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The unique skeleton of siliceous sponges (Porifera; Hexactinellida and Demospongiae) that evolved first from the Urmetazoa during the Proterozoic: a review

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Abstract. Sponges (phylum Porifera) had been considered as an enigmatic phylum, prior to the analysis of their genetic repertoire/tool kit. Already with the isolation of the first adhesion molecule, galectin, it became clear that the sequences of sponge cell surface receptors and of molecules forming the intracellular signal transduction pathways triggered by them, share high similarity with those identified in other metazoan phyla. These studies demonstrated that all metazoan phyla, including Porifera, originate from one common ancestor, the Urmetazoa. The sponges evolved prior to the Ediacaran-Cambrian boundary (542 million years ago [myr]) during two major “snowball earth events”, the Sturtian glaciation (710 to 680 myr) and the Varanger-Marinoan ice ages (605 to 585 myr). During this period the ocean was richer in silica due to the silicate weathering. The oldest sponge fossils (Hexactinellida) have been described from Australia, China and Mongolia and are thought to have existed coeval with the diverse Ediacara fauna. Only little younger are the fossils discovered in the Sansha section in Hunan (Early Cambrian; China). It has been proposed that only the sponges possessed the genetic repertoire to cope with the adverse conditions, e.g. temperature-protection molecules or proteins protecting them against ultraviolet radiation.

The skeletal elements of the Hexactinellida (model organisms Monorhaphis chuni and Monorhaphis intermedia or Hyalonema sieboldi) and Demospongiae (models Suberites domuncula and Geodia cydonium), the spicules, are formed enzymatically by the anabolic enzyme silicatein and the catabolic enzyme silicase. Both, the spicules of Hexactinellida and of Demospongiae, comprise a central axial canal and an axial filament which harbors the silicatein. After intracellular formation of the first lamella around the channel and the subsequent extracellular apposition of further lamellae the spicules are completed in a net formed of collagen fibers.

The data summarized here substantiate that with the finding of silicatein a new era in the field of bio/inorganic chemistry started. For the first time strategies could be formulated and experimentally proven that allow the formation/synthesis of inorganic structures by organic molecules. These findings are not only of importance for the further understanding of basic pathways in the body plan formation of sponges but also of eminent importance for applied/commercial processes in a sustainable use of biomolecules for novel bio/inorganic materials.

1 Introduction

The origin of the first ancestor of all metazoan phyla remained enigmatic until the identification of first sequences coding for informative proteins from a sponge (phylum Porifera) by application of molecular biological techniques (Pfeifer et al., 1993). Before it had been only speculated that the sponges are true metazoans that are composed of individually reacting and acting cells (see: Pechenik, 2000). With the isolation of the first sponge sequence, a galectin, it became overt that these animals have the genetic toolkit (reviewed in: Pilcher 2005) to allow their cells to differentiate from omnipotent via pluripotent to finally determined somatic cells (reviewed in: Müller, 2006); most of the functional analyses were performed with the sponges Suberites domuncula and Geodia cydonium (see: Müller et
With the first cell-matrix adhesion molecule, an integrin (Pancer et al., 1997; reviewed in: Nichols et al., 2006), it could be substantiated that the sponges represent organisms that are composed of cells expressing cell surface molecules allowing their cross talks and in turn also divisions and restrictions of their physiological functions (Müller and Müller, 2003). The individuality of a sponge specimen and the morphogenetic arrangement of its cells according to a defined body plan was underscored by the discovery of apoptotic as well as organizer-specific axis-forming molecules in S. domuncula (reviewed in: Wiens et al., 2004; Müller, 2005; Wiens and Müller, 2006). The next challenge was to understand the environmental factors which drove the evolution of the phylogenetically oldest animals, the sponges. One of the key elements, promoting the emergence of these animals was silicon, which displays both morphogenetic and also structural properties (Krasko et al., 2002).
2 Role of silicon and silicate

Most skeletal structures in Metazoa are composed of calcium-salts, e.g. calcite or apatite. However, as has been found in vertebrates, silica deposition is seen prior to the ossification process forming calcium-apatite bones (Carlisle, 1986). The morphogenetic role of silicon is not restricted to mammals, but has also been identified already in sponges (Krasko et al., 2000). In S. domuncula this element causes especially the expression of genes that are required for the formation of the skeletal elements (Krasko et al., 2002; Müller et al., 2004a; Müller, 2006). Silicon/silica is used in the two sponge classes Hexactinellida and Demospongiae, as starting substrate for the enzymatic synthesis of their silica-based spicules which are the key structures, allowing the formation/arrangement of differentiated cells within an individual according to a body plan (Wiens et al., 2006; Müller, 2006).

