-biter, comprised 93.6% of sand flies collected during an investigation of a bartonellosis outbreak in Huamalíes, Huánuco Department [52].

In neighbor Andean countries of Colombia and Ecuador, where *L. verrucarum* and *L. peruensis* are absent, other species are suspected as vectors, but insufficient evidence has been produced to implicate them with certainty. Dominant species in Colombian districts where human bartonellosis has been reported, include *L. gomezi*, *L. panamensis*, and *L. serrana*, all of which are anthropophilic but none of which has ever been associated with *B. bacilliformis* transmission [6]. *L. colombiana* is considered the most likely vector of *B. bacilliformis* in Colombia based on its highly anthropophilic behavior and occurrence in all three departments of the country from which bartonellosis outbreaks have been reported [6]. The species that transmits this pathogen in Ecuador remains unknown [6].

**Spatial and temporal dynamics of vector sand flies.** Three-year longitudinal studies of suspected vector sand flies associated with bartonellosis outbreaks in Ancash and Cusco departments of Peru revealed that both *L. verrucarum* and *L. peruensis* exhibit unimodal annual temporal distribution patterns with lowest population densities occurring in June and July, corresponding to the “winter” dry season; and highest population...
densities in November, prior to the onset of the “summer” rainy season in December (Figure 3) [53]. It was further determined that sand fly population densities are directly correlated with average minimum ambient temperatures and relative humidity (Figure 3) [53]. As nighttime temperatures and relative humidity increase, sand fly activity also increases (Figure 4) [53]. These findings are consistent with the notion that the height of the B. bacilliformis transmission season, March to May, [24,26] corresponds to the descending side of the seasonal sand fly population peak, February to June, when the aggregate population is oldest and when the highest percentage of flies have had an opportunity to take at least one blood meal and are thus more likely to be infected.

In the Ancash and Cusco regions of Peru, approximately 80% of the flies were captured inside case houses, mostly in sleeping quarters, and 35%–40% of females were blood-fed. Being weak fliers, sand flies breed in close proximity to their blood-meal source, feeding at dusk and during the evening, when ambient temperatures drop and relative humidity rises [53]. L. verrucarum and L. peruensis both rest inside human dwellings and feed readily indoors, preferentially on humans [24,53]. Results of bloodmeal analyses conducted in an endemic focus of cutaneous leishmaniasis in Chaute, Lima, showed that L. peruensis is more anthropophilic than is L. verrucarum [54]. It was also shown that multiple-host blood meals are common, including bloodmeals containing blood from animals only present within the houses mixed with blood from animals only present in nearby shelters, indicating movement of flies from animal shelters to the houses (or vice versa) to finish feeding. People living in endemic villages

![Figure 3. Monthly sand fly collection results from three villages in the Cusco Region, Peru. Results show a unimodal annual population distribution pattern with: (A) corresponding mean morning (blue line) and evening (pink line) temperatures and (B) corresponding mean morning (green line) and evening (red line) relative humidity. Collections were made during two nights per month at case homesteads from March 2001 to August 2004. The data gap between October 2001 and January 2002 is due to a cessation of activity mandated by the Peruvian Ministry of Health. doi:10.1371/journal.pntd.0002919.g003](image-url)
report that the sand fly bites occur indoors during the evening, directly before and during the hours of sleep, further confirming that transmission occurs inside the home during the evening and the night [24].

As entomological field studies have shown, vectors of infectious diseases, including vectors of bartonellosis, are sensitive to temperature, humidity, wind, and rainfall patterns; therefore, their abundance is influenced by climate variability [24,26]. During the El Niño weather event that occurred in Peru during late 1997 through mid-1998 weather cycles, the rainy season was extended, and the humidity and minimum monthly temperatures rose. This climatic anomaly allowed potential vector populations to flourish, and unusually high numbers of sand flies were collected [24,26]. The net result was that major epidemics of bartonellosis erupted in Peru, especially in Caraz and Cusco areas. The impact of weather extremes on vector sand fly ecology and transmission of *B. bacilliformis* warrants thorough study [26].

**Host Cell Parasitism, Angiogenesis, and Immune Response**

Colonization of the host: How, when, and where

*Bartonella* species infect an array of mammals with species specificity for their natural reservoir host(s); humans generally serve as incidental hosts. Notable exceptions include *B. bacilliformis* (human reservoir) and *B. quintana* (found in humans and nonhuman primates [55]). Infection of reservoir mammals is generally subclinical with persistent intraerythrocytic bacteremia. *Bartonella* are typically inoculated directly into blood by the bite of an arthropod vector such as fleas, ticks, and sand flies or indirectly when feces of the vector are scratched into the skin. *B. henselae* can also be transmitted via the scratch of a contaminated cat claw. The persistence of bartonellae in the natural host facilitates uptake and transmission by hematophagous arthropod vectors. Colonization of the host’s lymphatics and circulatory system may be enhanced by blood flow and perhaps motility in

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**Figure 4. Results of collection-bottle-rotator (CBR) trap collections of sand flies in Peru.** Results show that: (A) nightly sand fly activity is limited to early evening (1800–2000 hrs) from March through July, the coldest part of the year, which represents the Peruvian winter, and (B) as nighttime temperatures increase in late August through November (late winter and spring), sand fly activity extends throughout the night. “Inside” and “outside” refer to trap locations within and outside a domicile, respectively.

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flagellated species. For example, *B. bacilliformis* are highly motile by lophotrichous flagella (Figures 5 and 6), appendages consisting of approximately 42 kDa flagellin subunits that are resistant to protease treatment [56].

### Adherence to host cells

*Bartonella* infections involve interactions with both nucleated cells and erythrocytes. While erythrocyte adherence by different *bartonellae* is known to be host-specific, adhesion to nucleated cells is considered to be relatively generic. *B. bacilliformis* invades a variety of human cell types, including endothelial cells, dermal fibroblasts, and laryngeal epithelium [57]. Similarly, *B. henselae* and *B. quintana* have been shown to adhere and invade diverse human cell types, including epithelial cells [58], endothelial cells [59,60], and macrophages [61]. In each case, trimeric auto-transporter adhesin proteins (TAAs) are thought to play a leading role in adherence to nucleated host cells [62,63].

*B. bacilliformis* possesses three *brp* genes encoding *Bartonella* repeat proteins (Brps) [64]. These proteins, along with *Bartonella* adhesin A (BadA) or *B. henselae* and variably expressed outer membrane proteins (Vomps) of *B. quintana*, belong to the TAA family of type V secretion systems [62,63]. BadA is the best-understood *Bartonella* adhesin and consists of surface filaments with head, neck, and repetitive stalk domains assembled on a C-terminal anchor [65]. BadA’s head is crucial for host cell interactions and triggering a proangiogenic response [66]. Although *Bartonella* TAA s share remarkable features and functional similarities, they vary considerably in size, and multiple gene variants are often found in the genome [67]. There are at least two variants of the *badA* gene in *B. henselae*, and four genes encode the Vomps of *B. quintana* (*vompA-vompD*) [63]. *B. bacilliformis* Brps share common domains and structural features with TAA s (reviewed in [64]).

