How Cnidaria Got Its Cnidocysts

Stanley Shostak*

Department of Biological Sciences, University of Pittsburgh, USA

*Corresponding author: Stanley Shostak, Department of Biological Sciences, University of Pittsburgh, USA, Tel: 0114129156595, E-mail: sshostak@pitt.edu

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Abstract

A complex hypothesis is offered for the origins of cnidarian cnidocysts through symbiogeny. The two-part hypothetical pathway links the origins of tissues through an early amalgamation of amoebic and epithelial cells to and the later introduction of an extrusion apparatus from bacterial parasites. The first part of the hypothesis is based on evidence for morphological, molecular, and developmental similarities of cnidarians and myxozoans indicative of common ancestry. Support is drawn from Ediacaran fossils suggesting that stem-metazoans consisted of symbiogenic pairs of epithelial-like shells enclosing amoeba-like cells. The amoeba-like cells would have evolved into germ cells and cells differentiating as or inducing nerve, muscle, and gland cells. The second part of the hypothesis proposes that cnidocysts evolved in a cnidian/myxozoan branch of the metazoan tree through the horizontal transfer of bacterial genes encoding an extrusion apparatus to proto-cnidarian amoebic cells and consequently to the Cnidarian germ line. Evidence for bacterial genes in Cnidaria and transposable elements are cited in support of this part of the hypothesis.

Keywords: Cnidaria; Cnidocysts; Epithelial cells

Introduction

Ever since the modern synthesis, efforts to sort out the source(s) of unique phylogenetic characteristics start with the assumption of de novo novelty: “mutation did it.” Symbiosis offers an alternative source of cladogenic innovation albeit requiring compounding hypotheses with horizontal gene transfer. Setting aside this complication, the origins of cnidocysts in Cnidaria offers a unique opportunity for examining a hypothetical role for symbiosis in metazoan evolution.

Nowadays, macro- and micro-cnidarians are identified by the presence of cnidocysts (also known as nematocysts in Cnidaria and polar capsules in Myxozoa). Otherwise, cnidarians per se would seem to have little in common. Macro-cnidarians are classical anthozoans (anemones and corals) and medusozoans (medusas [jellyfish] and polyps), while micro-cnidarians are myxozoans, comprising myxosporeans (including actinosporeans) and malacosporean parasites. The question is, “How did cnidarians of either size acquire similar cnidocysts?”

Richard Christen et al. [1] offered an answer: “the unorthodox possibility that... metazoans (and possibly plants) were the results not of the aggregation of a single species of unicellular organisms, but the results of various symbiotic events between different types of protistan organisms.” In the early 1990s, several eukaryotic protists had already been nominated as candidates for sources of cnidocyst, but the notion of “symbiotic events” was only beginning to be taken seriously as a driving force of evolution [2,3].

Morphological similarities had long been recognized between cnidian cnidocysts, and microbial “cnidocysts” [4,5], specifically the “peduncle,” “rhizoid,” and “perforator” of dinoflagellates [6-8], the trichocysts of trypanosomes [9], zooflagellates [10] and mastigophorans [11], the “apicoplasts” (apical complexes) of “Sporoza,” the “polaroplast,” of microsporidian [12], and the “polar capsules” of myxosporidians [13-19]. Taking these similarities into account, Jiří Lom noted “homologies [were] perhaps too close to be considered only a convergency phenomenon” [20], and Pierre Tardent commented “The wheel didn't have to reinvent itself” [21].

Thus, I suggested that “any of several or even more than one 'sporoza' or predatory dinoflagellate... [might have served as] the source of cnidarian cnidocysts” [22], and Leo Buss and Adolf Seilacher famously “hypothesize[d] the existence of a cnidian ancestor that lacked cnidae” and was subsequently parasitized by a microsporidian bearing an extrusion apparatus. This “parasitism ultimately led to the integration of genes required for polar capsule syntheses into the cnidarian ancestor's genome” [23]. Likewise, Jason Holland et al. alluded to the possibility that “nematocysts did not originate in cnidarians... [but] could be explained by lateral organelle or gene transfer from protists that possess extrusable organelles similar to nematocysts [16]. These hypotheses, whether invoking sporozoans, microsporidians, or simply protists as the source of cnidocytes had fatal weaknesses, however, and alternatives were necessary to overcome these weaknesses.

