Convergence of Nutritional Symbioses in Obligate Blood Feeders
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Abstract

Symbiosis with intracellular or gut bacteria is essential for the nutrition of animals with an obligate blood feeding habit. Divergent bacterial lineages have independently evolved functional interactions with obligate blood feeders, but all have converged to an analogous biochemical feature: the provisioning of B vitamins. Although symbionts and blood feeders coevolved interdependence for millions of years, we emphasize that their associations are not necessarily stable. Ancestral symbionts can be replaced by recently acquired bacteria with similar biochemical features. This dynamic emerged through combination of phylogenetic and ecological constraints. Specifically, we highlight the lateral transfer of a streamlined biotin (B7 vitamin) operon, and conjecture that its extensive spreading across bacterial lineages may drive the emergence of novel nutritional symbioses with blood feeders.
Symbiosis resolved key challenges in obligatory blood diet

**Blood feeding** (See Glossary) is one of the most specialized diets found in animals [1]. Blood is nutritionally unbalanced with high levels of protein, iron and salt, but few carbohydrates, lipids and vitamins. Blood feeder genomes evolved large repertoires of genes related to vitamin and lipid shortage, haemoglobin digestion, iron managing or osmotic homeostasis to overcome these dietary challenges [1, 2]. Nevertheless, blood feeders cannot synthesize themselves essential cofactors and vitamins lacking in their diet [3, 4]. Facultative blood feeders, such as mosquitoes and fleas, usually need a blood meal to lay eggs but they also rely on other food sources over their life cycle and then avoid nutritional deficiencies.

However, obligate blood feeders (OBF), as ticks, lice, leeches and vampire bats, cannot (Box 1). To overcome this constraint, OBF have converged to analogous functional **microbiomes** with **nutritional symbionts** able to synthesize several **B vitamins**. An obligate blood feeding habit has independently emerged multiple times in animals including insects, arachnids, crustaceans, annelids and mammals, totalizing more than 7,400 species (Box 1). Accumulating studies demonstrate their ancient associations with B vitamin provisioning symbionts to condition the first appearance of OBF lineages that further radiated into current species [5-10]. The OBF microbiomes are functionally distinct from other animals: invertebrate OBF harbor typical low-complexity microbiomes (e.g. [10-14]), each dominated by one B vitamin provisioning symbiont (Box 1), while the only vertebrate OBF, vampire bats, harbor complex gut microbiomes with several potential B vitamin provisioning symbionts but that are distinct to microbiomes of insectivorous, carnivorous and frugivorous bats [15].
**B vitamin provisioning symbionts as essential partners**

Analogous B vitamins-based nutritional interactions appear to be strictly required for the survival and reproduction across the diverse OBF groups. We currently know little about vampire bats, as difficulties to maintain lab colonies make them fastidious experimental models. By contrast, experimental investigations on invertebrate OBF (including ticks, bed bugs, tsetse flies and kissing bugs) showed that, once deprived of their nutritional symbionts, they cease development, stop feeding, molting and reproduction (e.g. [13, 16-18]). They also exhibit physical abnormalities suggestive of a major vitamin deficiency, with dark and inflated bodies. Normal growth and development can be resumed only upon an artificial B vitamins supplementation or symbiont addition. Additional, albeit minor, contributions by nutritional symbionts exist, as exemplified in ticks by the production of the amino-acid L-proline [19].

B vitamin provisioning symbionts have evolved narrow associations with their hosts. They colonize only few organs of OBF, mostly bacteriomes, where they are hosted intracellularly in symbiotic cells termed bacteriocytes, but also gut caecae or Malpighian tubules in some species (Figure 1, Box 2). However, in triatomine bugs, symbionts live extracellularly in the lumen of the gut, although recent investigations also revealed complex microbiomes, including intracellular bacteria that could be additional B vitamin provisioning symbionts. In most cases, B vitamin provisioning symbionts are also heritable through successive generations via high fidelity maternal (usually transovarial, via oocyte infection) transmission (Figure 1, Box 2), and are further maintained during the life cycle of their hosts through transstadial transmission.
The emergence of B vitamin provisioning symbionts

B vitamin provisioning symbionts of OBF originated from at least two bacterial phyla (Proteobacteria and Actinobacteria) and within certain lineages of each phylum, notably the Enterobacteriaceae family (Box 1 and Table 1). We conjecture that their emergence in bacterial phyla depends on a combination of phylogenetic and ecological constraints through three mechanisms:

(i) The first mechanism implies that bacterial ancestors were already adapted to early mutualistic nutritional lifestyles, but with non-OBF organisms. Indeed, many Enterobacteriaceae are facultative symbionts, *ie*, non-required for host survival, but they are well known to influence animal nutrition and metabolism in diverse ways, and many are nutritional symbionts of non-OBF organisms. As such, some are extracellular symbionts inhabiting gut of vertebrates, others are intracellular symbionts of diverse arthropods, and most can produce B vitamins [20]. The combination of their broad distribution in animals and their biosynthesis capacity makes them a breeding ground for evolving nutritional symbiosis with OBF: the emergence of *Wigglesworthia* symbiont in tsetse flies, *Riesia* in lice and *Providencia* in leeches appears to occur through maintenance of ancestral B vitamin genes in these Enterobacteriaceae groups [6, 21, 22].

(ii) The second mechanism implies that bacterial ancestors were not adapted to a mutualistic nutritional lifestyle, but to a parasitic lifestyle with non-OBF organisms. This mechanism relies on maintenance of ancestral bacterial genes that encode for *pre-adaptations* to nutritional symbiosis, but that were primarily dedicated to another function, as best exemplified with the γ-proteobacterium
Francisella associated with ticks. It has emerged from a clade of virulent intracellular pathogens of vertebrates that includes the agent of tularemia F. tularensis [17, 23]. Francisella pathogens have evolved specific mechanisms to penetrate into phagocytes of mammals, and the self-production of biotin (B7 vitamin) is here a key factor that enables pathogen replication and ultimate escape from the phagosomes [24, 25]. In the Francisella genus, the biotin biosynthesis pathway has evolved in the context of pathogenesis before being coopted for nutritional symbiosis in ticks.

(iii) The third mechanism depends on lateral gene transfers. Several B vitamin provisioning symbioses have independently evolved following biotin gene uptakes in the Rickettsiales order (Alpha-proteobacteria): Wolbachia wCle in bed bugs, Midichloria in the castor bean tick Ixodes ricinus and Rickettsia buchneri in the black legged tick I. scapularis (Figure 2). All Rickettsiales are intracellular: some are pathogens, such as the agent of epidemic typhus Rickettsia prowazekii, while others are reproductive parasites of arthropods, such as Wolbachia, but only few harbour B vitamin genes [26-28]. Phylogenomic reconstructions revealed that three independent acquisitions of a streamlined biotin operon are at the origin of the Rickettsiales nutritional symbioses currently found in bed bugs and ticks [26-28]. While the acquisition of special ‘symbiosis’ genes is usually rare for nutritional symbionts [4, 29], the Rickettsiales nutritional symbioses show that foreign gene uptakes are key drivers of interactions with OBF. However, a different mechanism operates for folate (B9): the folate biosynthesis pathway was early present in the Rickettsiales ancestor but secondarily lost in most sub-lineages.
Remarkably, all the folate biosynthesis genes have been consistently maintained in Rickettsiales symbionts of bed bugs and ticks [26-28].

**Convergence of B vitamin provisioning symbioses**

Bacteria adapted to symbiotic (and specifically intracellular) lifestyles underwent massive genome reduction (**Box 3**). The gene set of B vitamin provisioning symbionts is largely a subset of the gene repertoires of their relatives: non-necessary genes have been pseudogenized or are missing completely, but B vitamin synthesis pathways have been conserved [6, 13, 17, 21, 22, 31-33]. However, depending on OBF symbiotic systems, certain B vitamin synthesis pathways have been maintained intact while others have been degraded or lost (**Table 1**).

Notably, the symbiont genomes consistently harbour biosynthesis pathways of biotin and, at lesser extent, folate and riboflavin (B₂): these three pathways form a set of core genes fitting with the nutritional need of OBF. Each of the B vitamins are required for key enzymatic reactions in animals: biotin is a coenzyme for carboxylase enzymes, needed for fatty acids synthesis, branched-chain amino acid catabolism, and gluconeogenesis; folate is a precursor essential for the synthesis of DNA, the modification of DNA and RNA, and is also an important cofactor for cellular metabolism; riboflavin is a precursor of flavin mononucleotide (FMN) and flavin adenine dinucleotide (FAD) coenzymes, which are needed for a variety of flavoprotein enzyme reactions, including activation of other vitamins.

