Modularity and Distribution of Sulfur Metabolism Genes in Bacterial Populations: Search and Design

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Abstract

Biological Engineering involves global DNA sampling and modular design from genetic parts. A new approach reflected by natural history is based on the recognition of interchangeable DNA fragments that move around the world due to horizontal gene transfer. According to the large scale, metagenomics provide opportunities to sequence whole genomes within environmental populations. Annotated gene sequences, protein structures, and metabolic data can be used to design small biosystems from interchangeable genetic parts, the same as from functional modules.

To illustrate this, the 21 genes for sulfur metabolism were inferred from the genome of bacterium Vesicomyosocius okutanii HA, and the distribution of two gene clusters (dissimilatory sulfite reductase – dsr and sulfur-oxidation – sox) within environmental samples was investigated. The correlation between the dsr and sox clusters for the experimental set of 41 stations was R = 0.86 which demonstrates the complementarity of dsr and sox metabolic pathways in environmental populations. Hypothetical functions were assigned using comparisons with known proteins. The 18 reads from symbions of gutless worm Olavius algarvensis showed a high identity to large AprA protein from V.okutanii. In addition, comparative 3D modeling of hypothetical DsrB protein revealed sulfite reductase ferredoxin-like half domain, sulfite reductase 4Fe-4S domain, and a repressor of phase-1 flagellin.

The simplistic reconstruction of sulfur metabolism from parts and examples of hierarchical modularity in nature are given. The origin of modularity is considered in the context of minimal cell and horizontal gene transfer. The role of ancient sulfur metabolism in modularization is discussed under the umbrella of iron-sulfur world theory (Wächtershäuser, 1988), deep-hot biosphere model (Gold, 1992), and radiolysis hypothesis (Garzón and Garzón, 2001). The reverse engineering approach based on natural genetic modules is proposed for understanding early life.

Keywords: Constructive biology; Sulfur metabolism; Metagenomics; Gene geography; Horizontal gene transfer; Minimal genome; Origin of life; Compositional evolution

Introduction

Instead of small gradual changes like point mutations, the mechanisms of compositional evolution combine interdependent genetic modules that have evolved previously in parallel (Watson, 2006). Examples of compositional mechanisms in nature include hybridization (Rieseberg et al., 2003), horizontal gene transfer (HGT) (Doolittle, 1980; Jain et al., 2002), and symbiotic encapsulation (Margulis, 1970; Merezhkovsky, 1909), as exhibited in the history of major evolutionary transitions (Maynard Smith and Szathmáry, 1995). Both gradual and compositional mechanisms of biological evolution are mediated by natural selection (Darwin, 1859). On the other hand, progress in the development of complex man made systems, like software products, has demonstrated a benefit of modular design that sometimes leads to an ignorance of evolution. In this light, the search for natural building blocks in DNA records and attempts to design artificial biological systems from those complementary modules might help us appreciate the beginning of life and the dynamics of evolutionary progress up to the modern life forms. As Richard Watson wrote: ‘the existence of modularity in nature is now becoming testable’ (Watson, 2006). To prove this statement, metagenomics deals with the global gene set where each genome is studied as part of a biological community and DNA sequences are analyzed from the viewpoint of ecology.