Silicon/silicate is usually precipitated passively onto inorganic matrices, e.g. Roman glasses (Fig. 1A), or on organic matrices like in diatoms (Fig. 1B). In sponges, however, the formation of “polymerized”/condensed silicate is governed by an enzyme termed silicatein (Cha et al., 1999). Even after this information is available, it remains to be studied which morphogenetic factors allowed the formation of up to two meters large and highly elaborated sponges, e.g. the hexactinellids Monorhaphis chuni (Fig. 1C) or other hexactinellids like Hyalonema species (Fig. 1D) and also demosponges, e.g. Lubomirskia baicalensis (Fig. 1E). The question arises: When and in which environment did the sponges appear/evolve?

3 Evolution during the Proterozoic: evolution of sponges in the silicon ocean

It is surprising that 542 million years ago [myr], near the Ediacaran-Cambrian boundary, a rapid appearance of different animal types occurred. Exciting examples of these emergences were the so-called Ediacaran biota, which are older than the Cambrian animals (Knoll and Caroll, 1999). The development and divergence of the major animal clades were surely driven by the genetic tool kits available at that time. The major, or perhaps even only, metazoan phylum with hard skeleton that existed at the border from Ediacaran to Cambrian (approximately 543 MA) which did not become extinct are the sponges. Consequently, sponges were also termed “living fossils” (Müller, 1998): they represent the evolutionary oldest, still extant taxon which testifies the developmental level of the animals that lived in the Neo-Proterozoic eon (1000 to 520 MA); Figs. 2 and 3. This must be especially mentioned since two major “snowball earth events” occurred, the Sturtian glaciation (710 to 680 myr) and the Varanger-Marinoan ice ages (605 to 585 myr), which very likely caused the covering of the earth by a continuous ice layer (Hoffmann and Schrag, 2002). It was proposed that as a consequence of these ice ages most species went extinct, perhaps more than 85% (Hoffman et al., 1998).

The primordial earth surface comprised initially insoluble silicates and carbonates as well as, to a small extent, phosphates. During the silicate weathering-carbonate precipitation cycle, prior or simultaneously with the glaciations, a dissolution of these surface rocks composed of insoluble silicates [CaSiO₃] resulted in formation of soluble calcium carbonate [CaCO₃] and soluble silica [SiO₂], under
consumption of atmospheric CO$_2$ (Walker, 2003). The resulting soluble minerals leached into the waters of the rivers, lakes and oceans. There, they were again re-precipitated into new minerals, as part of the sedimentary rocks. Such processes depend upon temperature, pH and atmospheric carbon dioxide. Passively, the minerals are transformed diagenetically to secondary structures.

In contrast to passive re-precipitation, biogenic deposition of minerals by metazoans is first seen in sponges. The oldest sponge fossils (Hexactinellida) have been described from Australia, China and Mongolia (from more than 540 MA) (Gehling and Rigby, 1996; Brasier et al., 1997; Li et al., 1998). Hence, the Hexactinellida are the oldest group of sponges as documented there and later in fossil records of the Sansha section in Hunan (Early Cambrian; China; Steiner et al., 1993; Steiner, 1994). In both lower and upper levels of the Niutitang Formation more or less completely preserved sponge fossils, e.g. Solactiniella plumata (Fig. 4A-a to Fig. 4A-c), have been discovered (Steiner et al., 1993). The occurrence of almost intact sponge fossils also in the basal part of this formation is real (Rigby and Guang, 1996) and is stratigraphically equivalent to the Chengjiang assemblage in Yunnan (China). So far the base of the Niutitang Formation has been correlated with the Tommotian black shales, “Badaowan” Member. However, the transgression that deposited the black shale of the Niutitang Formation and the “Badaowan” Member was a diachronous event, which first became evident in the basin environment, Hunan/SE – Guizhon/S-Anhui, and progressed across the platform, Yunnan/Sichuan (Steiner, 1994). Therefore, the base of the
Niutitang Formation, which contains basinal phosphate concretions and cherts, may be correlated with the Tommotian or even the Nemakit-Daldynian. The important point is that the sponges at the Sansha section definitely occur at lower levels than the famous, diverse fauna at Chengjiang. Except for one questionable demosponge, all the sponges described from the Sansha section can be attributed to the Hexactinellida.