Fatal Weaknesses

Buss and Seilacher's [23] choice of microsporidians as the parasitic source of cnidocysts was perfectly reasonable especially, since at the time, microsporidians and myxosporeans were considered closely related: They both lacked flagella, and had an extrusible organ, the polaroplast or polar capsule, respectively. The conspicuous cnidian “penetrant” or stenotele resembled the microsporidian polaroplast's “spore extrusion apparatus (EXA)... equipped to explosively discharge a tube through which the infective sporoplasm is sent into a host cell” [24], while the sticky or ensnaring tube of most cnidarian cnidocytes more nearly resembled the thread averted by myxosporean's polar capsules.

Differences in the cellularity of microsporidians and of cnidarians suggested that the similarities between polaroplasts and cnidocysts
were due to convergence, however. Eventually, “profound differences in biology and ultrastructure were demonstrated” [25], and Microsporidia fell out of the running for ancestor of cnidarian cnidocysts.

Myxosporidians were also eliminated as the sources of cnidarian cnidocysts, but their fatal flaw was not the failure of homologies. Rather, their problem was one of timing: Which came first?

Historically, metazoan characteristics were recognized in cells or plasmodia formed during myxozoans’ parasitic life cycles, for example, the metazoan-like separation of somatic cells (with diploid nuclei, tight junctions, and collagen) from germ cells (with haploid nuclei and intercellular bridges). But the place of myxozoans in the Metazoa was shrouded in ambiguity. For example, Polypodium hydridorme, the infamous parasite of caviar bore Cnidar-like cnidocyst, while, Buddenbrockia plumatellae, the parasitic “worm” of freshwater bryozoans was thought to be a myxozoan. Ultimately, arguments on behalf of Polypodium’s place among the cnidarians (Narcomedusae [18]) were strengthened with molecular evidence [19] and Polypodium was established as a parasitic cnidarian that lives as an inverted stolon (instead of a stalked plasmodium). The resulting polyps, equipped with cnidocysts, form gonads containing (presumably) infectious germ cells (analogous to myxozoan spores. Likewise, molecular affinities were discovered between the “cnidarian” Buddenbrockia plumatellae and the malacosporean Tetracapsuloides byzooideae [14,19,26-28]. Thus, the knot tying Myxozoa to Cnidaria continued to tighten.

Microscopic evidence also supported the homology of myxozoan polar capsules and cnidarian cnidocysts. Cnidocysts had long been recognized as intra-cellular “organoïds” [29] or encapsulated cellular organelles. Ultrastructurally, cnidarian cnidoblasts (aka nematoblasts) produce cnidocysts within an enlarged golgi apparatus [30], and myxozoans’ capsulogenic cells produce polar capsules within the golgi apparatus (from vesicles arising in the rough endoplasmic reticulum [20]). Furthermore, “both myxozoan polar capsules and cnidarian nematocysts consist of a capsule whose wall is continuous with a coiled tubule that everts from its apical end” [31].

Since the most fundamental evidence of homology is molecular, the discovery of “nematocyst” proteins (nematogalectins and mimicollagens) shared by cnidarian cnidoblasts and myxozoan polar capsules [17, reviewed 25] clinched the case for the homology [13-20,26-28,31]. Consequently, Myxozoa was moved out of Protista into a parasitic cnidarian cnidocyst! Sequence comparisons of complete small subunit ribosomal RNA coding regions demonstrate that “myxozoans emerged after the cnidarians, and thus could not give rise to a cnidarian endobiont that eventually might have evolved into the nematocyst” [15, emphasis added]. In other words, homologies notwithstanding, cnidarians came first! Myxozoan polar capsules arose from cnidarian cnidocysts and not vice versa.

Status of the Myxozoa/Cnidaria Relationship

Myxozoa does not sit comfortably on the metazoan tree. “The unexpected diversity in the genomic organization” [36] presented difficulties for ascertaining where the myxozoans resided among metazoans. Ambiguity surrounded comparative analyses of HOX gene clusters [37,38] and small subunit ribosomal RNA (SSU rRNA), 18S and 16S ribosomal RNA (rDNA) and 5S and 5.8S rRNA [15,39-41]. Conflicting evidence pointed toward a closer relationship of microbial eukaryotes with Bilateria (metazoans excluding Porifera, Cnidaria, Ctenophora, and Placozoa [15,19,42,43]) or with Radiata (aka, Coelenterata: Cnidaria, Ctenophora, and Placozoa) [14,16-18]. This conflict may be resolved, however, were Cnidaria reassigned to the Bilateria (not unjustified morphologically given some anemones’ mirror-image symmetry around the mid-sagittal plane).