The role of BadA has been well studied in *B. henselae*, where it is essential for autoagglutination, adhesion to host cells, and extracellular matrix proteins such as fibronectin and collagens, inhibition of phagocytosis, and induction of a proangiogenic response in host cells [62,68,69]. The Vomps of *B. quintana* have also been shown to play a similar role in host cell interaction and autoagglutination [63,70,71]. Thus, while the biological role of *B. bacilliformis* Brps has not yet been investigated, it is plausible that the Brps play similar roles in pathogenesis.

Erythrocytes are another major target cell during bartonelloses. Optimal adherence to erythrocytes by *B. bacilliformis* occurs in vitro at approximately six hours after co-incubation [34] and is thought to be energy-dependent [72]. Studies have shown that *B. bacilliformis* and *B. henselae* recognize five or six proteins, respectively, of human erythrocyte membranes [73]. A polar tuft of fibrous projections resembling peritrichous flagella on *B. bacilliformis* was observed to make contact with the erythrocyte membrane during adhesion [72], and antiserum to flagella was shown to significantly reduce *B. bacilliformis* association with red cells as compared to controls [56]. These data suggest that flagella may possess adhesive qualities and/or increase bacteria–host cell collisions. It has been proposed that Trw T4SS and flagella expression are mutually exclusive phenotypes (Figure 6), and the Trw system replaces the erythrocytic adhesion role of flagella among non-flagellated *Bartonella* [74]. Specifically, TrwL and TrwJ adhesins have been shown to result in adherence to erythrocytes. TrwJ1 and TrwJ2 were shown to localize to the bacterial cell surface and serve as species-specific adhesins for erythrocytes [75,76].

### Invasion of host cells

*Bartonella* species are facultative intracellular bacteria, and growth within host cells is typical. Virulence studies with *B. bacilliformis* and cultured epithelial or endothelial cell monolayers demonstrated that the bacterium can induce host cells to reconfigure the cytoskeleton and enhance uptake. Internalization
is significantly reduced (approximately 30% of untreated controls) if actin filament formation is inhibited with cytochalasin D, or if bacteria are pre-treated with anti-\textit{B. bacilliformis} antibodies [57]. These inhibition studies indicate that the bacterium plays an active role during internalization and the process involves a surface-exposed molecule(s) that is accessible to antibody.

Parasitism of erythrocytes (hemotrophy) is an uncommon strategy practiced by a handful of bacterial pathogens, including 

**Figure 6. Genomic structure of seven \textit{Bartonella} chromosomes.** (A) Chromosomes (Chr) are arranged in size from largest (\textit{B. tribocorum}) to smallest (\textit{B. bacilliformis}). Plasmid (Pla) sizes are listed, if present. \textit{B. quintana}'s genome encodes the least number of proteins (1,142), and \textit{B. claridgeiae} has the lowest GC% (35.7%). Several virulence-related ORFs have been used to infer phylogeny (\textit{fla – vbh}) and black circles indicate their presence in a particular species. (B) Multiple alignment of seven complete genomes using pM. Location, orientation and position of locally collinear syntenic blocks (LCBs) shared amongst all chromosomes are color-coded and connected by lines. User can analyze location, orientation, and size of LCBs in multiple chromosomes simultaneously (red arrowheads). Local rearrangements, duplications, and inversions are easily identified. Abbreviations correspond to the \textit{Bartonella} species shown in (A).

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Deformin is an extracellular factor that generates trenches and pits in erythrocyte membranes (Figure 7) that resemble those seen on infected red cells [34]. The invaginations are thought to provide entry portals for bacterial colonization. However, even with deformin, bacteria must motile to access the cytosol [78]. Initial studies showed that deformin was a secreted, homodimeric protein (130 kDa in its native state). Activity was enhanced by pretreatment of erythrocytes with trypsin or neuraminidase and abrogated if cells were pretreated with phospholipase D. Deformin-induced invaginations were reversible by treatment with vanadate or dilauroyl phosphatidylcholine (DLPC), or if intracellular Ca\(^+\) levels were increased with ionophores [78,79]. Subsequent work contradicted earlier results and showed that deformin was a small, hydrophobic molecule of approximately 1400 Da with high affinity for albumin [80]. Finally, a third study implicated a set of 36 kDa proteins in the culture supernatant as being required for maximal activity [81]. Despite these discrepant observations, the nature of deformin and its role in hemotrophy warrant further investigation, especially considering the marked morphological changes the factor produces in erythrocyte membranes.

Flagellum-mediated motility also appears to play an important role in hemotrophy. If *B. bacilliformis* is treated with antiflagellin antibodies, in vitro invasiveness for human erythrocytes is nearly abrogated [56]. *B. bacilliformis* also possesses an invasion-associated locus comprised of two genes, *ialA* and *ialB*, that can confer an invasive phenotype on *Escherichia coli* for human red cells in vitro [82]. The *ialA* gene codes for a (di)nucleoside polyphosphate hydrolase that may function to reduce stress-induced dimucleotide “alarmones” encountered in the host cell and therefore enhance pathogen survival [83]. The *ialB* gene codes for a protein with similar molecular mass and considerable amino acid sequence similarity to the adhesion and invasion locus (Ail) protein of *Vibrio cholerae* [84] and the resistance to complement killing (*Rck*) protein of *Salmonella typhimurium* [85]. Both Ail and Rck are implicated in host cell attachment, invasion, and serum resistance. Whether IaA and IaB mediate or facilitate invasion of other host cell types is unknown. A signature-tagged mutagenesis study using *B. bacilliformis* strain displayed a reduced ability to invade erythrocytes in vitro [75]. These results suggest that the *ialAB* locus may be required for optimal intracellular invasion by many *Bartonella* species.

A major difference between *B. bacilliformis* and other common, pathogenic *Bartonella* species, such as *B. henselae* and *B. quintana*, is the absence of VirB T4SS (Figure 6). VirB and the coupling protein VirD4 constitute a machine that delivers T4SS effectors (Beps) to the host cell’s cytosol. In turn, translocated Beps subvert cellular functions, including: (1) rearrangement of the actin cytoskeleton, leading to formation of and internalization by invasions [59,56]; (2) activation of a pro-inflammatory response via NF-κB [82,88]; (3) inhibition of apoptosis [88]; and (4) capillary sprout formation in endothelial cells cultured in a collagen matrix [89]. During invasionosome formation, *B. henselae* are contacted and moved rearward to form an aggregate on the leading lamella of the endothelial cell. The bacterial mass is subsequently internalized by membrane protrusions that are rich in cortical F-actin, ICAM1, and phosphotyrosine. Chemical inhibition studies suggest that invasionosome activity is actin-dependent and microtubule-independent. Not surprisingly, *B. bacilliformis* cannot trigger invasionosome formation because it lacks a VirB T4SS.