Very likely the oldest isolated spicules were also found in China, observed in thin sectioned material from the Dengying Formation, “Shibantan” Member, Hubei Province (Steiner et al., 1993). These spicules are mainly monaxial spicules, but some are also rather definite “crosses”, the evidence of triaxones from hexactinellids. The “Shibantan” Member in Hubei is of Late Proterozoic age and stratigraphically equivalent with the Ediacara (South Australia), which is famous because of its exceptionally preserved Vendian fossils. The Ediacara fauna has normally been considered to begin at about 600 MA, although new stratigraphic data place the base of the Ediacara fauna at ≈565 MA, and suggest that it ranges up to the Precambrian/Cambrian boundary (Grotzinger et al., 1996). Still, the Ediacara-type Vendian fossils are the oldest megafossils which may, at least partly, be interpreted as metazoa, although this interpretation is still controversial (Seilacher, 1989; Morris, 1993; Morris, 1994; Retallack, 1994). It is thus justified to call the Porifera the oldest organismic group in the Earth’s history, which can now be proven as definitely belonging to the animal kingdom, and which furthermore has survived successfully until present times. In contrast, to the marine Hexactinellida and Demospongiae, the freshwater demosponges are much younger; one of the first fossil freshwater sponges, *Spongilla gutenbergiana*, has been described from the Middle Eocene (Lutetian); (Müller et al., 1982; Fig. 4B-a to Fig. 4B-c).

The earliest evidence of demosponge fossils have been found in 750 my old strata of Nevada (Reitner and Wörheide, 2002). Hence, the records of Demospongiae and Calcarea presence of isolated spicules in thin sections of rocks from Early Cambrian (Atabatian) Archaeocyath mounds of the Flinders Ranges (South Australia) (reviewed in: Li et al., 1998) are younger. The Calcarea documented from the Flinders Ranges Archaeocyath mounds are both, isolated spicules and also more or less complete rigid skeletons of small calcaronean sponges sitting on Archaeocyaths. It had been suggested that a fourth class of sponges, the Sclerospongiae, exists composing both siliceous and calcareous skeletal elements; however these “coralline sponges” apparently have arisen several times and are not closely related. The spicules are of the triaene, equal-angular type characteristic of the modern Calcarea, but quite different from those typical of the Paleozoic Heteractinellida. Very similar, perhaps identical, spicules were described from the Early

**Fig. 4.** Fossil marine and freshwater sponge. (A) One of the oldest fossil marine sponge in body preservation are known from the Lowermost Cambrian Sansha section (Hunan, China); *Solactiniella plumata* (Hexactinellida) (A-a). This fossil is composed of highly intact spicules (A-b). (A-c) Some spicules are broken and expose the internal axial canals. (B) One of the oldest fossil freshwater sponge; *Spongilla gutenbergiana* from the Middle Eocene (Lutetian) near Messel (Darmstadt, Germany). (B-a) Spicule assembly, reminiscent of a complete animal. (B-b) Oxeas in these nests. (B-c) Oxeas with the flashing centers, representing the axial canals. Size bars are given.
Cambrian of Sardinia as *Sardospongia triradiata* and attributed to the Heteractinellida (Mostler, 1985). The complete calcarean sponge *Gravestockia pharetronensis* (Reitner, 1992) from the Flinders Ranges possessed a rigid calcitic skeleton with affinity to the modern Pharetronida. As one of the earliest families of demosponges the Geodiidae have been described on the basis of their sterrasters (Reitner, 1992) from the Flinders Ranges possessed a rigid calcite skeleton with affinity to the modern Pharetronida. As one of the earliest families of demosponges the Geodiidae have been described on the basis of their sterrasters (Reitner and Mehl, 1995).