The presence of other B vitamin genes is more variable, and some pathways are missing one or more genes, or are entirely absent (**Table 1**). This pattern may actually depend on the symbiont lifestyle and the specific nutritional need of certain OBF. Indeed, the symbiont *Rhodococcus rhodnii* of kissing bugs has complete gene sets for the eight B vitamins (**Table**
139 1), suggesting that all are potentially needed either for kissing bug life cycle, or for \textit{R. rhodnii} growth, or both. As an extracellular gut symbiont, \textit{R. rhodnii} is exposed to fluctuating environments, in and out of host (it is transmitted by feces; \textbf{Figure 1}). This lifestyle is reflected in its large genome (4.3Mb) which exhibits important gene clusters dedicated to antimicrobial molecules and metabolic plasticity, and the eight B vitamin pathways may contribute to this plasticity [34]. By comparison, most B vitamin provisioning symbionts of OBF are intracellular; they live in a more stable, predictable and protected environment. As such, their metabolic needs are more limited and they have small genomes, lacking genes in almost all functional categories [29, 35, 36]. Hence, most intracellular symbionts have maintained intact the pathways for only few B vitamins other than biotin, ribflavin and folate (Table 1). However, some intracellular symbionts, as \textit{Wigglesworthia} symbiont in tsetse flies and \textit{Riesia} in lice, can produce most B vitamins (Table 1), suggesting that their hosts need this provisioning to their own growth and reproduction.

152 \textbf{The fragile stability of B vitamin provisioning symbioses}

B vitamin provisioning symbionts and OBF evolve a narrow interdependence, and each cannot survive without the other. Thanks to this \textbf{coevolution}, OBF nutritional symbioses have been traditionally envisioned as stable associations lasting for millions of years and resulting in \textbf{co-cladogenesis}, as shown in tsetse flies, and certain clades or genera of bat flies, louse flies, ticks and lice [5, 7-9, 13]. However, recent observations reveal that OBF nutritional symbioses are much more dynamic. Notably, nutritional symbioses can break down: recently acquired symbionts can replace ancestral B vitamin provisioning symbionts and provide similar benefits to the host. Such a pattern was observed in louse flies with the recent
acquisition of a *Sodalis* symbiont [38]. In sucking lice, there are up to six independent lineages of B vitamin provisioning symbionts [6, 7, 39-43], which suggests recent origin of each lineage and therefore several replacement events. In ticks, there are at least four independent lineages of B vitamin provisioning symbionts (*Coxiella, Francisella*, *Rickettsia* and *Midichloria*) [9, 16-17, 19, 23, 26, 28, 31, 33]. Notably, *Coxiella* symbioses are ancestral in some tick genera but recent replacements by *Francisella* in some species appear across the tick phylogeny [5, 9]. Genome sequencing otherwise confirmed that *Coxiella* and *Francisella* have roughly similar B vitamin biosynthesis capabilities: the recently acquired *Francisella* provides the same B vitamin benefit to ticks as the ancestral *Coxiella* (*Table 1*) [17, 19, 23, 31, 33].

Why ancestral and co-evolved nutritional symbionts are replaced in OBF remains unresolved, but several mechanisms can be proposed. Indeed, ancient symbionts suffer Muller’s ratchet, with fixation of deleterious mutations through genetic drift, and they may just have over degraded genomes (*Box 3*). In this context, the comparison of OBF with sap-feeding insects is instructive: they host symbionts compensating for nutritional deficiencies of the sap diet. In sap-feeders, the most severely reduced of symbiont genomes are missing genes usually considered to be essential and harbor the tiniest known bacterial genomes. However, these nutritional symbioses do not collapse thanks to diverse mechanisms recently observed, including evolution of novel traits by hosts to compensate for symbiont gene losses, acquisition of another symbiont to supplement (or replace) functions that are lost in the older symbiont, or DNA uptake from environmental microbes to replace lost symbiont genes. These mechanisms are all potentially applicable to the symbioses with OBF. In ticks, such a pattern was observed for *Coxiella* with loss of essential genes for their replication, offering the
opportunity to another member of the microbiome to out-competing them [31, 33]. In addition, recently acquired symbionts may have higher biosynthetic capability than ancestral symbionts, and then supply additional benefits to OBF. In ticks, some ancestral Coxiella have genomes of only 0.66 Mb [33] while the recently acquired Francisella have bigger genomes (>1.5 Mb) that may have higher biosynthetic capability [17, 23]. This degeneration–replacement model has been proposed for other nutritional symbionts of arthropods [44-46], but are difficult to observe since replacements are expected to be transient [29]. However, the recent observation of a few tick species with co-infections by ancestral Coxiella and recently acquired Francisella may correspond to this transient state before extinction of the ancestral symbiont [5].