**Intracellular growth**

*Bartonella* invade and multiply within cells of their natural hosts. It is thought that seeding of the bloodstream occurs from a primary niche, including the endothelium and migratory cells, such as lymphocytes and phagocytes, or perhaps hematopoietic progenitor cells [74,90]. Regardless, it appears that these bacteria have increased intracellular growth rates [91]. This is puzzling, considering the absolute requirement of *bartonellae* for heme and the iron-limiting nature of the intracellular niche. Two independent systems have been described whereby bartonellae bind and transport heme. The hemin-binding proteins (Hbp’s) are a group of porin-like, beta-barrel proteins that are surface-exposed [92,93]. Hbp’s are encoded by paralogous gene families [94], and the number of genes varies among species (ranges from three in *B. bacilliformis* to eight in *B. australis*). *B. bacilliformis* encodes three closely linked, tandem *hbp* genes (e.g., BAR-BKCK585_1213-1215 and BbINS_05732, 05727, and 05722). Several roles for Hbps have been proposed, including: (1) coating of the bacterium with heme to serve as a nutritive reservoir [93], (2) providing a heme-based antioxidant barrier against reactive oxygen species [93], (3) decreasing oxygen levels around the bacterium to provide a more favorable microaerophilic environment [93,95], and (4) serving as an adhesin to bind fibronectin, heparin, and HUVECs [96].

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**Figure 7.** Scanning electron micrographs of deformin-induced invaginations and pits on erythrocyte membranes. *B. bacilliformis* colonization of cell membrane deformations is readily apparent. Reprinted by permission from [34].

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Bartonella species also employ a more typical hemin utilization (Hut) system consisting of a hemin receptor, an ABC transporter (including a permease, ATPase, and a periplasmic heme-binding protein) and a cytosolic heme-degrading/storage protein [97]. In *B. quintana* and *B. henselae*, the hutB gene is located immediately upstream and in opposite orientation to the hut locus and encodes HutB, a protein that energizes outer membrane processes, presumably including Hut-mediated heme transport. Interestingly, *B. bacilliformis* lacks an obvious *TonB* homologue, begging the question as to the nature of its functional surrogate (note: ExbB and ExbD are found in all three bacteria). Regulation of the hut locus is accomplished through the iron response regulator (Irr) and promoter regions containing an H-box, a motif known to interact with Irr [97]. Other than Irr’s involvement, the heme transport system in *Bartonella* species is remarkably similar to those of other gram-negative bacteria. *B. bacilliformis* is generally accepted as nonhemolytic on blood agar. However, incomplete β-hemolysis can be observed on thin blood agar plates that are cultured for at least four days. The hemolysin is extracellular and can pass through 0.2 μm filters into underlying medium, where it produces a hemolytic zone that silhouettes the colony’s shape [98]. A similar, incomplete β hemolytic activity with a delayed appearance (four days’ growth) was also observed with *B. elizabethae* [99]. Finally, a contact-dependent hemolysin has been described from *B. bacilliformis* that is proteinaceous, requires prolonged incubation periods, acts independently of deformin, and is most active at 37°C [100]. The molecular nature of the hemolysins and their potential role in virulence have not been investigated thoroughly, to date. It is tempting to speculate that hemolysins participate in escape from vacuoles and cytosolic membranes of host cells.

**Angiogenesis**

The clinical manifestation and histological appearance of VP are similar to bacillary angiomatosis (BA), a pathology that can occur in immunocompromised individuals infected with *B. henselae* or *B. quintana* [101–103]. VP and BA are characterized by tumor-like lesions or nodules filled with immature capillaries and swollen endothelium due to chronic colonization by bacteria and infiltration of polymorphonuclear leukocytes and macrophages [104,105]. BA caused by *B. henselae* has been observed primarily on skin, although the bacteria can disseminate and affect other organs such as eyes, liver, and heart [102,106]. In contrast, VP lesions are rarely observed in organs other than skin and subcutaneous tissue [10]. As in tumor angiogenesis, *Bartonella*-triggered neovascularization follows a series of steps that involve disruption of the normal pattern of the extracellular matrix and basal membrane, endothelial cell migration and proliferation at the site of angiogenic stimuli [107].

In an effort to study angiogenesis induced by *B. bacilliformis*, early studies employed nonhuman primate models, including rhesus monkeys [108]. Resulting angiogenic lesions varied depending on the susceptibility of the monkeys as well as the virulence capacity of the bacteria. Intradermal injection of young rhesus monkeys with *B. bacilliformis* obtained from the blood of a patient with a fatal case of OF gave rise to granulomatous nodules filled with new capillaries similar to VP [47]. Nodules varied in size (0.5–2 cm in diameter and 2–10 mm high), appeared at the site of inoculation within three weeks, and increased in size until 60 days post-inoculation, where they remained stationary before starting to regress [47]. The pathogen was readily found inside angioblasts and endothelial cells of nascent capillaries. Noguchi demonstrated that *B. bacilliformis* that had been passaged in rhesus monkeys, as well as those that were taken from VP tissues led to more severe clinical manifestations of infection [109]. In those test animals, numerous miliařy, cherry-red verrucae appeared within one week around the site of inoculation (eyebrows) and were scattered over remote areas such as the groin and legs.

Despite these initial studies, there are few reports describing the molecular basis of *B. bacilliformis*-induced angiogenesis. This is largely due to the lack of an inexpensive in vivo animal model. Although BALB/c mice resist infection by *B. bacilliformis* [110], a SCID/SCID model for studying *Bartonella* infection may prove useful, but has not been tested in this regard [111]. Garcia et al. investigated the effect of *B. bacilliformis* extracts on endothelial cells in vitro as well as in an in vivo model of angiogenesis consisting of subcutaneous implantation of sponge discs containing bacterial extracts into adult Sprague-Dawley rats [103]. The study revealed that a mitogenic factor in bacterial extracts caused vascular proliferation in a dose-dependent manner. However, this factor did not induce a pro-inflammatory response as it did not lead to an increase in the number of phagocytes and lymphocytes. A subsequent report found that *B. bacilliformis* GroEL had mitogenic activity toward endothelial cells undergoing serum starvation-induced apoptosis [112], but it did not affect caspase 3 activity. *B. henselae* was also shown to stimulate endothelial cell proliferation, but to a much lesser extent than *B. bacilliformis* extracts [112]. Subsequently, GroEL of *B. henselae* was found in a secreted fraction of the bacterium and caused endothelial cell proliferation and acted by triggering release of intracellular calcium stores [113]. It is possible that GroEL of both species functions as a chaperonin to stabilize a yet unidentified mitogenic factor. However, in contrast to *B. bacilliformis*, an extract from *B henselae* was shown to have an anti-apoptotic factor that reduced apoptosis of endothelial cells by inhibiting caspase 3 and caspase 8 activities [114]. Interestingly, the anti-apoptotic activity was enhanced in the presence of live bacteria.