Those taxa that survived the “snowball earth” episodes must have had the genetic repertoire to cope with the adverse conditions, meaning: (i) proteins to protect against adverse temperature, (ii) possibility to survive food restriction and (iii) – not to forget – also the protection machinery against ultraviolet radiation. During the pre-Sturtian period, atmospheric CO$_2$ had been consumed and removed from the atmosphere resulting in extreme temperature amplitudes. Sponges have cryoprotective proteins, e.g. galactose-specific lectins (Pfeifer et al., 1993; Wiens et al., 2003) which display cryoprotective protein-membrane activity, or, interestingly, also βγ-crystallins, proteins that contain only low amounts of water and hence are resistant to adverse protein folding (Krasko et al., 1997). Food restriction was probably compensated by the establishment of a symbiotic relationship with microorganisms (Müller et al., 2004b), such as gram-positive (Thakur et al., 2005) and gram-negative bacteria (Wiens et al., 2005) as well as fungi (Perovic-Ottstadt et al., 2004a). This eukaryotic-prokaryotic “labor-division” allowed the sponges a flexible and rapid adaptation to the changing environment. It is furthermore amazing that sponges have an unexpected variety of protection systems against mutagens, including also ultraviolet radiation. Several protection systems against radiation have been described; e.g. the (6–4) photolyase system (Krasko et al., 2003b) or the SOS-response-like mechanism (Krasko et al., 1998), and numerous is the literature on protection systems against environmental stress (e.g. Efremova et al., 2002). Perhaps the greatest fortune for the sponges was their ability to utilize silicic acid as substrate for their skeleton. When sponges emerged, the insoluble silicate minerals were converted into monomeric, soluble silica providing the sponges with an advantageous basis for survival and diversification, with respect to the number of the species and their abundance. In this context it should be mentioned that with the diversification and evolution of sponges (at least valid for endemic freshwater species) an increase in the gene number for silicateins occurred (Müller et al., 2006d). Early assumptions postulated that those taxa which survived mass extinction, e.g. the Foraminifera, became ecological and morphologically generalized species (Cifelli, 1969). Furthermore, the number of species, the diversity of the biota, increased rapidly after each period of extinction (Futuyma, 1986). Hence, the sponges (survivor taxon) became the beneficiaries of the glaciation crises and received the chance to colonize those habitats which had been de-populated (see also: Butterfield, 2007).

The urmetazoans/sponges (Müller et al., 2001) had already the basic genetic toolkit for all derived metazoans which emerged during the “Cambrian Explosion” (Figs. 2 and 3). This statement implies that the genetic repertoire of the sponges, which survived the glaciations, gave the frame, potentials/potentialities but also the limits of the body plan construction seen in higher derived taxa and which exist in the present day animal phyla. The derived taxa utilized the pre-existing molecules and pathways for their diversification of patterning and for an increase in the genetic network complexity. It can be postulated that during the progress of evolution the degree of redundancy decreased on the expense of an increase in complexity. This progressive “perfection” might be detrimental to the stability and survival of most of the species which are evolutionary younger.

The oxygen level in the atmosphere and in the water during the early Proterozoic was lower than at present (Towe, 1970; Hayes, 1994). It had been postulated that with the rise of oxygen level the synthesis of collagen became possible (Towe, 1981); sponges contain and express collagen genes (Garrone, 1998). The ability to form these extracellular fibrils had been considered to be a crucial prerequisite for the origin of multicellular animals and the establishment of a metazoan body plan (Towe, 1981). Until now, no experimental evidence for an existence of a primordial, blood oxygen-transporting system in sponges has been found. Therefore, we assume that most of the oxygen required by these animals to allow intermediary metabolism is provided by diffusion. Perhaps oxygen is partly transported/generated via the tyrosinases (Müller et al., 2004b). However, this apparent disadvantage was surely of benefit for the sponges to survive the oxygen deficiency in waters because of an ice cover during the glaciations.