Presently, Myxozoa has two classes, Myxosporea and Malacosporea (comprising Tetracapsuloides and Buddenbrockia [44]) (The previous classes, Myxosporea and Actinosporea [20], are probably “alternating life stages of a single organism” [13]). Myxozoa, as such, constitutes an “unranked subphylum” within Cnidaria [45], a “sister to Medusozoa within Cnidaria” [31], or a class “within the phylum Cnidaria, on the medusozoan lineage” [27]. Placement is yet to be made and the question remains open whether “myxozoans are highly degenerate cnidarians” [17] or extreme members of a parasitic branch of Cnidaria [18,46,47].

Comparative morphology

Robert Weill [4] drew attention to the protist/cnidarian comparison while describing seventeen recognizable cnidocysts (now nearly thirty morphological types or as few as four proteomic types [based on mimicollagen genes]). Different types of cnidocyst [5-8] are adapted to different functions from procuring prey, defence, aiding digestion, and (if rarely) supporting structure. The averted tube may deliver a dose of venom, ensnaring glue, promote mixing currents, or reinforce the wall of burrows [48-53].

In as much as “it is very dangerous at present to lay too much stress on the value of nematocysts, pending a thorough investigation of variability and occurrence” [54], cnidocyst morphology has played a limited role in cnidarian taxonomy. Even the identity of particular cnidocysts may be uncertain. For example, the atrichous isorhizers (-a: without; trich: spines; iso-: same; rhiza: tube: a uniformly slender tube that everts from its apical end” [31].

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Attrichous isorhizas are also present in the Medusozoa (Staurzoa, Scyphozoa, Hydrozoa, and Cubozoa). Their presence in both cnidarian subphyla (Anthozoa and Medusozoa) recommends them as originary, although holotrichous isorhizas (holo-: uniform), present throughout the Hexacorallia, are rarely present in the Medusozoa. Both cnidarian subphyla harbor a family of cnidocysts consisting of varieties of isorhizas and mastogophores, but Filifera and Capitata (Anthoathecata, Hydrozoa) alone harbor a second family consisting of microbasic euryteles (eury-: broad; tele: end: tube dilated at end of butt [rare in thecate hydrozoans]) and desmonemes (desmos- bond; nema: thread: binding thread) or volvents (volvere: to turn or twist: thick tube forming an ensnaring corkscrew at discharge), while stenoteles (steno: short) or penetrants (butt of tube with small spines in spiral pattern, “thorns” and long slender basal “styles”) occur (almost) exclusively in the Capitata hydrozoans [57,58]. Among anthozoans, hexacorallians uniquely harbor spirocysts (may not be homologous to other
Cnidoblasts (individually or in small clones) derived from amoebic cells or, in the anemone Nemastomella vectensis from “cuboidal-shaped epithelial cells... detected both, in the ectodermal and entodermal tissue layers” [65] produce cnidocysts, migrate to tentacles (or other sites, such as the mesenterial filaments and acoitia of anemones and gastrodermis of corals). Having once begun differentiating their cnidocyst, cnidoblasts settle into extracellular pockets. In the epidermis, cnidoblasts become cnidocytes when they form sheet-like filaments de-somose with specialized epidermal battery cells and become competent for discharging their cnidocyst. [30,43,50]. Epithelial cells provide “launch sites” and proximity to sensory and ganglionic cells. Early pansporoblasts contain eight primordial spore envelopes, each with a polar capsule connected to a primordial sporoplasm. Sporoplasmic masses coalesce with spore envelopes in mature spores as polar capsules differentiate in their capsulogenic cells.

In Aurantiactinomyxon, pansporoblast cells surround inner cells whose meiotic divisions produce gametes (shedding polar bodies in the process). Following fertilization (nuclear fusion), diploid sporoblasts organize into future sporoplasm, valvogenic, endospore, and capsulogenic cells. Early pansporoblasts contain eight primordial spore envelopes, each with a polar capsule connected to a primordial sporoplasm. Sporoplasmic masses coalesce with spore envelopes in mature spores as polar capsules differentiate in their capsulogenic cells.