Our current understanding of *Bartonella*-induced angiogenesis suggests the process involves at least three mechanisms that work synergistically: (1) mitogenic triggering of endothelial cell proliferation, (2) inhibition of apoptosis (contact-dependent or independent), and (3) angiogenic reprogramming of infected host cells (with pro-inflammatory activation of cytokines) [67,103]. Analysis of VP lesions revealed that infection induces production of VEGF, VEGFR1, VEGFR2, and angiopoietin-2 transcripts [115]. Interestingly, endothelium was shown to produce very little VEGF, whereas overlying epidermis was the major source of VEGF. To our knowledge, the mechanism whereby *B. bacilliformis* induces production of these pro-angiogenic factors has not been investigated, to date. Expression of hypoxia-inducible factor 1 (HIF-1) and subsequent induction of VEGF have been well documented in tumor angiogenesis [116,117].

*B. henselae* infection studies using in vitro cell culture models and BA tissues showed that bacteria induced HIF-1 activation and VEGF secretion [68,69]. This angiogenic response was shown to be dependent on expression of BadA and its interaction with β1-integrin mediated by fibronectin binding [62]. Moreover, *B. henselae* infection was shown to trigger a VirB/Bep-dependent angiogenic phenotype in human endothelial cell models. This includes inhibition of endothelial cell apoptosis and a robust induction of capillary sprout formation mediated by BepA [89,118], and NF-κB-dependent stimulation of a pro-inflammatory response leading to production of IL-8, expression of adhesion molecules such as E-selectin and ICAM-1, and up-regulation of macrophage chemoattractant protein-1 (MCP-1) [105,119]. It is reasonable to speculate that *B. bacilliformis* does not display
effector protein-dependent cell proliferation or anti-apoptotic activity, because it lacks a T4SS. Nevertheless, it may be of interest to determine if *B. bacilliformis* Brps are involved in interaction with β1-integrin and elucidate what role they might play in inducing a pro-angiogenic response.

**Host immune response, protection**

Lymphadenopathy is a frequent complication. Immunosuppression is a common manifestation of *B. bacilliformis* infection and probably results from acute anemia and leukocytosis. In fact, a compromised immune state is thought to contribute to the increased incidence of secondary infections seen in infected individuals.

Lifelong humoral immunity apparently results from a *B. bacilliformis* infection [2,120]. In endemic areas, up to approximately 60% of the population can be seropositive (mainly IgM isotype), and many of these individuals are healthy [121]. Nevertheless, seropositive people who are asymptomatic or post-eruptive for verruga are often blood culture-positive for *B. bacilliformis* [122]. An extraordinary carrier rate, up to 15%, has been reported in some areas of Peru [123], and it is this infected population that presumably serves as the reservoir [1,3,77]. An inverse correlation between patients’ chronological age and incidence or disease severity in endemic regions suggests that humoral immunity, as a result of recurring or chronic infections, confers partial immunological protection [24,36]. Because bartonellae are found in various body fluids and cells, participation by both humoral and cellular arms of the immune system would undoubtedly need to be invoked to resolve the infection. However, for unknown reasons, this does not occur in many afflicted individuals, and the infection is persistent.

**Phylogeny and Genomics: Clues to the Evolution of *B. bacilliformis***

**Bartonella comparative genomics**

*Bartonella* genomes consist of single, circular chromosomes ranging in size from 1.45 Mb in *B. bacilliformis* to 2.64 Mb in *B. tribocorum*, and are largely collinear with a Guanine+Cytosine (GC) content of 38%–42% [124]. Adenine+Thymidine (AT)-richness of these genomes is a universal attribute of host-associated bacteria. Several *Bartonella* species also harbor circular plasmids [125–128], although none have been reported for *B. bacilliformis*. The relatively small size of *Bartonella* genomes is a common attribute among endosymbionts and host-associated bacteria, and possibly reflects genome reduction as a result of decreased selection pressure in the intracellular niche and in the context of a narrower range of hosts [129]. It appears that the number of vertically inherited genes within *Bartonella* genomes is relatively consistent (approximately 1,000), but the number of horizontally acquired (imported) genes varies significantly; thus the genomes of species such as *B. tribocorum* and *B. grahamii*, which both exceed 2 Mb, contain more than 400 imported genes, whereas those of species such as *B. bacilliformis* and *B. quintana*, which are 1.45 and 1.6 Mb respectively, contain 150 or fewer imported genes [127].

Comparative genomics has been used to survey the distribution of coding sequences in extant *Bartonella* species, then correlate this distribution to the phylogenetic juxtapositions of these species as a means to infer patterns of gene acquisition and loss during evolutionary radiation of the genus [124,130]. The focus of considerable interest is on genes encoding virulence factors that mediate interactions with the host, such as the virB and *true* gene clusters and numerous others described elsewhere in this review, as they are likely key determinants of both host specificity and divergent modes of parasitism that characterize the genus (Figure 6).

**Taxonomy/phylogeny of the genus**

The genus *Bartonella* was proposed almost a century ago by Strong to accommodate *B. bacilliformis*, following its description in red blood cells of OF patients by Alberto Barton in 1909 [131]. The advent of 16S rDNA sequencing in the late 1980s suggested that *B. bacilliformis* was genetically similar to the trench fever agent, then *Rochalimaea quintana* [132], and also a newly emerging pathogen that was subsequently named *Rochalimaea henselae* [133,134]. The genera *Bartonella* and *Rochalimaea* were later unified with one another [135] and with the genus *Grahamella* [136], taking the name of the longest established taxon, *Bartonella*. Over the past 13 years, the genus has gradually expanded, such that it now includes 31 “validated” species and subspecies [http://www.bacterio.net/index.html] along with many more informal taxa and partially characterized strains. *B. bacilliformis* is the type species of the genus.

Phylogeny of the genus remains somewhat unresolved. The evolutionary relationship of *Bartonella* to *Brucella* and other genera within the *Agrobacterium/Rhizobium* group of τ-proteobacteria is well established, but delineation of clusters of species within the genus remains somewhat uncertain. The value of intra-genus phylogenetic inference derived from 16S rDNA alignments was limited due to its conservation amongst *Bartonella* species [137,138], thus alignments of other loci have been more frequently exploited. The first of these to be widely adopted was *gltA* [139], and phylogenetic inference derived from this locus indicated that *B. bacilliformis* was the most outlying species, in a lineage deeply divergent from other members of the genus. Phylogenies derived from *ftsZ*, *groEL*, and *tpdB* genes also inferred this peripheral position for *B. bacilliformis* [140]. However, the advent of phylogenies derived from alignment of concatenated sequence data from multiple loci cast doubt on *B. bacilliformis*’ position as an outlier. Reconstructions by Gundl et al. [141] strongly supported a specific evolutionary relationship with *Bartonella claridgeiae*, a position that was reaffirmed by Inoue and colleagues in 2010 [142]. This study also inferred that *B. bacilliformis* and *B. claridgeiae* lay within a well-supported cluster that also contains *B. bartlesii*, *B. bovis*, *B. capreoli*, *B. chomelii*, and *B. schoenbuchensis*.