4 Unique formation and degradation of biosilica in sponges: silicatein and silicase

Sponges are sessile filter-feeding organisms; their body is composed of an epithelial layer which surrounds a mesohyl compartment; this is reticulated in a highly organized manner by a canal system. The main structural and functional novelties, which evolved during the major evolutionary transitions to the Porifera, are summarized in Fig. 3. One characteristic trait of the demosponges and hexactinellids sponges are the spicules (the sclerocytes) which stabilize the sponge bodies and provide the platform on which the body plan can develop (reviewed in: Müller et al., 2004a). In the center of the spicules is a hollow canal of varying diameter around which the silica is deposited under formation of concentric layers (Uriz et al., 2000; Uriz et al., 2006). An organic filament, called axial filament, around which the axial canal had been formed is composed of the enzyme silicatein (Fig. 5). Interestingly, the spicules with their characteristic features, the axial canals, can be identified
in fossil sponges (Fig. 4A-c). The inorganic silica phase of the siliceous spicules contains 6–13% water, yielding an approximate formula of \((\text{SiO}_2)_{2.5} \cdot \text{H}_2\text{O}\) (reviewed in: Sanford, 2003). High resolution magnetic resonance microimaging studies revealed that this water is largely present in a “mobile” form at least in certain freshwater sponges (Müller et al., 2006b). In addition, spicules contain trace amounts of other elements, mainly S, Al, K, and Ca, but also Cl, Fe, Na, Zn, and Cu (reviewed in: Uriz et al., 2003; Sanford, 2003).

One major breakthrough in the understanding of spicule formation came after the discovery of the key enzyme involved. The group of Morse discovered that the organic filament in the central axial canal of spicules is composed of a cathepsin L-related enzyme, which they termed silicatein (Shimizu et al., 1998; Cha et al., 1999); Fig. 5. They cloned two of the proposed three isoforms of silicateins, the \(\alpha\)- and \(\beta\)-forms, from the marine demosponge Tethya aurantium (Cha et al., 1999). In subsequent years these molecules were cloned also from other sponges, among them the marine sponge Suberites domuncula and the freshwater sponge Lubomirskia baicalensis (Schröder et al., 2004b; Wiens et al., 2006).

From S. domuncula the cDNA encoding silicatein was isolated and characterized; the predicted translation product of 330 amino acids has a calculated size of \(M_r 36306\). Silicatein is a new member of the cathepsin subfamily (Cha et al., 1999; Müller et al., 2003; Müller et al., 2005). The three amino acids Cys, His and Asn, which form the catalytic triad of cysteine proteases, are present in the sponge cathepsin at the characteristic sites: Cys\(_{125}\), His\(_{164}\) and Asn\(_{184}\). Furthermore, silicatein comprises one cluster of characteristic hydroxy amino acids (serine). We could demonstrate that silicatein occurs in the axial filament in several isoforms (Schröder et al., 2004b) and by two-dimensional gel electrophoretic analysis of the axial filament (silicatein) we could show that in this compartment silicatein undergoes stepwise phosphorylation.

A further enzyme of potential interest for application in nanobiotechnology is the silicase. This enzyme is able to depolymerize amorphous silica (Schröder et al., 2003a). The cDNA encoding the silicase has been identified in primmorphs from S. domuncula, applying the technique of differential display of transcripts. The expression of the silicase gene is strongly upregulated in response to higher silicon concentrations in the incubation assays (Schröder et al., 2003b), like the expression of silicatein. The deduced polypeptide has an \(M_r 43131\). Silicase is a member of the family of carbonic anhydrases. An alignment of the sponge silicase sequence with the human carbonic anhydrase II shows that most of the amino acids which are characteristic for the eukaryotic-type carbonic anhydrase signature are also present in the sponge silicase (Schröder et al., 2003b). The recombinant silicase was found to display besides carbonic anhydrase activity the ability to dissolve amorphous silica under formation of free silicic acid (Schröder et al., 2003b).

