L-form bacteria, cell walls and the origins of life

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1. Summary

The peptidoglycan wall is a defining feature of bacterial cells and was probably already present in their last common ancestor. L-forms are bacterial variants that lack a cell wall and divide by a variety of processes involving membrane blebbing, tubulation, vesiculation and fission. Their unusual mode of proliferation provides a model for primitive cells and is reminiscent of recently developed in vitro vesicle reproduction processes. Invention of the cell wall may have underpinned the explosion of bacterial life on the Earth. Later innovations in cell envelope structure, particularly the emergence of the outer membrane of Gram-negative bacteria, possibly in an early endospore former, seem to have spurred further major evolutionary radiations. Comparative studies of bacterial cell envelope structure may help to resolve the early key steps in evolutionary development of the bacterial domain of life.

2. Uncertain evolutionary origins of the bacterial sub-kingdom

Bacteria occupy virtually every conceivable ecological niche on the planet and proliferate in vast numbers. The geological record and estimations from evolutionary clocks suggest that bacteria first appeared over 3 billion years ago [1]. Huge catalogues of rDNA sequences, supplemented by a rapidly increasing number of complete genome sequences, have provided crucial insights into the phylogenetic space occupied by bacteria, and led to the identification of about 20 major phyletic groups [2]. The new taxa expand on earlier groupings based largely on morphological and physiological traits. Figure 1a shows a typical tree [3], with most of the major recognized groups indicated. No attempt has been made here to put a scale on the length of the branches and the exact order of branching should be viewed as tentative. This is because at least two important factors confound attempts to define the root of the tree and the early order of branching. First, the majority of bacterial genes seem to have been subject to horizontal gene transfer and so generate inconsistent tree structures. Second, at the deepest phylogenetic levels the extent to which sequences are conserved disappears beneath the threshold at which statistical methods give reliable outcomes. Thus, how the major groups of bacteria emerged from the pre-cellular ‘primordial soup’ and began to differentiate from each other remains obscure [6–8].

3. The cell wall as a unifying trait among bacteria

Bacteria share a number of traits that distinguish them from the other major group of prokaryotes, the archaea and from eukaryotes. Most prominently, these include ‘information storage and processing’ functions: especially the DNA replication, transcription and translation machineries [9,10]. Another major feature thought to be unique to bacteria is the peptidoglycan (PG) cell
Figure 1. Outline phylogenetic tree of the bacteria and the distribution of key cell envelope features. (a) An outline phylogenetic tree for the bacteria, based on the major phylogenetic groups recognized by Wu et al. [3]. The detailed branching order is tentative for reasons outlined in the text. Groups traditionally described as Gram-negative and Gram-positive are indicated in red and blue text, respectively. (b,c) Distribution of genes required for PG (b) or Lipid A (c) synthesis, based on the output from STRING v. 9.0 [4] using murC and lpxC as markers. Similar results were obtained with other genes from the PG or the OM pathways (not shown). The number of organisms in each group are shown. Plus symbols (+) denote greater than 90% of organisms contained a likely gene homologue; plus or minus symbol (±), 50–70%; minus symbol (−), less than 5%. (d) Comments on current knowledge of cell envelope structures of various groups based mainly on information from Bergey's Manual of Systematic Bacteriology [5].

wall. This important structure provides an external protective shell that serves several critical roles for those organisms that possess it. Traditionally, bacteria have been divided into two major subdivisions depending on whether they possess a second membrane (the outer membrane, OM) outside the PG wall. Gram-negatives or ‘diderms’ (red in figure 1a) have an OM, whereas Gram-positives (blue) lack this layer, often having a particularly thick PG layer instead. As summarized in figure 1ad, there are a number of variations on this simple scheme, and envelope structure does not align completely with sequence-based taxonomies. Nevertheless, the cell wall is a crucial structure in almost all bacteria. Their growth and division are limited by the necessity to enlarge and then divide the wall. These processes are usually regulated spatially and temporally by important cytoskeletal proteins, MreB and FtsZ, that are distant relatives of actin and tubulin, respectively [11]. The biosynthetic pathways for PG precursor synthesis are well worked out and 10 or so genes encoding the key enzymes needed to make a functional wall can readily be identified. The levels of conservation of these various proteins and their respective genes are variable. However, several of the proteins appear to have excellent sequence conservation right across the bacterial sub-kingdom. Figure 1b shows an example of the conservation of a PG precursor synthesis gene, murC. Although a few groups of bacteria do not have a cell wall, in the well-characterized examples (e.g. Tenericutes, including Mycoplasma, Phyttoplasma, etc.), it is clear that these organisms are closely related to major groups of bacteria that all possess a wall [12,13]. This strongly suggests that the exceptional organisms emerged by loss of the wall at some point in evolution. On the basis of the near ubiquity and conserved synthesis and structure of the PG cell wall, it seems reasonable to assume that the last common ancestor (LCA) of all bacteria (if such a single ancestor ever existed) already had a wall. What, then, was the nature of cellular life before the invention of the wall? Recent discoveries with ‘L-form’ (cell wall-deficient) bacteria may provide clues as to the general cellular organization of the primitive progenitors of bacteria and especially how they proliferated.