Whole genome sequence (WGS) data of several *Bartonella* species have now been released, and those of many more are either in draft form or in progress at this writing. Among these, a complete genome is available for one *B. bacilliformis* strain (KC583), whereas data in the form of contigs have been released for a second strain (INS), and projects on nine other strains are currently in progress [http://www.ncbi.nlm.nih.gov/genome/genomes/524]. WGS data are being increasingly used to infer a robust phylogeny amongst *Bartonella* species.

Engel and colleagues [130] generated concatenated sequences from 478 loci within genomes of representatives of eight *Bartonella* species (*B. bacilliformis*, *B. claridgeiae*, *B. grahamii*, *B. henselae*, *B. quintana*, *B. rochalimae*, *B. schoenbuchensis*, *B. tribocorum*, and two unnamed taxa), and from their alignment inferred a phylogeny with four lineages, where *B. bacilliformis* is in a lineage separate from all other *Bartonella*. Most recently, Guy and colleagues [124] described two new, fully resolved genomes (*B. australis* and *B. vinsonii* subspecies *berkhoffii*) and four near-complete genomes (*B. bovis* [2 isolates], *B. vinsonii* subspecies *berkhoffii*, and *B. schoenbuchensis*). The group then compared 428 orthologous core genes amongst the 16 *Bartonella* taxa described.
above. This phylogeny conflicted with that proposed by Engel et al. [130], in that B. australis rather than B. bacilliformis, was the earliest diverging species and that B. bacilliformis clustered with B. bovis and B. schoenbuchensis. Subsequent addition of WGS data for an isolate of an informal taxon referred to as “Bartonella tamarie” to this dataset and a revision of the genes included did not result in a repositioning of B. bacilliformis within the inferred phylogeny, although “B. tamarie” replaced B. australis as the earliest diverging species [124].

Placement of B. bacilliformis within a cluster containing B. bovis and B. schoenbuchensis, rather than the earliest diverging species of the genus, has led to reevaluation of hypothetical evolutionary processes that have shaped Bartonella genomes. The small genome size of B. bacilliformis could be explained by reductive genome evolution following a recent host switch; a model akin to that proposed for B. quintana [143]. As all other members of the cluster containing B. bacilliformis are ruminant-associated, it has been proposed that B. bacilliformis descended from an ancestral strain that successfully made a host shift from ruminants to humans [124]. The alpaca (Vicugna pacos) and the lama (Lama glama), the two most common domesticated ruminant species in the endemic region, are perhaps a source of this ancestral strain. Domestication of these ruminants began some 6,000 years ago, providing a time frame for a hypothetical B. bacilliformis host-shift.

Genetic diversity among B. bacilliformis strains

Few studies exploring the genetic diversity of B. bacilliformis have been published [144,145]. These studies employed a variety of different typing methods, including pan-genomic approaches such as amplified fragment-length polymorphism and infrequent restriction site PCR and/or comparison of nucleotide sequences at loci including gltA, salB, and, most frequently, the 16S-23S rDNA internal transcribed spacer (ITS). All these approaches have delineated genotypes within the species and have been useful in characterizing the molecular epidemiology of bartonellosis [144,145]. Most recently, a multilocus sequence typing (MLST) scheme was developed for B. bacilliformis [146]. Application of this procedure to 46 isolates collected in Peru yielded interesting findings. Perhaps the most significant was that B. bacilliformis may not be a single species; the MLST data provided clear evidence that two isolates recovered from patients with OF had diverged from other B. bacilliformis isolates. These isolates are therefore likely members of a novel Bartonella genospecies. The two divergent isolates were not epidemiologically linked (i.e., they were obtained nine years apart and from locations that lie 150 km from one another). Furthermore, two additional isolates that are also potential members of this new genospecies have been encountered elsewhere [10]. Taken together, results suggest that novel B. bacilliformis-like genospecies have a wide distribution in Peru. The study by Chaloner et al. [146] delineated eight genetic lineages among the isolates surveyed and, in line with many previous studies, these lineages correlated to some degree with the epidemiological provenance of isolates. Thus, strains associated with new foci of Carrión’s disease clustered together and were distinct from the most common genotypes, which were encountered across endemic regions of Peru. However, extrapolation of these data to explore the general epidemiology of Carrión’s disease may be somewhat premature given the relatively small size and opportunistic nature of the collection examined. This shortfall can only be accurately addressed with systematic surveys, although MLST appears to be an appropriate genotyping method to employ if such studies can be instigated.

In an effort to explore more comprehensively the extent of genetic variation among B. bacilliformis strains, we compared the KC503 WGS with genome sequence data available in the form of contigs for the INS isolate (the only genomic data currently available for another B. bacilliformis strain [147]), using progressive Mauve (mM). A total of 20 contigs, containing approximately 1,444,079 base pairs of data, were analyzed. Remarkably, our efforts revealed the two datasets differed by only 27 single-nucleotide polymorphisms (SNPs), with no evidence of any indels or repeat motifs. We mapped the position of these 27 SNPs, finding that 19 lay in putative open reading frames. Of these, 13 were non-synonymous and two introduced a stop codon. Taken together, these observations indicate that the two B. bacilliformis strains, despite sharing no specific epidemiological relationship (being isolated roughly 50 years apart from patients inhabiting geographically remote locations) possess, for all intents and purposes, indistinguishable genomes. Such is the extraordinary nature of these findings that their credibility must be questioned; the upcoming release of genomic data from an additional nine B. bacilliformis isolates will provide a broader insight into intraspecies genomic diversity and put the results described above into better context.

Diagnostics and Identification

Diagnostic symptoms of OF include an extraordinarily low erythrocyte count (approximately 500,000/mm³) [33,77,148] and numerous infected erythrocytes in blood smears prepared with Wright’s, Giemsa, or eosin/thiazine stains [32,121] (Figure 1A). Blood smears have served as a standard diagnostic tool for Carrión’s disease for decades, due to low cost and ease of use. However, the test suffers from low sensitivity, despite its high specificity [4]. Diagnoses can also be made based on blood culture, PCR, immunological tests, or a combination thereof (see below). Diagnosis of VP is made by the occurrence of cutaneous angiomatous skin lesions. As in BA, the histopathology of verrugae reveal bacteria, albeit with difficulty [40], when sectioned and stained with Warthin-Starry silver stain [149] or Giemsa stain [32]. A patient’s age (especially <5 years old) and whether bartonellosis has recently occurred within the family are also useful predictors of a B. bacilliformis infection [24].