**Invasion of a streamlined biotin operon**

Accumulating genomic sequences confirmed that lateral transfer of a compact, streamlined, biotin operon is rampant in OBF nutritional symbioses: related biotin operons (i.e., that diverged recently from the same operon ancestor) were detected in diverse B vitamin provisioning symbionts of OBF (**Figure 2**, [26, 27, 43]). Related operons were also found in other intracellular bacteria, mostly in symbionts of non-OBF arthropods, such as the reproductive parasite Cardinium [47]. The incongruence between bacterial and operon phylogenetic trees underlines that these streamlined biotin operons experience recent (and likely ongoing) transfers between distantly related bacterial lineages (**Figure 2**). Its invasive nature may have contributed to major evolutionary innovations through the emergence of novel OBF nutritional symbioses. What is yet to be established are the mechanisms underlying the invasive nature of this streamlined biotin operon above other biotin operons.
These mechanisms may operate on different levels: the primary acquisition of the operon depends on the opportunity to DNA uptake from other bacteria, while its success to spread in the symbiont population rather depends on selection acting on the benefit it provides.

Lateral gene transfer is usually thought to be rare in intracellular symbionts as they reside in confined and isolated environments. However, according to the ‘intracellular arena’ hypothesis [48, 49], coinfections of different symbionts within the same host cell, and the propensity of some symbionts to switch between arthropod hosts, have created freely recombining intracellular bacterial communities [48-51]. The detection of the streamlined biotin operon in *Wolbachia, Rickettsia* and *Cardinium* [26, 27, 47] that are three of the most common intracellular symbionts of arthropods [52, 53], corroborates the ‘intracellular arena’ hypothesis and its role in emergence in novel OBF nutritional symbioses.

Once acquired, the further maintenance and spread of the operon in the symbiont population may be indicative of positive selection acting on it. A possibility is that the streamlined biotin operon is more efficient in producing biotin than others. In other operon systems, there are selective pressures for efficient specific gene orders and reduced intergenic regions to optimize the expression and functionality of operon in general [54, 55]. Others mechanisms may include the streamlined nature of this operon itself: its compact gene structure may favor its transfer in a single genetic block to other bacteria. Alternatively, the streamlined biotin operon may have genomic features favoring its transposition, but such a mechanism has not been detected to date.

**Concluding remarks**
That feeding specialization to strict blood diet is driven by nutritional symbioses is now beyond doubt. The capacity to synthesize B vitamins is widespread in bacteria but the nutritional symbionts of OBF have all converged to analogous, critical, interactions with their respective hosts. This convergence consists in severe degeneration of bacterial genomes accompanied by preservation of some B vitamin biosynthesis pathways, or in some cases, by a secondary acquisition of the streamlined biotin operon. The OBF nutritional symbioses can be ancient and highly co-evolved. However, we have now to consider that they are also influenced by a dynamic and complex web of interactions by which symbionts move between hosts and genes move between symbionts. We postulate that a better characterization of this web of interactions is now required if we are to understand the mechanisms driving the different aspects of nutritional symbiosis with OBF, such as convergence of B vitamin biosynthesis capacity, instability of association, extinction of ancestral symbiont, acquisition of novel symbionts or invasive spread of the streamlined biotin operon (see Outstanding Questions). Ecological opportunities, along with phylogenetic constraints and selective pressures acting on symbiotic systems, may altogether explain common and divergent evolutionary patterns in OBF nutritional symbiosis. In this context, comparative ecological and genomic approaches will be highly valuable in enhancing understanding of OBF nutritional specialty via symbioses.

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**Glossary**

**Bacteriocyte:** Specialized giant cell (also known as mycetocytes) of certain insects as tsetse flies, bat flies, bed bugs but also aphids or weevils. It specifically contains endosymbionts which provide essential amino acids and other molecules to their hosts, including B vitamins.

**Bacteriome:** Microbes containing cell clusters and surrounding cells which form altogether a symbiotic organ. Depending on species, bacteriomes are found near the gut, or in the gut wall, the gonads or nested among other organs within abdominal body cavity.