Cell wall-deficient variants of bacteria that normally possess a wall have been described many times in the literature, since the original report of Klieneberger [14] nearly 80 years ago. We recently embarked on a molecular genetic analysis of the L-form state and showed that conversion of Bacillus subtilis into a form that can replicate reasonably efficiently in the absence of a cell wall (i.e. as an L-form) requires only two genetic changes [15] (R. Mercier, Y. Kawai & J. Errington 2012, unpublished data). Remarkably, despite the limited mutational changes required, the new L-form cells completely abandon the normally essential cell division machinery used by virtually all extant bacterial cells, and proliferate instead by a mechanism of membrane tubulation or blebbing (figure 2a). We showed, at least for B. subtilis, that this process is completely independent of the cell wall precursor synthetic pathway and the major cytoskeletal proteins, MreB and FtsZ. A recent report on Listeria L-forms highlighted a somewhat different process but which nevertheless again
involves membrane dynamics (‘vesiculation’ in figure 2a) [17].
A wide range of bacteria are thought to be readily able to enter the L-form state, including both Gram-positive and -negative lineages [18]. Where their proliferation has been described, it generally resembles the mechanisms illustrated in figure 2. We have suggested that the membrane blebbing process used by *B. subtilis* L-forms represents an early mode of proliferation used by primitive cells before the invention of the cell wall [15,16]. The mechanism may have been retained by modern cells as a back-up process for use when cell wall synthesis is compromised. The risk to cell wall damage is probably ancient, given the widespread production of cell wall active antibiotics, such as β-lactams, glycopeptides and lipopeptides, by various ancient groups of bacteria [20,21].

5. Importance of membrane composition and dynamics for L-form proliferation

The finding that L-form proliferation is independent of cytoskeletal systems highlighted the question of its mechanism. Having established that blebbing and tubulation of L-forms was not dependent on known cytoskeletal systems we attempted to identify genes required specifically for this curious form of proliferation. The clearest mutant phenotype obtained, in which L-form proliferation was virtually abolished but growth of walled cells was unaffected, turned out to work by impairing branched chain fatty acid synthesis. Various lines of evidence point to this phenotype being manifested by a requirement for high membrane fluidity. Mutant cells with reduced membrane fluidity were able to grow in size and undergo some shape perturbations but they did not undertake the final membrane scission step needed to generate separate progeny cells [16]. The requirement for a particular state of membrane fluidity highlights the importance of the biophysical properties of membranes in L-form proliferation.

6. L-form-like proliferation under simple conditions in vitro

Meanwhile, as thoroughly reviewed recently by Briers *et al.* [22], in vitro experiments designed to reconstruct key steps required for the evolution of early forms of cellular life have generated outcomes remarkably reminiscent of the proliferation of L-form cells. One of the most important theoretical requirements for the evolution of life is a mechanism for encapsulating nucleic acids and the products of replication and gene expression [23]. A related requirement is for the cell envelope to replicate, while retaining and segregating the cell contents. Several recent studies have created plausible solutions to this problem by providing controlled lipid vesicle growth and fission under relatively simple *in vitro* conditions [19,24–28]. The theoretical basis for membrane vesicle fission is also well established [29,30]. In outline, shape perturbations leading to fission can be generated simply by increasing the vesicle surface area to volume ratio. A range of vesicles with different configuration and composition have been studied: unilamellar or multiple layered, and made up of simple fatty acids, as were probably abundant in the prebiotic era [31], through to more physiologically relevant phospholipid mixtures. Surface growth can be driven by ‘feeding’ with amphipathic molecules, such as fatty acids, either in solution or as micelles, which can intercalate or fuse with the vesicles to increase the surface area of the outermost leaflet or bilayer. Provided that the internal volume of the vesicle does not equilibrate rapidly, the mismatch between surface area and volume directly drives shape distortions that can lead to vesicle fission [19,25,28]. Similar outcomes can be generated by use of osmotic upshift to reduce interior volume at fixed surface area [24]. An example of the remarkable similarity in appearance between *in vitro* derived vesicle fission processes and L-form division is shown in figure 2b–d. Shape distortions and fission can also be promoted by intravesicular
nanoparticles or macromolecules [26,27], suggesting a possible role for cell constituents, particularly the nucleoid, in promoting fission. Folding of the membrane around a nucleoid would directly promote the formation of a genetically viable progeny cell. The development of increasingly complex in vitro proliferation systems, more closely mimicking living cells, in parallel with further top-down dissection of L-form proliferation, appears an exciting area for future studies.