Growth requirements and physiology

Bartonellae can be isolated from patient blood by preparing samples with a lysis-centrifugation method [150] or a freeze-thaw protocol [151]. Primary isolates of Bartonella species have also been obtained using a BACTEC blood culture system (Becton Dickinson) and Bartonella alphaproteobacteria growth medium (BAPGM) [152,153]. Homogenized tissues may also be used as an inoculum for blood plates or endothelial cell monolayers [154].

The slow growth rate of B. bacilliformis (approximately 7-hour generation time in vitro [34]) can complicate identification and requires that isolates be cultured several days to visualize colonies. Most bartonellae are microaerophilic, capnophilic (5% CO₂), and grown at approximately 37°C, whereas B. bacilliformis requires cooler temperatures (25°C–30°C) and ambient CO₂ for culture. In addition, humidity of the incubator should be near 100%. Like other Bartonella species, B. bacilliformis cannot synthesize protoporphyrin IX (PPIX) or heme (Fe²⁺-PPIX), and the essential nutrient is obtained from heme (Fe³⁺-PPIX), hemoglobin, and host erythrocytes. Interestingly, Bartonella cannot utilize heme-rich scavenger molecules of the host, such as lactoferrin or transferrin, unlike many pathogens. Commonly used growth media for Bartonella species include chocolate agar, hemin agar,
or various blood agars (for recipes, see [155]). The basal medium of these formulations varies and includes Brucella medium, trypticase soy, GC, heart infusion, or Columbia agars. Media supplements often include erythrocytes (sheep, rabbit, etc.), hemoglobin, IsoVitalex (Becton Dickinson), and serum (sheep). The medium should be pH 7.0–7.5 for optimal growth.

**Colony and cellular morphology, structure**

*B. bacilliformis* is a gram-negative, non–acid fast, pleomorphic coccobacillus. Bacteria stain well with Giemsa or Gimenez stains. Most cells measure 0.5 μm wide by 1.0 μm long. Electron micrographs of *B. bacilliformis* are shown in Figure 5. *B. bacilliformis* colonies are typically very small (initially pinpoint but growing to 0.2–1 mm in diameter after several days), round, and lenticular with an even margin and a raised center in older colonies, and range from translucent to opaque. Although colonies from primary isolates can take weeks to become visible, growth is much more rapid (three to seven days to maximal size) in primary cultures. Most cells measure 0.5 μm long. Electron micrographs of *B. henselae* and *B. rochalimae* [39,56,136]. Twisting motility, as observed in low-passage *B. henselae* and *B. quintana* [58,134], has not been reported. Capsules and endospores are not present in *B. bacilliformis*, in keeping with other *Bartonella* species.

On the whole, the cellular architecture and composition of bartonellae are typical of gram-negative bacteria [157]. Notable exceptions include the unusual fatty acids (CFAs) (see “Cellular Fatty Acid Analysis,” below) and architecture and immunogenicity of its lipopolysaccharide (LPS). A “deep rough” LPS chemotype (little or if any O-chain polysaccharide is present), was first reported for *B. bacilliformis* [158,159]. Later, similar chemotypes were reported for *B. quintana* [160] and *B. henselae* [161]. Further investigation of *B. henselae*’s LPS showed that it contains a unique inner oligosaccharide core of two keto-deoxyoctulosonic acids (KDOs) and one glucose molecule [161]. Its pentaacyl lipid A backbone is also unusual; instead of the typical glucosamine sugars, it contains 2,3-diamino-2,3-dideoxyglucose dissaccharide (GlCN2N) with both sugars phosphorylated. Interestingly, lipid A of *B. henselae*’s LPS possesses a long-chain fatty acid (C26:0 or C20:0) similar to other intracellular pathogens, including *Legionella* and *Chlamydia* species [161]. It is tempting to speculate that *B. henselae*’s LPS is representative of all *Bartonella*, especially considering the uniform LPS chemotype for the genus. Finally, work has shown that *Bartonella* endotoxin is considerably less toxic relative to LPS from other gram-negative bacteria, and its activation of TLR-4 pathways is significantly (1,000–10,000-fold) lower than *Salmonella*’s LPS [161]. In fact, recent work has shown that *B. quintana*’s LPS is a potent antagonist of TLR-4-mediated activation of human monocytes by *E. coli* LPS [162]. Whether a relatively low level of endotoxicity and TLR-4 antagonist activity holds for *B. bacilliformis*’s LPS is unknown, but would not be surprising.

The cellular and subcellular protein content of *Bartonella* species has been analyzed by several groups over the years. Early work showed that the *B. bacilliformis* outer membrane fraction contains 14 proteins of 11.2 to 75.3 kDa in mass [159], as compared to nine surface-exposed proteins in *B. henselae* [163]. The proteomes of whole cells and subcellular fractions of *B. quintana* and *B. henselae* have been analyzed by two-dimensional gel electrophoresis and MALDI-TOF-MS of the protein spots. For example, two studies identified 44 [164] and 53 [165] proteins associated with the *B. henselae* outer membrane fractions. A third study analyzed the *B. quintana* membrane subproteome and found 60 proteins [166]. Finally, analysis of *B. henselae*’s total proteome identified 191 different proteins (431 spots) [167]. Unfortunately, a comprehensive analysis of the *B. bacilliformis* proteome has not been done, to date.

**Biochemical tests**

*Bartonella* are nonfermentative, aerobic (microaerophilic) bacteria with a relatively unremarkable physiology. Thus, conventional biochemical tests are not particularly helpful in presumptive identification of species. One problem with using standard biochemical tests on *Bartonella* species is that hemin is normally not included in the medium, and results must be judged cautiously. Notwithstanding this caveat, standard biochemical tests show that *B. bacilliformis* is strictly aerobic and does not utilize carbohydrates by preformed or de novo enzymes. Tests for oxidase, Voges-Proskauer, catalase, indole production, nitrate reduction, urase activity, hippurate hydrolysis, succinate, alkaline phosphatase, tetrazolium reductase, pyrazinamidase, tributyrin, o-nitrophenyl-β-D-galactoside, esculin hydrolysis, and arginine dihydrolase are all negative [136,156,168,169]. However, tests for peptidase activity on L-proline and L-lysine (acid) are useful in presumptive, differential identification of *B. bacilliformis*, which is negative for proline and positive for lysine utilization [136,156,168,170]. Screens are typically done in multistest formats designed for anaerobes, such as the MicroScan Rapid Anaerobe identification panel (Dade Behring).