Capsulogenic cells perform the function of cnidoblasts. Ultimately, the “structure of the polar capsules, and more importantly, of the capsulogenic cells, is exactly like that of the nematocysts in coelenterates” [i.e., the cnidocyst in Cnidaria] [20]. And like cnidoblasts, capsulogenic cells locate themselves strategically in the environment of shell valves aimed at a discharge canal in the pansporoblast. Although the launch mechanism is unknown, the polar tubule wends its way toward the host.

Developmental biology

Cnidarian cnidoblasts first appear among inner cells of embryos before the emergence of polarity. In embryos of the marine hydrozoan, Pennaria tiarella, “the inner cells will give rise to endoderm and interstitial cells, while the outer cells will form larval ectoderm” [63] consisting of epithelial and mucous (gland) cells. Soon thereafter, however, only a “few developing interstitial cells and nematoblasts are found in the central cores of the embryos” [63] and fewer still remain central when the mesoglea has formed and separated epidermis from gastrodermis. By 82 hours post-fertilization, interstitial cells (i.e., amoebic cells), nematoblasts (cnidoblasts), and ganglionic cells constitute thirty-one percent of the pre-metamorphic planula’s outer cells. These cells have moved from the inner cell mass into the epidermis.

In polyps, amoebic cells are individual, small, densely basophilic, and with relatively large nuclei and nucleoli [64]. In hydra, these cells tend to populate interstices at the base of epidermal cells (hence, “interstitial cells”), while in other cnidarian polyps and medusas, amoebic cells occupy the expanded inter-epithelial compartment as well.

Cnidoblasts, therefore, correspond to transitional amplifying cells on mammalian cells in culture” [72]. Thus, cnidarian amoebic cells do not correspond to embryonic stem cells (ESCs) or induced pluripotent stem cell (iPSCs) [73]. Rather, amoebic cells correspond to vertebrate adult stem cells (ASCs) with limited potential. Cnidoblasts, therefore, correspond to transitional amplifying cells (TACs) already differentiating along determined lines.

Epithelial cells have even more restricted competence. These cells have been shown numerous times to lack the potential to modulate into amoebic cells or any of their derivatives. For example, in hydra treated in any of a number of ways (colchicine, nitrogen mustard, hydroxyurea, urethane, or lowered temperature), interstitial cells are lost and with them germ, nerve, ganglionic, and gland cells [74,75]. When the loss is complete, the specialized cells do not regenerate, and behaviour suffers. The hydras do not move or eat, but if they are fed mechanically they grow, bud, and add supernumerary tentacles.

Similarly, Pennaria tiarella planulas treated with colchicine, podophyllotoxin, or vinblastine sulfate suffer the “complete elimination of interstitial cells, nematoblasts, and sensory-motor interneurons” [76]. In depleted planulas “the remaining cells... become organized into a distinct epidermis and gastrodermis separated by a mesogela” albeit the successfully treated planulae die without metamorphosing into polyps [76].

If the missing amoebas are restored to hydras through grafting with normal head or foot tissue, the epithelial hydras re-acquire the missing specialized cells (germ, secretory, and, at least, some nerve cells [74,75], sperm [77] and egg [78]). The results with similarly grafted Pennaria planula are comparable except “reconstituted planular head pieces contain interstitial cells, ganglionic cells and a reforming neural plexus but few nematoblasts/nematocytes. Reconstituted planula tail pieces contained interstitial cells and nematoblast/nematocytes but no ganglionic cells” [79].

Myxosporidians have no definitive embryonic stage, but their sporoegenic cells (sometimes called ova) are comparable to cnidarian amoebic cells by way of performing the role of adult stem cells and giving rise to cells of different types. For example, in Myxobolus, following a nuclear proliferative phase, multiple polyplid vegetative and diploid generative cells are carved out of a plasmodium. Generative cells pair and one cell becomes the sporoegenic cell, while the other becomes a percyte that envelopes the sporoegenic cells. The percyte divides and forms the pansporoblast, the mature spore’s outermost envelope. The sporoegenic cells divide meiotically to form valvogenic cells, the sporoplasmic sporoblast and capsulogenic cell.