On the basis of the recent in vitro experiments and their similarity to L-form morphology and behaviour, it seems entirely plausible that membrane dynamics and purely biophysical processes could have allowed early cells to proliferate. It then follows that L-form proliferation could be a relic of this primitive mechanism originally used before the invention of the cell wall, but which remains accessible to modern cells.

7. L-form-like growth and proliferation would have supported massive genetic flux or horizontal gene transfer

Essentially, all modern organisms share a common genetic code, showing that some basic features of biological life became fixed relatively early. However, as discussed earlier, it is well known that archaea and eukaryotes differ significantly from the bacteria, especially in terms of the machinery used to replicate and express the information in DNA [9,10]. These differences are consistent with the notion that the mechanisms underlying various key cell functions were still in a state of flux when the early ancestors of the archaea and bacteria separated from each other.

We have observed L-forms to undergo spontaneous fusion, potentially generating heterokaryons or chimaeric genomes, under certain conditions (P. Domínguez-Cuevas & J. Errington 2012, unpublished data). Moreover, protoplasts (which are cells that are transiently converted into an L-form-like state but which do not normally undergo prolonged growth or division) have long been used by geneticists in fusion experiments to cross different bacterial strains, allowing the selection of recombinant genomes [32]. Assuming that ancient cellular organisms had a similar vesicular structure and used an L-form-like mode of proliferation, it seems likely that their primitive genomes would have been susceptible to continuous genetic flux. Fusion of vesicles containing dissimilar genomes would have allowed them to recombine or re-assort, and blebbing or tubulation would allow propagation and spread of the recombinant progeny. L-forms tend to have an unusually wide range of sizes and frequently contain multiple genomes [15,17]. The coincidence of multiple genomes in a single membrane bound vesicle would have facilitated genetic exchange.

8. Invention of the peptidoglycan wall and the bacterial radiation

It is generally assumed that a key step in the evolution of life must have involved a transition from an early acellular form of life in which horizontal gene transfer was rife, to more recognizably modern cellular organisms that could undergo tree-like evolutionary progression, leading to the elaboration of increasingly sophisticated forms. Woese [33] called this transition the ‘Darwinian Threshold’. However, there is presently no consensus view as to the possible basis for the emergence of stable cellular life forms. In line with a previous suggestion [34], I suggest that invention of the PG cell wall could have been a pivotal step in the evolution of bacteria.

Emergence of a wall would have conferred several important advantages to the genome that encoded it. First, the wall provides an external rigid or semi rigid ‘shell’ that offers protection from all kinds of damage, as well as the ability to explore and tolerate a much wider range of habitats, particularly of differing osmolarities, than their simple membrane-bound predecessors. Second, the wall would provide much tighter control of cell size, shape, division, growth orientation, etc., especially in conjunction with a cytoskeleton to provide spatial regulation. Certainly, walled B. subtilis cells grow much faster than their isogenic L-form counterparts [15]. Third, the wall would almost certainly have had a dramatic effect on genome stability, protecting cells from the rampant horizontal gene transfer that probably occurred in cells with a naked cytoplasmic membrane. This would in turn have allowed a much more sophisticated integration of the genetic operating system (replication, transcription and translation) with the informational genes specifying the day-to-day life of the organism and its interactions with the environment.

The advantages offered by invention of the wall could have enabled an explosion of successful new forms all sharing the PG wall system but featuring increasingly elaborate variations able to explore and exploit new environments (figure 3).
9. Emergence of archaea by invention of a different cell envelope structure(s)?

It is well known that archaea tend to have different cell envelope structures to bacteria [35]. Informatic analysis of the large number of archaeal genomes now available confirms that they completely lack the multi-step biosynthetic pathway for PG precursors (J. Errington 2012, unpublished data). Unlike bacteria, archaea display a variety of different protective surface layers, most commonly, a paracrystalline proteinaceous shell called an S-layer [35]. Several families of archaea have a PG-like layer called pseudomurein, but the precursors for assembly of this kind of wall are synthesized by a completely different enzymatic pathway from that of PG. In the light of the above discussion, it seems possible that the archaea emerged from the ‘protocell soup’ by invention of one or more different cell envelope structures. The genome captured by the successful enveloped archaeal progenitors had a related but distinct operating system, explaining the differences in replication, transcription and translational apparatus between bacteria and archaea.