**Polymerase chain reaction (PCR)**

PCR is sensitive and specific for identifying *Bartonella* species and can be used to confirm a presumptive identification. PCR can be done directly on affected tissues and/or blood or on enrichment cultures [171,172]. In early work, PCR was used to amplify the 16S rRNA gene or fragments to generate a sequencing template. Resulting data were compared to known 16S rRNA gene sequences [154,173–175]. However, the utility of the 16S RNA gene as a means of differentiating *Bartonella* species was limited due to conservation [137,138]. More useful PCR-based strategies for genus and species identification have been described for serine protease (*hrxA*) [176], glut4 [*139,177], divisome protein (*ftsZ*) [178], riboflavin synthase (*ribC*) [179], NADH dehydrogenase gamma subunit (*nuoG*) [180], heme-binding protein A/Pap31 (*hbpA/pap31*) [181], tmRNA (*ssrA*) [182], RNA polymerase beta subunit (*rpoB*) [183], and the 16S–23S rRNA ITS [138,184]. One recent report employed PCR, with *rpoB* as the target, and pyrosequencing of the resulting amplicons for rapid species identification, including *B. bacilliformis* [185]. Another used *uidA* as a PCR target and showed high sensitivity and specificity in 15 patients with OF [186]. Finally, a recent study demonstrated the utility of collecting dried-blood spots from patients in endemic areas, followed by quantitative PCR (qPCR). Results showed that 24.6% of 65 children studied were positive by qPCR on spots, whereas only 3% were blood-culture positive [187].

**Cellular Fatty Acid (CFA) analysis**

CFA analysis by gas chromatography has been used to identify *Bartonella* to the genus level. This is made possible by the group’s atypical fatty acid composition. For example, predominant fatty acids of bartonellae include cis-11-octadecanoic acid (C18:1 cis-11) (approximately 54%) and hexadecanoic acid (C16:0) (approximately 20%). However, *B. bacilliformis* is unusual in that it contains little octadecanoic acid (C18:0) (approximately 2%) compared to *B. elizabethae*, *B. henselae*, *B. clarridgeiae*, and *B. quintana* (approximately 20%). *B. bacilliformis* also has an unusually high amount (approximately 20%) of cis-11-hexadecanoic acid.
(C_{16:1}) \text{ relative to other } Bartonella \text{ species (typically } \leq 1\%) [99,150,156,188,189].

Immunological tests

Immunological tests are, for the most part, sensitive and specific for Carrion’s disease. However, positive results must be judged cautiously, as they may not indicate an active infection, but rather a persistent humoral immune response. The indirect fluorescence antibody (IFA) test has been used as a specific and sensitive diagnostic tool [190]. In addition, recent work with ELISAs containing recombinant LppB (43-kDa lipoprotein) showed a high degree of sensitivity and specificity in 27 confirmed patients with either chronic or acute disease [191]. Finally, an immunoblot-based protocol incorporating proteins from sonicated B. bacilliformis showed good sensitivity with sera from ten cases of acute disease and very high sensitivity using sera from 32 cases of chronic disease [192]. Likewise, work by Maguia et al. showed a high degree of sensitivity in immunoblots probed with nine and 11 sera from patients with acute and chronic disease, respectively [193,194]. It is clear that immunological tests are useful diagnostic tools for Carrion’s disease, especially when combined with other techniques such as PCR and blood culture. However, immunodiagnoses for this disease are relatively underdeveloped.

Current Status of Treatment, Prevention and Control

Syndrome-dependent courses of therapy

Analysis of antimicrobial susceptibilities of four strains of B. bacilliformis showed that the bacterium is extremely sensitive to a wide array of drugs in vitro, including aminoglycosides, beta-lactams, macrolides, tetracyclines, fluoroquinolones, chloramphenicol, clorimoxazole, and rifampin [195]. However, outcomes of antimicrobial therapy in patients with Carrion’s disease have not been as clear-cut because of several confounding factors, including patient history, immunosuppression, frequent secondary infections, and persistent or recurring bartonellosis despite therapeutic intervention.

Antimicrobial therapy for Carrion’s disease is predicated on whether disease is acute or chronic. Acute disease has been treated with chloramphenicol for decades [2,14,193,196–199], sometimes in combination with other antimicrobials [193]. One apparently effective mixture is chloramphenicol plus a beta-lactam [197]. The choice of chloramphenicol was undoubtedly made popular due to its effectiveness, low cost, and broad spectrum of activity against common co-infectious agents, such as salmonellae [37,193]. Tetracyclines (doxycycline), beta-lactams (ampicillin, penicillin G), macrolides (erythromycin, roxithromycin), trimethoprim-sulfamethoxazole, and fluoroquinolones (norfloxacin, ciprofloxacin) have been used as alternatives to treat OF [15,41,196,197]. Although ciprofloxacin holds great promise for treating acute disease, it should be used judiciously, as resistance mutants arise rapidly in vitro [200,201], and intrinsic resistance to the drug has been reported in strains that were obtained prior to widespread use of fluoroquinolones [202]. Interestingly, in addition to antimicrobial therapy, blood transfusions have been employed for treating severe anemia of OF [36,203].

Chronic disease is usually treated with rifampin, although streptomycin is also effective and was the drug-of-choice prior to 1975 [5,193,197,199]. Of note is that VP does not respond to either chloramphenicol or penicillin treatment [15,32]. Ciprofloxacin, azithromycin, and sulfamicillin, combined with deflazacort, have been used as alternatives to treat VP [4,3,204].

Prevention and control measures

No vaccine is currently available for B. bacilliformis. Early work investigated a composite vaccine prepared from four strains of the bacterium that were grown axenically and formalin-inactivated (0.4% formalin in physiological saline) [205]. Immunizations were administered (one to three times subcutaneously) to 22 men stationed for one month at a Peruvian military outpost located in the “verruga zone” near Mucurana, Peru. Nineteen of the men seroconverted during the study. Twelve men (55% of test group) became blood-culture positive over the course of exposure and eventually presented with symptoms of bartonellosis that were, for the most part, relatively mild. Unvaccinated control groups, prior to and after the study, showed approximately 75% (anecdotal) and 90% infection rates, respectively, with many incapacitated due to severity of illness [205]. Although preliminary in nature, this pilot study suggests that antibodies generated against formalin-killed B. bacilliformis reduce disease severity in humans but do not provide full protection against infection. Reasons for the partial protection are unclear, but may be due to strain-specific antigenic variation, a humoral immune response bias against the vaccine formulation, or modification of important protective antigens by formalin cross-linkage.