The amoebic cells of polyps and medusas are self-renewing [70] and the source of individual or small clones of cnidoblasts [71], but cells lack “myc, nanog, klf4, Oct4, and Sox2 genes that confer pluripotency
Origins of Cnidaria/Myxozoa

Several fossils from the Precambrian, Ediacaran Doushantuo Formations have the appearance of myxospora in the form of cellular spheres enclosed in larger spheres. Specifically, phosphorite specimens from Weng-an, “in addition to the blastomere-like cells… [contain] one or more spheroidal to ellipsoidal multicellular structures, here termed matryoshkas in reference to their similarity to nested Russian dolls. The matryoshkas are of variable size (30–350 μm) but they are generally larger than blastomere-like cells. They themselves are multicellular, consisting of tightly packed cells (9–14 μm in size) that are significantly smaller than the blastomere-like cells. Measurements show that the matryoshkas do not follow a palintonic cell division pattern…. [Rather,] matryoshkas are growing structures, with cytoplasmic growth after each division to restore cell size” [80].

The similarity of matryoshkas to sporoplasm within sporoblasts is simply uncanny. Thus, the fossil record suggests that microscopic forms of cnidarians (as well as the macroscopic forms [81]) may have existed in the Ediacaran/Vendobionta.

This new image of Buss and Seilacher’s cnidocyst-less ur-cnidarian [23] goes back to the Ediacara [82]. It is not a “choanoblastaeae” [83], an aggregate of collar cells, but epithelial shell. The image is reminiscent of Placozoa [84], and suggests that placozoan-like forms might have constituted a common stem with cnidocyte-free cnidarians.

Represented today by Trichoplax adhaerens, Placozoa is “the most basal metazoan known… Genetic evidence also points to a position close to the last common metazoan ancestor” [85]. The placozoan’s plate-like or compressed-sphere structure consists of epithelial-like layers (with marginal belt desmosomes but without adhering junctions or basal lamina) joined across an interspace by a vaguely connected (syncytial?) meshwork of stellate contractile cells. The thin upper layer of flagellated cells contrasts with the thicker lower layer of cylindrical cells and non-flagellated gland cells. Stellate, contractile fibre cells are found in the interspace between the layers [86].

Despite its protein shape, Trichoplax is not bereft of the rudiments of symmetry. A “full coding region, spatial expression and function of Trox-2, the single Hox/ParaHox-type gene identified in Trichoplax… is expressed in a ring around the periphery of Trichoplax, in small cells located between the outer margins of the upper and lower epithelial cell layers. Inhibition of Trox-2 function, either by uptake of morpholino antisense oligonucleotides or by RNA interference, causes complete cessation of growth and binary fission” [86].

Under crowded conditions in the laboratory, Trichoplax is capable of producing egg (and sperm?), suggesting that germ cells arose in the ancestral animal. Oocytes are derived from ventral epithelium, but they are surrounded by a dense wicker-work of fibre cells that develops in the interspace. At the same time, large numbers of small non-flagellated round presumptive sperm appear in the interspace amidst fibre cells. In the laboratory, embryos are expelled from the degenerating “mother” at or before the 64-cell stage, although further development presumably occurs in the wild [87].

If the radial/bilaterian branch(es) of the metazoan tree exhibit a pattern of accumulation of amoeba-cells and diversification within epitheli-like shells, symbiogenetic integration [88] would seem to begin with amoebic cells in Trichoplax adhaerentes (however sparse and rudimentary) becoming competent for germ cell differentiation. The more abundant amoebic cells populating ctenophores induce nerve and give rise to muscle, as well as differentiate into germ cells. Cnidaria exhibits an even greater accumulation of amoebic cells to the extent that, in contemporary hydrozoans, cnidocytes alone comprise almost half the cell population (e.g., estimates based on Hydra attenuata [89]) and along with germ cells, nerve, sensory cells, muscle, and gland cells comprise the major cellular component of modern cnidarians [90]. Cell differentiation is also more sophisticated. For example, the box jellyfish Tripedalia cystophora’s sensory clubs (rhopalia) are equipped with statocysts, two lens eyes, and two pairs of simpler pit and slit eyes [91]. Bilaterians show even greater symbiogenic diversification (i.e., head ectoderm taking on mesodermal qualities [the epithelial to mesenchymal transition] and mesoderm taking on epithelial qualities [the mesenchymal to epithelial transition]) [90].