10. Evolutionary development of the bacterial cell envelope

The idea that bacteria underwent a major radiation following the invention of the wall predicts that it should be possible to discern a pathway of emergence of the major modern bacterial groups, in parallel with an elaboration of wall structures. Unfortunately, sequence-based homology detection methods have not yet provided a robust and reliable means of discerning branching order at the deepest levels of bacterial evolution [2]. However, Jensen and co-workers [36] recently provided a major new insight into this problem by discovery of a likely origin, perhaps for all of the Gram-negative bacteria, within the Veillonellaceae group of the Firmicutes (essentially the major group of low G+C Gram-positive bacteria). The relationship between the Gram-negative and Gram-positive bacteria has been the matter of much speculation and discussion. Some have viewed the diders as likely to have evolved from the monoders by evolution of a more complex cell envelope [37]. Gupta suggested that antibiotic selection provided the driving force for development of this additional protective layer. By contrast, Lake [38] suggested that the OM arose following an ancient endosymbiotic fusion of actinobacterial and clostridial forms. A diametrically opposing view is that the diderm form was ancestral and that monoders emerged by loss of the OM [39]. Schemes such as the one shown in figure 1a do not help in distinguishing between the two models because the Gram-positive and -negative groups appear interspersed. But as explained earlier, this could be due to lack of statistical resolution in the deeper branches of the tree.

The main problem with the first of the above scenarios was the question of how the OM could have been invented. A solution to this problem, strongly supporting the likelihood of a monoderm → diderm order of emergence, was provided by recognition that even though the Veillonellaceae clearly belong to the ancient Firmicutes lineage, they are Gram-negative in organization. Furthermore, their OM originates in the OM of the endospore, which is a specialized cell-type characteristic of the Firmicutes. Figure 4 shows the well-characterized steps of endospore formation, as largely worked out through decades of research on B. subtilis and its relatives. Engulfment of the prespore by the mother cell leads to a ‘cell within a cell’ and in particular, generates a prespore with a double membrane. In B. subtilis and many other spore formers, the OM disappears during spore development and is not visible when the spore eventually germinates. However, in the Veillonellaceae or at least some members of this group
The major Gram-negative phyla are now suggested to have emerged from within the ancestors of the Tenericutes group by complete loss of the wall. The evolutionary relationships between modern Firmicutes, Actinobacteria, Chloroflexi and the Deinococcus-Thermus group remain unclear. The major Gram-negative phyla are now suggested to have emerged from within the ancestors of the Firmicutes by retention of the endospore OM. The Tenericutes probably emerged from the same ancient group by complete loss of the wall.

 especialmente, Acetotena longum [36]), the OM remains and is retained through to spore outgrowth and beyond, as normal vegetative growth is restored. Detailed informatic searches support the idea that the Acetotena group (Veillonellaceae) are truly core members of the phylum Firmicutes. If this idea is correct it has profound implications for the evolutionary development of bacteria because the whole diderm radi-ation—including many or all of the red ‘Gram-negative’ bacteria listed in figure 1a would be descendants of a branch in the Firmicutes lineage (figure 5). The root of the whole bacterial lineage would then lie somewhere in the ancestors of a relatively small group of monoderm bacteria.

Molecular markers specific for OM biogenesis are limited because many of the proteins, e.g. Omps, show poor sequence conservation. The lpxC gene presently seems to be the best surrogate [36]. It is a well-conserved cytoplasmic protein carrying out a key specific step in the synthesis of LipidA, the major lipid component of the OM of many diderm bacteria. As shown in figure 1c, lpxC tracks well with the diderm state. Included are several major subgroups of organisms of previously uncertain origin: Cytophagobacteria, Chlorobi, Bacteroides, Planctomycetes and Chlamydia. We cannot rule out the alternative hypothesis, that these organisms derived lpxC and multiple genes needed to make an OM by lateral gene transfer, but this seems unlikely.

The explosion of genome sequencing is providing a gold mine of information on the diversity and versatility of the bacteria. Nevertheless, there are still major gaps in our knowledge of how the bacterial (and archaeal) groups emerged and then diverged. Sequence-based analytical methods are still inadequate for defining the early steps in the bacterial radiation. Several groups of bacteria that may be representative of pivotal steps in the evolutionary sequence are poorly characterized. Comparative studies of the structure and synthesis of the cell envelopes of a wider range of organisms may provide important clues about the evolutionary development of the bacteria. Thermotoga provides an interesting example. Its defining feature is a loose outer layer called a toga. This was thought to be related in some way to the OM of diderm bacteria. However, as shown in figure 1a, these organisms lack an lpxC gene and therefore can probably not synthesize LipidA. Nevertheless, the toga was recently shown to contain classical Omp-like proteins [40], consistent with these organisms having a diderm ancestry. It seems likely that further detailed studies of the cell envelopes of deeply rooted bacterial groups will provide important insights into the early evolutionary history of the bacteria.

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References