Carbonic anhydrases are a family of zinc metal enzymes (Sly and Hu, 1995). In the S. domuncula silicase the three conserved His residues which bind the zinc ion are found in the deduced protein at aa\(_{181}\), aa\(_{183}\) and aa\(_{306}\). The proposed mode of action of the silicase to depolymerize amorphous silica follows the reaction mechanism known for other zinc-dependent enzymes involved in ester hydrolysis. The zinc ion is a Lewis acid that interacts with water, a Lewis base. The zinc-bound hydroxide ion formed by splitting the water molecule mediates a nucleophilic attack at one of the silicon atoms linked by oxygen bond(s). In the next step the zinc-complex binds, within the polymeric silicate, to one silicon atom under cleavage of the oxygen linkage. Through consumption of \(\text{H}_2\text{O}\), the silicic acid product is released and the zinc-bound hydroxide is regenerated allowing the start of the next catalytic cycle. The enzyme has been expressed in Escherichia coli using the glutathione S-transferase (GST) fusion system (Schröder et al., 2003b).

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**Fig. 5.** Phylogenetic relationship of the silicateins, the enzyme which catalyzes the polymerization process of biosilica in the sponge spicules. Four deduced silicatein sequences of the isoform silicatein-\(\alpha\) \([\alpha-1, \alpha-2, \alpha-3\) and \(\alpha-4\)] from Lubomirskia baicalensis (SILICAa1_LUBAI; SILICAa2_LUBAI; SILICAa3_LUBAI; SILICAa4_LUBAI) and the two cathepsin L sequences (CATL_LUBAI; CATL2_LUBAI) were aligned with silicatein-\(\alpha\) from Suberites domuncula (CATL_SUBDO) and the two cathepsin L sequences (CATL_LUBBAI; CATL2_LUBBAI) aligned with silicatein-\(\alpha\) from S. domuncula (SILICAa_SUBDO). Two major groups of silicateins, Tethya aurantium (T. aurantium) (SILICAa_TETYA) and with the \(\beta\)-isoenzymes from S. domuncula (SILICAB_SUBDO) and Tethya aurantium (SILICAB_TETYA), as well as with the cathepsin L sequences from sponges Suberites domuncula (CATL_SUBDO), Geoidea cydonium (CATL_GEOCY, Y10527) and Aphrocallistes vastus (CATL_APHRVAS) as well as from the cnidarian Hydra vulgaris (CATL_HYDRA; AA065603). An unrooted phylogram was constructed after the alignment of these sequences. The numbers at the nodes are an indication of the level of confidence for the branches as determined by bootstrap analysis (1000 bootstrap replicates).

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Recent studies have shown that during spicule formation in primmorphs of *S. domuncula*, the expression of both enzymes, silicatein and silicase, is seen at those sites within the 3D-cell aggregates where the growth/formation of the siliceous structures starts (Eckert et al., 2006). The role of the silicateins as synthesizing enzymes also was elucidated by in situ hybridization studies (Schröder et al., 2004a). In a recent study immunofluorescence analyses have been applied to demonstrate that in primmorphs silicatein is associated with fibrous structures in primmorphs that represent the primordial spicules. In parallel with the application of anti-silicatein it could be shown that also the silica catabolizing enzyme [silicase] is localized at the same structures within the primmorphs. This result had been supported also by immunofluorescence studies of sections through tissue samples.

5 Hexactinellida: first approaches to understand spicule formation

The monophyletic group of Hexactinellida is, according to the form and organization of its spicules, divided into two subclasses, Hexasterophora and Amphidiscophora. One member of this subclass is the family Monoraphididae which includes also the three species – *Monoraphis chuni*, *M. dives* and *M. intermedia*. Since the discovery of these sponges during the first German Deep Sea Expedition (RV "Valdivia") and the first descriptions of these animals by Chun (1900) and Schulze (1904), only little additional information has been added about the form and construction of their skeletal systems. Nothing has been published about the synthesis of their giant spicules. Worldwide *M. chuni* has been documented only at few sampling sites, suggesting a distribution in the deep sea in the Indian Ocean and in the West Pacific. One outstanding feature of *Monoraphis* is the anchoring spicule which can reach lengths of up to 3 m with a maximum diameter of 8.5 mm (Schulze, 1904). *M. chuni* thus produces the largest biosilica structure known on Earth. Polished cross sections confirm the existence of up to 500 highly regular concentric rings which are arranged around a 150 µm thick, less structured core (Schulze, 1904). In the middle of this core, a square axial channel with an approx. 2 µm thick proteinaceous filament is visible, which
runs through the entire spicule from one end to the other (Fig. 6A, C and D). The silica layers are composed of smaller granules and spheres within a network of (perhaps) proteinaceous material (Fig. 5B).