**Blood feeding:** The practice, either occasional or exclusive, of certain animals to feed on blood of vertebrates. Also known as blood sucking, hematophagy or sanguinivory.

**B vitamins:** A group of chemically diverse water-soluble vitamins, commonly synthesized by microorganisms, that are essential for cell metabolism and cannot be produced by animals.

The eight types of B vitamins are: thiamine (B₁), riboflavin (B₂), nicotinic acid, also known as
niacin (B₃), pantothenic acid (B₅), pyridoxine (B₆), biotin (B₇), folate (B₉) and cobalamin (B₁₂).

**Co-cladogenesis**: The parallel process of speciation between host and symbiont, such that phylogenetic trees of each partner are congruent.

**Coevolution**: The evolution of two or more species which reciprocally affect each other through the process of natural selection.

**Lateral gene transfer (LGT)**: The movement of genetic material (including genes, operon, genomic islands, or more broad genomic regions) between unrelated organisms that may belong to different genera, families, orders, classes, phyla or even kingdoms, and that usually do not broadly exchange their genetic material. Also known as Horizontal gene transfer (HGT).

**Malpighian tubules**: Excretory and osmoregulatory organs of arthropods that lie in the abdominal body cavity and empty into the junction between midgut and hindgut.

**Microbiome**: The community of commensal, mutualistic and pathogenic microorganisms that reside in an environmental niche (such as an animal for instance).

**Nutritional symbionts**: microbes, usually bacteria and yeasts, that synthesize key nutrients lacking in the hosts’ diet.

**Pre-adaptations**: The possibility of a certain characteristic trait to adopt a new biological function without evolutionary modification, or to have the same function in a different environmental context.

**Reproductive parasite**: Maternally inherited microorganisms that spread within arthropod populations by manipulating the host reproductive processes to enhance their own transmission. Manipulations involve biasing the sex ratio of infected females towards the
production of daughters through parthenogenesis, feminization of genetic males or male
killing. Some reproductive parasites also induce reproductive incompatibility (cytoplasmic
incompatibility) between males infected with a particular strain of bacteria and females not
infected with this strain.

**Transovarial transmission**: Transmission via oocyte infection.

**Transtadial transmission**: Transmission during ontogeny from one life cycle stage to the
next.
Obligate blood feeding (OBF) habit has independently emerged multiple times in animals. There are more than 7,400 OBF species currently known worldwide [60]. Most of the OBF species are found in insects including:

- Sucking lice (Anoplura), with more than 5000 species. The Anoplura are all blood-feeding ectoparasites of mammals. A few species are parasites of humans, including the human body louse *Pediculus humanus humanus*, and the human pubic louse *Pthirus pubis*. Body lice are vectors for the transmission of the human diseases: epidemic typhus, trench fever, and relapsing fever.

- True bugs (Heteroptera), with at least 100 species of cimicids (bed bugs and relatives), 32 species of Polycatenidae bat bugs and more than 130 species of kissing (aka assassin, triatomine or vampire) bugs. Most species of cimicid are specialised on insectivorous bats or birds, while a few species, the common bed bug *Cimex lectularius* and its tropical relative *C. hemipterus*, feed on humans. Kissing bugs share shelter with nesting arboreal vertebrates, from which they suck blood. They are mainly found and widespread in the Americas where they are vectors of the Chagas disease parasite *Trypanosoma cruzi*.

- True flies (Diptera) with tsetse flies (>20 species), bat flies (>500 species) and louse flies (or keds, >150 species). They belong to the superfamily Hippoboscoidea and are obligate parasites of mammals and birds, often with a crab-like (louse fly) or a spider-like (bat fly) appearance. They reproduce through adenotrophic viviparity (see Box 2). Tsetse flies are vectors of trypanosomes, which cause human sleeping sickness and animal trypanosomiasis.

Other OBF species exist in arachnids:
• Ticks (Acari: Ixodidea), with more than 900 species. Depending on species, they feed on mammals, birds, but also reptiles and amphibians. Best-known tick-borne disease is the Lyme disease that is nowadays a widespread infectious disease in the northern hemisphere.

• Other Acari groups, with an undetermined number of species, such as the poultry red mite *Dermanyssus gallinae* (vector of avian spirochaetosis), and the snake mite *Ophionyssus natricis* (vector of ophidian paramyxovirus).