For Katja Seipel and Volker Schmid, however, “it appears that the most parsimonious hypothesis taking into account the recent molecular, cellular, and developmental data is based on a motile life form with mesodermate-like development as a common ancestor of Ctenophora, Cnidaria, and Bilateria” [92]. This “mesodermate hypothesis” proposes “a motile tri-layered cnidarian ancestor and a monophyletic descent of striated muscle in Cnidaria and Bilateria. As a consequence, diploblastic evolved secondarily in cnidarian larvae and polyplps” [92].

However neatly the mesodermate hypothesis accounts for the origin and broad occurrence of muscle, the hypothesis raises another problem: Whatever happened to Cnidaria that limited its evolutionary potential to two subphyla plus Myxozoa (of whatever status) in contrast to the proliferation of bilaterian stocks? Conceivably, Placozoa and Ctenophora were limited in their potential by relatively sparse amoebic cells, but cnidarians did not suffer a similar disadvantage. Could it be that Cnidaria’s stunted evolution is grounded in their amoebic cells having uniquely acquired cnidocytes?

Acquiring Cnidocytes

Assuming that the absence of cnidocytes in fossils of scyphozoan-like medusas appearing in the Vendian biota [81,93,94] is not due to taphonomy and diagenesis but genuinely to the absence of cnidocytes, one assumes that a cnidarian/myxozoan branching occurred somewhere between six hundred and four hundred fifty million years ago. Thus, cnidarian origins presumably occurred in the preCambrian prior to if not simultaneous with the origins of bilaterians.

A consensus view of these radiations of metazoa remains elusive due to problems of sufficient sampling of non-bilaterian taxa and appropriate out group choices. Presently, however, broad taxon sampling of genomes supports the branching of Porifera, Placozoa (each with germ cells), and “eumetazoa” (containing nervous systems: Ctenophora, Cnidaria, and Bilateria) [95,96].

The question of whether Cnidaria branched off stem-eumetazoa or early bilaterians remains problematic, since “genome mining and molecular phylogenetic approaches” demonstrate that “a muscle protein core set, including a Myosin type II Heavy Chain motor protein characteristic of striated muscles in vertebrates (MyHC-st), was already present in unicellular organisms before the origin of multicellular animals” while no “protein correlates with the evolutionary origin of muscle… suggest[ing] that the core contractile apparatus in eumetazoa muscles antedates the origin of the animal kingdom and that lineage-specific innovations underlie muscle evolution in cnidarians and bilaterians.” Indeed, “tree topology strongly indicates that the myhc-st and myhc-nm genes had already
separated in the last common ancestor of all animals and the aforementioned protists, with the latter having later lost myhc-st" [97].

Similarly, "[p]hylogenetic analyses indicate that the MHCIa / MHCIb duplication is more ancient than the divergence between extant metazoan lineages." Indeed, "Class II myosins originated in unikonts, i.e., eukaryotes ancestrally bearing a single flagellum or no flagellum, including the amoebozoans, fungi and holozoans (e.g., choanoflagellates and multicellular animals or Metazoa)" [98]. Thus, evidence does not exclude the possibility that the convergence of muscle proteins is traceable to pre-metazoan evolutionary source(s) even in non-muscle proteins.

Occon's razor would seem to shave down the muscle "synapomorphy (shared derived character) of the Eumetazoa (Cnidaria + Ctenophora + Bilateria), together with nerve cells" [98] to postbranching evolution due to mutation, duplication, and selection leading to a convergence in bilaterians. The origins of muscle proteins such as Class II myosins, however, would seem to have predated the branching of cnidarians and other eumetazoans. Indeed, the presumptive evolution of muscle proteins does not exclude the possibility that cnidoblastic genes present in symbiotic amoeba moved by horizontal gene transfer into the genome of cnidarian symbiont(s).

The analysis of the "composition of venomous and structural proteins forming one of the most sophisticated organelles in the animal kingdom" [99], namely, cnidarian cnidocyst, likewise, favours neither the possibility of evolution within cnidarians nor origins from a foreign source stem genes. Indeed, the "final injected nematocyst payload comprises a mixture of dynamically evolving proteins involved in the development, maturation, maintenance, and discharge of the nematocysts, which is unique to each organism and potentially to each nematocyst type" [100].