An additional hexactinellid *Hyalonema sieboldi* likewise possesses long stalk spicules that attach them to the substratum; the animals live in a depth of more than 600 fathoms (1000 m); (Wyville Thomson, 1874 and also Müller et al., 2006c). The long stalk spicules are composed of distinct siliceous layers which are also superposed in a stratified pattern around a central axial filament (Schulze, 1904). It could be demonstrated that these giant spicules from the root-tuft of *H. sieboldi* transmit light with high efficiency. Surprisingly, however, the blue light with a wavelength between 400 and 600 nm is filtered out. Data elaborated in our group suggest that the spicules from *H. sieboldi* act as optical absorbent in a novel photoreception system (Müller et al., 2006c).

Very recently, Aizenberg et al. (2005) published a detailed structural analysis of the spicule formation in the hexactinellid *Euplectella*. They demonstrated structural hierarchies of the spicule synthesis starting from the nanometer-sized particles of silica to the final mature spicules. The compositions of the proteins which are associated with and found in the spicules have not been described. In a recent study Ehrlich et al. (2005) dissolved the basal spicules of the hexactinellid *H. sieboldi* in alkaline solution for 14 days and showed that the abundant structural protein, associated with the spicules might be collagen.

We performed microscopic analyses of the spicules from *M. chuni* with major emphasis of the large-sized giant spicules (giant basal spicules or basalia; size of 1 m) and also the large comitalia (small di- or tri-actine spicules of a size of around 60 mm). The focus of the study was put on the organic components of these spicules, including the collagen fibrils which surround them. Electron microprobe analysis data show a regionally different distribution of sodium and potassium within the spicules. With respect to the organic components, it should be highlighted that after dissolution of the spicules several proteins could be visualized; (i) one of them cross-reacted with antibodies raised against silicatein (Wang et al., 2007; Müller et al., 2007), while (ii) others displayed proteolytic activity (to be published).

After having substantiated, by application of molecular biological techniques, that also Hexactinellidae have at least one gene coding for collagen (the cDNA was isolated from the hexactinellid *Aphrocallistes vastus*) a schematic outline of spicule growth both in longitudinal and transversal direction could be given (Fig. 7). In the center of the spicules the axial canal is filled with an axial filament whose protein composition is highly similar to that described for demosponges. After formation of a first silica layer, assumed to be the result of silicatein or of a silicatein-related protein, the next layer is formed through centrifugal and centripetal synthesis of a silicatein(-related) protein. In analogy to Demospongiae (Schröder et al., 2006) it is adopted that the silicatein(-related) protein is associated with a lectin which forms a tube-like cage around the existing silica lamella. After deposition of newly formed silica clusters followed by a solid silica lamella the existing proteins are hydrolyzed by protease(s). In continuation a third lamella of silica is formed. The model also proposes that collagen guides the silicatein(-related) protein/lectin associates concentrically along the spicules.

### 6 Demospongiae: silica deposition during spicule formation

The biochemical basis for the understanding of the morphology of the spicules has been analyzed recently using the marine demosponge *S. domuncula* (reviewed in: Müller et al., 2006a). From this species primmorphs were prepared and applied for the analysis of spicule formation. Primmorphs, a special type of 3D-cell aggregates, containing proliferating and differentiating cells, allow to study the differentiation of archaeocytes to sclerocytes during incubation in medium supplemented with silicic acid. The spicules were analyzed...
Fig. 8. Morphology of demosponge spicules showing the zonation of the silica shell of the spicules from Superites domuncula. (A to C) SEM analysis of broken spicules in primmorphs, after an incubation period of 20 days. Around the axial canal (ac) two layers of lamellae (l1 and l2) can be distinguished. In B the spicule have been etched order to visualize the lamellae. (C and D) Immunostaining of spicules with antibodies, raised against silicatein (PoAb-aSILIC). (D) Immunostaining of cryosections through tissue of S. domuncula. 8-µm thick frozen sections were reacted with polyclonal silicatein antibodies. Spicules (sp) and one canal of the aquiferous system (ca) are marked. (E and F) Reaction of antibodies against silicatein with partially digested (hydrofluoric acid) spicules. The preparations were placed onto glass slides and treated with the antibodies. (F) The immuno-stained specimen showed that both the surface of the spicules (sp) and also the axial filaments (af) were brightly stained.