Only few other OBF species exist in non-arthropod groups:

• Leeches (Hirudinea), with more than 700 species. The salivary glands of these annelids produce an anticoagulant peptide, hirudin, used to treat some blood-clotting disorders.

• Fish lice (crustacean subclass Branchiura), with approximately 120 known species. They are the most widespread ectoparasites of freshwater fish in the world and can cause the severe disease state argulosis. All life stages of both sexes are parasitic.

• Vampire bats (Chiroptera: Desmodontinae), with only three species. All are native to the Americas. Although rare, vampire bat bites can infect humans by rabies.

Phylogeny of representative B vitamin provisioning symbionts associated with these obligate blood feeders is depicted in Figure I.

**Figure I. Evolutionary relationships of major examples for B vitamin provisioning symbionts associated with obligate blood feeders.** The phylogeny is based on widely supported findings from studies listed in the citations. B vitamin provisioning symbionts associated with obligate blood feeders are shown in red. The bacterial phylogeny was drawn from iTOL ([https://itol.embl.de/](https://itol.embl.de/)).
Box 2. Localization and transmission of B vitamin provisioning symbionts

Reliable transmission mechanisms of B vitamin provisioning symbionts are necessary to stabilize nutritional associations between generations of OBF, and, more broadly, over evolutionary time. The efficiency of symbiont transmission can be enhanced by the evolution of co-adapted traits that ultimately lead to greater interdependence. Each lineage of OBF exhibits specific traits that condition what mechanisms of symbiont transmission is more favorable. In most OBF harboring bacteriocytes, such as bed bugs and lice, B vitamin provisioning symbionts are transmitted vertically through transfer from the maternal bacteriocyte to the ovaries of the female host, and thence to the eggs, a process known as transovarial transmission (see also Figure 1). In other obligate blood feeders harboring bacteriocytes, such as tsetse flies, bat flies and louse flies, the transmission of B vitamin provisioning symbionts arise latter during the insect development. Females of tsetse flies, bat flies and louse flies retain each egg within her uterus to have the offspring develop internally during the first three larval stages, a method called adenotrophic viviparity. During this time, the female feeds the developing offspring with a milky substance secreted by a modified gland in the uterus. The milky substance contains B vitamin provisioning symbionts which are then transmitted to the developing insect larvae. In triatomine bugs, the B vitamin provisioning symbionts show an exception among invertebrates as they are found in the gut, free of host cells and are transmitted to offspring via the feces either by egg shell contamination or by coprophagy, or even via cannibalism. The excretion of large quantities of water following the quick processing of the enormous volume of ingested blood, combine with the gregarious behavior of the triatomines, may have facilitated the evolution of this transmission mode. In vampire bats, we currently do not know how individuals acquire their
B vitamin provisioning symbionts, but it seems probable that transmission takes place at the time of birth when a newborn is exposed to a mother’s microbiota.
Box 3. Genomic decay in bacterial symbionts

Nutritional symbionts required for host survival harbor unusual genome modifications, including extreme reduction, rapid protein evolution, low GC-content and codon reassignments [4, 29, 36]. The process of genome reduction is initiated as a consequence of loss of selection on multiple gene functions when a free-living bacterium becomes host-associated [61]. The within-host environment, particularly inside a host cell, is relatively stable and nutrient rich, and the need for motility, regulation, secondary metabolite biosynthesis and defense is largely lost: over time the genes encoding such functions are pseudogenized and further definitely lost [4, 29]. For nutritional symbionts required for host survival, this process can continue even further, leading to tiny genomes where even genes considered essential are lost [4, 29, 36]. Symbiont genome decay however affects genes in all functional categories, even those involved in beneficial interactions with hosts [4, 29, 36]. Symbiont genomes continuously accumulate deleterious mutations and their degeneration may ultimately lead to maladaptation and then limit their beneficial contributions to their hosts. A main driving force for this process comes from strict clonality and small population size of nutritional symbionts during transmission, on which Muller’s ratchet effect, leading to elevated rates of fixation of deleterious mutations irreversibly. Genetic drift favors the fixation of neutral or deleterious mutations that cause gene inactivation, gene loss, or inefficiency of gene products. Indeed, gene products of nutritional symbionts have lower efficiencies and reduced thermal stability than their homologs in free-living relatives. Once a bacterium has proceeded down the irreversible path into such obligate symbiosis, there is little opportunity to exit and the nutritional symbiosis can break down and collapse [29, 36].
Figure 1. Major examples for tissue tropism and mode of inheritance B vitamin provisioning symbionts associated with obligate blood feeders. Red dots: B vitamin provisioning symbionts in major hosting organ; Violet dots: B vitamin provisioning symbionts in gonads (for maternal transmission, excluding kissing bugs); Violet organs: gonads/milk glands (lice flies); Yellow organs: midgut/Malpighian tubules (ticks). EB-esophagus bacteriome; MGT- midgut; MGL-milk glands; MT-Malpighian tubules; OA-ovary ampule; OB- ovarian bacteriome; Ov- ovaries; SD- stomach disk; TB- testis bacteriome. Drawings are based on representative work of each taxon: Lice [32, 40]; ticks [17, 56]; Bedbugs [13]; Louse flies [57]; kissing bugs [58]; leeches [22, 59].