In the absence of a vaccine, control of the sand fly vector has been the fallback course of action in endemic areas. As early as World War II, Peruvian army personnel were provided with bed nets and given Vaseline-citronella ointments as a means of warding off sand flies [205]. Early work in endemic areas of Peru showed that DDT insecticide (5% w/v in kerosene) sprayed in and around dwellings was effective at killing sand flies and kept the insects at very low levels for up to four months [206]. The reduction of sand fly numbers “to the vanishing point” using DDT also correlated to a “virtual cessation of bartonellosis” in construction camps within endemic areas of Peru, although exact numbers were not reported [206]. More recent, unpublished results of sand fly control using residual pyrethroid insecticides in homes near Ancash, Peru are promising, as the duration of control was increased to a year (as reported in [4]). Moreover, others have reported good success with residual pyrethroids against old-world sand flies [207]. In any case, control measures using insecticides may prove to be challenging, especially considering that Lutzomyia larvae develop in largely unknown terrestrial habitats, and adult flies, when not resting in houses or animal sheds, dwell in deep cracks and crevices during daylight hours; sites that are not readily accessible to superficial application of insecticides.

In Colombia, other interventions including pyrethroid-impregnated bednets and curtains were used as practical means of controlling sand flies entering houses [208]. In Italy and Burkina Faso, permethrin-impregnated curtains reduced the numbers of endophilic phlebotomine sand flies by 90% and 95%, respectively [209,210]. However, because peak biting activity occurs shortly after sundown and prior to the time when human inhabitants usually retire for the night, impregnated bednets and curtains may have minimal effect in interrupting sand fly–human transmission [211]. Furthermore, it has been shown that sand flies can pass through the mesh of an impregnated bednet and bite a host before succumbing to insecticide exposure (Personal communication, Dr. Edgar Rowton, unpublished data). Nevertheless, if the use of impregnated bednets and curtains can shorten the lives of sand flies, and thereby suppress the vector population, the risk of infection may be reduced [211]. Varying results of field studies dictate the need for trials against specific target vectors.
Future Prospects

B. bacilliformis for before adopting this type of intervention as a means of control for B. bacilliformis vectors.

Field trials conducted in Israel and Kenya showed that sand flies will feed on sugar sprayed on plants in the flies’ natural habitats, suggesting a possible means for controlling sand flies with toxic baits [212]. This finding prompted a pilot study in Kenya to control sand flies resting in rodent burrows and eroded termitaria by spraying vegetation with sugar solution containing Bacillus sphaericus and a dye to reveal that the flies had fed on the sugar [213]. Marked flies were caught entering the rodent burrows and termitaria, and the number of flies emerging from the rodent burrows declined significantly [213]. More recently, scientists at Louisiana State University evaluated several chemical insecticide-treated sugar baits and concluded that they could be efficacious in controlling adult sand fly populations with targeted use in specific habitats [214].

In many foci, sand flies are closely associated with rodents and have been recovered entering and exiting rodent burrows (personal observation, PGL). Burrows of many rodents are the suspected habitats of the immature stages [215]. In fact, sand fly larvae have been observed feeding on rodent feces inside a burrow [216]. Both adult and immature stages benefit from the microhabitat created within rodent burrows [211]. Recently, novel research has focused on sand fly control methods that capitalize on the close ecological interaction between sand flies and rodents. Feed-through or systemic insecticides such as ivermectin or spinosad incorporated into rodent baits have been shown to cause 100% mortality of bloodfeeding sand flies for at least one week [217,218]. In follow-on studies, it was shown that rodents fed on fipronil produced feces that were toxic to coprophagous larvae for up to 21 days. An approach using rodent baits containing fipronil could significantly suppress vector populations that originate in the vicinity of rodent populations, eliminating the portion of the adult population that feeds on the rodents and immature stages that feed on the toxic feces [219].

Future Prospects

In many respects, Carrión’s disease is a “poster-child” for neglected tropical diseases. The illness is endemic to remote, rural areas of South America, and afflicted populations are typically indigent and lack ready access to transportation, modern means of communication, and healthcare. Not surprisingly, few people outside the Andes region are even aware of the disease and the potential public health crisis posed by its spread. Climatic change and vectoring of Carrión’s disease by alternate sand fly vectors is a sand fly–vectored bacterium responsible for an emerging infectious disease of humans that is endemic to South America. B. bacilliformis is the most virulent Bartonella species and causes two disparate syndromes; a life-threatening, acute infection of erythrocytes (OF) and a comparatively benign, chronic infection of the vascular endothelium (VP).

The genome of B. bacilliformis has undergone extensive reduction in response to specialization related to its human-only parasitism and reservoir. Virulence determinants are understudied but include adhesins (e.g., Bartonella repeat proteins), flagella, a deformation factor for erythrocyte membrane (deformin), an invasion-associated locus encoding two proteins (ialAB), a mitogenic protein that may facilitate angiogenesis, and two heme-acquisition systems (Hut and Hbp’s).

Improved diagnostics, better sand fly control measures, enhanced monitoring and surveillance for outbreaks, and a protective vaccine are urgently needed to control Carrión’s disease in South America.

Key Learning Points

- B. bacilliformis is a sand fly–vectored bacterium responsible for an emerging infectious disease of humans that is endemic to South America.
- B. bacilliformis is the most virulent Bartonella species and causes two disparate syndromes; a life-threatening, acute infection of erythrocytes (OF) and a comparatively benign, chronic infection of the vascular endothelium (VP).
- The genome of B. bacilliformis has undergone extensive reduction in response to specialization related to its human-only parasitism and reservoir.
- Virulence determinants are understudied but include adhesins (e.g., Bartonella repeat proteins), flagella, a deformation factor for erythrocyte membrane (deformin), an invasion-associated locus encoding two proteins (ialAB), a mitogenic protein that may facilitate angiogenesis, and two heme-acquisition systems (Hut and Hbp’s).
- Improved diagnostics, better sand fly control measures, enhanced monitoring and surveillance for outbreaks, and a protective vaccine are urgently needed to control Carrión’s disease in South America.

Top Five Papers in Field

1. Foundations: Benson LA, Kar S, McLaughlin G, Ihler GM (1986) Entry of Bartonella bacilliformis into erythrocytes. Infect Immun 54: 347–353.
2. Angiogenesis: Minnick MF, Smitherman LS, Samuels DS (2003) Mitogenic effect of Bartonella bacilliformis on human vascular endothelial cells and involvement of GroEL. Infect Immun 71: 6933–6942.
3. Epidemiology: Chamberlin J, Laughlin LW, Romero S, Solórzano N, Gordon S, et al. (2002) Epidemiology of endemic Bartonella bacilliformis: a prospective cohort study in a Peruvian mountain valley community. J Infect Dis 186: 983–990.
4. Phylogeny: Chalonier GL, Ventosilla P, Birtles RJ (2011) Multi-locus sequence analysis reveals profound genetic diversity among isolates of the human pathogen Bartonella bacilliformis. PLoS Negl Trop Dis 5: e1248.
5. Control: Sanchez Clemente N, Ugarte-Gil CA, Solórzano N, Maguiña C, Pachas P, et al. (2012) Bartonella bacilliformis: a systematic review of the literature to guide the research agenda for elimination. PLoS Negl Trop Dis 6: e1819.
obstacle to overcome to realize this goal is a considerable quantity of capital for research and development.

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