by electron microscopical procedures, e.g. scanning electron microscopy (SEM) analysis of spicules, transmission electron analysis (TEM) and electron immunogold labeling. The skeleton of S. domuncula is composed of only two types of megascleres, monactinal tylostyles and a smaller fraction of diactinal oxeas. The spicules reach up to 450 µm in length and diameters of 5 to 7 µm (Fig. 8A to C); they grow through apposition of lamellar silica layers. While the two ends of the oxeas are pointed, the tylostyles have one pointed end and one swollen knob. All spicules have an axial canal of 0.3 to 1.6 µm in width in the center. As shown in SEM images (Fig. 8A to C) the central canal is surrounded by lamellated layers approximately 0.3 to 1 µm thick. To monitor the location and distribution of silicatein in and around the spicules, polyclonal antibodies were raised against purified filaments from spicules in rabbits (Müller et al., 2005). Immunofluorescence studies with cryosections through tissue revealed that the surfaces of the spicules in the tissue (Fig. 8D) and of isolated spicules (Fig. 8E and F) are surrounded by silicatein. From these studies we conclude that silicatein is not only present in the axial filament of the spicules, but is also located on their surface.

According to our results the process of spicule formation should be divided into the following phases; the initial intracellular steps and the extracellular final and shaping phase (Müller et al., 2006a and Eckert et al., 2006). (i) Intracellular phase: Silica is taken up actively by a Na+/HCO₃⁻[Si(OH)₄] cotransporter. Very likely during the transport through the endoplasmic reticulum and the Golgi complex, silicatein undergoes phosphorylation and is transported into vesicles where it forms rods, the axial filaments (Fig. 8A); around them the first layer(s) of silica is (are) formed. Silica deposition occurs in two directions; first from the axial canal to the surface (centrifugal orientation) and second from the mesohyl to the surface of the spicule (centripetal). Finally the spicules are released into the extracellular space where they grow in length and diameter by appositional growth (Fig. 8B).

(ii) Extracellular phase (appositional growth): Silicatein is present also in the extracellular space. It came surprising that also there the silicatein molecules are organized to larger entities. Immunogold electron microscopical analysis showed that the silicatein molecules are arranged along strings, which run parallel to the surfaces of the spicules (Schröder et al., 2006). In the presence of Ca²⁺, silicatein associates with galectin and allows the appositional growth of the spicules (Fig. 8B). Since the surface of a new siliceous spicule is also covered with silicatein, the appositional growth/thickening of a spicule hence proceeds from two directions (centrifugal and centripetal).

(iii) Extracellular phase (shaping): In the next step, the galectin-containing strings are organized by collagen fibers.
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tic acid is a fermentation product, and thus proved the basis
for the discipline biochemistry. The causal-analytical un-
derstanding of organic reactions in biological systems be-
came possible after the deciphering of the genetic code and
the subsequent elucidation and application of molecular bi-
ological techniques. It took, however, until recently when first strategies had been formulated and ex-
perimentally proven, to outline the biosynthesis of inorganic
structures formed in uni- and multicellular organisms. At
present, in a self-accelerating progress, the matrices (tem-
plates), e.g. collagen, and the organic catalysts (enzymes),
e.g. silicatein, required for the synthesis of those inorganic
structures and skeletal elements have been illuminated with
the help of inorganic/organic chemists, biochemists, molec-
ular biologists and material scientists, to establish the disci-
pline “bio-inorganic material science”. The first opportuni-
ties have been touched in biomedicine and electronics pro-
viding us with the chances, power and potential of silica-
technology (see: Schr¨oder et al., 2007 and Wang et al., 2007).

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