Figure 2. Evolutionary relationships, origin and structure of the streamlined biotin operon. The figure is based on widely supported findings from previous studies. Red, B vitamin provisioning symbionts associated with obligate blood feeders; Blue, intracellular symbionts of other arthropods; Black, intracellular pathogens of mammals. Filled arrows and white arrows indicate intact genes and pseudogenes, respectively.
Table 1. Biosynthetic pathways for B vitamins in 11 KEGG curated* genomes of symbionts associated with OBF.

| Phyla          | Order          | Family        | Strain                                | Obligate blood feeder host               | KEGG code | Vitamin B1 | Vitamin B2 | Vitamin B3 | Vitamin B5 | Vitamin B6 | Vitamin B7 | Vitamin B9 | Vitamin B12 |
|----------------|----------------|---------------|---------------------------------------|-----------------------------------------|-----------|------------|------------|------------|------------|------------|------------|------------|-------------|------------|
| γ-proteobacteria | Legionellales  | Coxiellaceae  | Coxiella str. CRt                     | Tick (Rhipicephalus turanicus)          | cey       |             |             |             |             |             |             |             |             |
|                 |                |               | Coxiella str. CeAS-UFV                | Tick (Amblyomma sculptum)               | cend      |             |             |             |             |             |             |             |             |
|                 |                |               | Coxiella str. CLEAA                   | Lone star tick (Amblyomma americanum)  | cee       |             |             |             |             |             |             |             |             |
| γ-proteobacteria | Thiotrichales  | Francisellaceae | Francisella persica                   | Soft tick (Argas arboreus)              | fper      |             |             |             |             |             |             |             |             |
| γ-proteobacteria | Enterobacterales | Erwiniaceae  | Wigglesworthia glossinidina           | Tsetse fly (Glossina brevipalpis)       | wbr       |             |             |             |             |             |             |             |             |
|                 |                |               | Wigglesworthia glossinidina           | Tsetse fly (Glossina morsitans)         | wgl       |             |             |             |             |             |             |             |             |
| γ-proteobacteria | Enterobacterales | Morganellaceae | Arsenophonus lipopteni                | Deer keds (Lipoptena fortisetosa)        | asy       |             |             |             |             |             |             |             |             |
| γ-proteobacteria | Enterobacterales | Enterobacteriaceae | Candid. Riesia sp. GBBU | Gorilla louse (Pthirus gorillae) | rig       |             |             |             |             |             |             |             |             |
|                 |                |               | Candid. Riesia pediculicola str. USDA | Human body louse (Pediculus humanus humanus) | nmp        |             |             |             |             |             |             |             |             |
| α-proteobacteria | Rickettsiales  | Midichloriaceae | Candid. Midichloria mitochondrii      | Castor bean tick (Isodes ricinus)       | nnm       |             |             |             |             |             |             |             |             |
| α-proteobacteria | Rickettsiales  | Anaplasmataceae | Wolbachia str. wCle                   | Bed bug (Cimex lectularis)              | wcl       |             |             |             |             |             |             |             |             |

*KEGG pathways used for B1, B2, B3, B5, B6, B7, B9 and B12 are 00730, 00740, 00760, 00770, 00750, 00780, 00790, 00860, respectively. Black squares, putatively functional pathways; Grey squares, incomplete pathways with pseudogenes or missing genes; white squares, pathways absent.

** Only flavin reductase (fr) present. An enzyme that converts Aquacob(III)alamin into vitamin B12.