Bacteria with a protrusion apparatus would seem excellent candidates for genes encoding cnidocyst precursors. Of course, one can only hypothesize that proto-cnidarian amoebic cells were hosts to histozoic symbiotrophic bacteria equipped with an infectious apparatus.

Imagine infectious prokaryotes losing their virulent edge and becoming adapted to a symbiotic lifestyle within amoebic cells in proto-cnidarians. Ultimately, bacterial genes would have to have moved into the nucleus of amoebic cells thereby establishing the genomic foundation for cnidarian cnidocysts. Subsequently, through intra-organismic competition and selection, cnidoblasts would have evolved with all the variations found among cnidocysts coupled to the unique migratory behaviour to epithelial sites and triggering mechanisms.

Anticipating this possibility, Richard Christen et al. [1] commented, "We should note that the mixing of genetic material between two different organisms was probably not as difficult more [than] a billion years ago than it is now, and that transfection of genetic material still exists between present day symbionts such as a plant and Agrobacterium." The notion of gene transfer and incorporation would seem consistent with the abundance of transposable elements and other evidence of horizontal gene transfer in cnidarians as well as the presence of "non-metazoan genes among cnidarian ESTs [expressed sequence tags]" [101]. Indeed, "a flip gene [has] entered a medusozoa genome from the genome of a unicellular organism... in the lineage that gives rise to the gern line (i.e., the interstitial cell lineage)" [102].

Several features of cnidarians relationships with bacteria may be relevant here. First, not surprisingly, like other animals, cnidarians bear their quota of epithelial-bound, stably colonizing microbial communities of various phylogenotypes [28,103]. Indeed, "microscopic analysis... revealed numerous bacteria within all epithelial cells in all Hydra oligactis polyps analysed irrespective of whether the animals were taken from the long-term laboratory culture (n = 15) or directly from the wild (n = 5)" [71], and some "[n]on-metazoan genes among cnidarian ESTs... are candidates for horizontal gene transfer [HGT]" [71] including "seventy-one Hydra gene models showing closer relationships to bacterial genes than to metazoan genes based on sequence similarities and phylogenetic analysis" [71]. Another "90 transposable elements... were potentially horizontally transferred into the Hydra genome" [71].

What is especially intriguing as well as surprising is that budding in brown hydras (if not in algal-laden green species) ceases when bacterial loss is induced experimentally. On the other hand, budding is restored when the bacteria are re-introduced [104]. This intimacy of bacterial symbiosis to cnidarian sexual reproduction would seem to implicate bacteria in cnidarian evolution albeit not necessarily as the source of cnidocysts.

Finally, the notion of cnidian amoebic cells having been infected and subsequently colonized by bacterial genes raises the possibility that despite their wealth of amoebic cells, proto-cnidarians did not undergo evolutionary ramifications along the lines of bilaterians or evolve a central nervous system and complex organ systems (beyond gonads, gullet, mesenteries, canals., medusoid statocysts and rhopalia) because the production of cnidocysts prevented cnidian evolution and diverted it largely toward the elaboration of opportunistic hunters in the mold of polypl, medusas, and parasites.

Summary and Conclusions

Cnidaria and Myxozoa are linked by molecular, morphological, and developmental homologies begging for a phylogenetic explanation. Microsporidians are not sufficiently homologous to Cnidaria to account for cnidocysts, and Myxozoa bearing cnidocysts in the form of polar capsules arose from Cnidaria and could not have given rise to cnidian cnidocysts. Rather, Ediacaran precursors are imagined to have accumulated amoebo-like cells within epithelial-like shells and evolved in several directions. In contrast to metazoans acquiring mesoderm, germ layers, and bilateral symmetry, cnidocyst-less cnidarians may have acquired cnidocysts from infectious bacteria equipped with an extrusion apparatus. Bacterial genes subsequently may have moved to the cnidian genome programming amoebic cells for cnidocyst production. The branching of Cnidaria from bilaterians, thus, may have been due to cnidarians specializing in cnidocyst evolution while failing to elaborate the complex organs characteristic of bilaterians.

References