Let us consider sulfur metabolism in a symbiotic community as an inspiration for a ‘protocell’. This approach will help us to recognize modules in complex systems. For instance, hemoglobin in the giant tube worm Riftia pachyptila binds hydrogen sulfide (H₂S) which allows the worm to reduce toxicity and transmit the sulfide into symbiotic bacteria which supply the host with nutrition (Flores et al., 2005; Chabasse et al., 2006). In general, reduced inorganic sulfur compounds can be used as electron donors. Oxidation of sulfide to sulfate takes place in the bacterial cytoplasm by using dissimilatory sulfite reductase (DsrAB, EC 1.8.1.2) operating in reverse, adenosine-5’-phosphosulfate reductase (APS reductase, EC 1.8.99.2) and ATP sulfurylase (EC 2.7.7.7) (Pott and Dahl, 1998). Genes involved in oxidative sulfur metabolism and CO₂ fixation via the Calvin-Benson-Bassham cycle were characterized in some metagenomics projects (Beller et al., 2006; Scott et al., 2006; Kuwahara et al., 2007). These genes are mainly localized in clusters. The large dsr cluster encodes the dissimilatory sulfide oxidation and provides the assembly of electron transport complex III (Dahl et al., 2005; Pires et al., 2006). The sox genes cluster is used to oxidize thiosulfate taking place in the periplasmic space (Friedrich et al., 2001). Some genes encoding sulfide quinone oxidoreductase, rhodenase, and ATP sulfurylase are under peculiar transcriptional control (Beller et al., 2006). Sulfate, a final product of the sulfur oxidation under anoxic conditions, is
global ocean sampling (gos) expedition puts together microbial databases and for visualization. for instance, the 'sorcerer ii' clear, camera is a metagenomic data set with tools for querying ecology research and analysis – camera (rusch et al., 2007). to be retrieved from community cyberinfrastructure for advanced marine microbial information – ncbi (benson et al., 1998), expert protein analysis system – expasy (appel et al., 1994; gasteiger et al., 2003), kyoto encyclopedia of genes and genomes – kegg (kanehisa andgoto, 2000) and integrated genomics – ergo (overbeek et al., 2003), were used to find proteins and gene assemblies in order to infer sulfide oxidation metabolic networks that may be small enough for optimization, synthesis, and experimental evaluation. data obtained were compared with environmental samples available from community cyberinfrastructure for advanced marine microbial ecosystem research and analysis – camera (rusch et al., 2007). to be clear, camera is a metagenomic data set with tools for querying the databases and for visualization. for instance, the 'sorcerer ii' global ocean sampling (gos) expedition puts together microbial communities every 200 miles around the globe. in addition, camera v1.3.2 contains a vertical profile and deep-water samples of marine microbial communities collected in hawaii and in the eastern mediterranean, 150 genomes of ocean microbes, symbionts of mediterranean gutless oloquiochaeta olavius algarvensis and hydrothermal vent polychaete alvinella pompejana, as well as other collections including samples from the acid mine drainage biofilm and rich farm soil (sheshadri et al., 2007). these sequence data were scanned for genes coding sulfur metabolism.

the aim of this paper is to map proteins involved in sulfur metabolism within various bioisous. molecular reconstructions are tested for signs of modularity. the article poses a problem of origin of modules in biological systems and particularly addresses the compatibility of modules due to the lateral gene transfer. the methodology combines the analysis of environmental databases and the design of metabolic pathways. the scheme includes a transport of hydrogen sulfide, accumulation of polysulfide, sulfide, and thiosulfate oxidation into sulfate, and finally a sulfate export. produced electrons will be used to atp generation via a respiratory electron-transport chain. this 'energetic module' will be installed on an imaginable 'minimal cell' with a set of housekeeping genes (gabaldon et al., 2007; gibson et al., 2008; gil et al., 2004; glass et al., 2006; koonin, 2003). the gene assembling includes sox and dsr clusters from the sulfur oxidizing bacterium candidatus vesicomyosocius okutanii ha with a small completely sequenced genome (~1 mb), which is a symbiont of a deep-sea clam calyptogena okutanii (kuwahara et al., 2007). the artificial system assumes a particular advantage for symbiotic bacteria and their host. some ideas for this model were retrieved from (dahl et al., 2005; pires et al., 2006), even though the symbiotic relationships might be more complex (arnold et al., 2001). preliminary data from global dna sampling and their analysis, as well as common ideas concerning biological design and extension of life based on natural interchangeable modules have been considered.

materials and methods
databases, such as the national center for biotechnology information – ncbi (benson et al., 1998), expert protein analysis system – expasy (appel et al., 1994; gasteiger et al., 2003), kyoto encyclopedia of genes and genomes – kegg (kanehisa and goto, 2000) and integrated genomics – ergo (overbeek et al., 2003), were used to find proteins and gene assemblies in order to infer sulfide oxidation metabolic networks that may be small enough for optimization, synthesis, and experimental evaluation. data obtained were compared with environmental samples available from community cyberinfrastructure for advanced marine microbial ecosystem research and analysis – camera (rusch et al., 2007). to be clear, camera is a metagenomic data set with tools for querying the databases and for visualization. for instance, the 'sorcerer ii' global ocean sampling (gos) expedition puts together microbial communities every 200 miles around the globe. in addition, camera v1.3.2 contains a vertical profile and deep-water samples of marine microbial communities collected in hawaii and in the eastern mediterranean, 150 genomes of ocean microbes, symbionts of mediterranean gutless oloquiochaeta olavius algarvensis and hydrothermal vent polychaete alvinella pompejana, as well as other collections including samples from the acid mine drainage biofilm and rich farm soil (sheshadri et al., 2007). these sequence data were scanned for genes coding sulfur metabolism.

to reduce the complexity of a system and to model dynamics of biochemical networks, the jdesigner v2.1 simulator was used (see link to manual in references). the networks were described by first order differential equations (ode) with arbitrary parameters allowing a qualitative interpretation. camera fragment recruitment viewer enabled the comparison of complete bacterial genomes with environmental databases (rusch et al., 2007). a family of blast programs was applied for quick searches in large databases (altschul et al., 1997). the significance of hits was estimated by a bit score and by an expectation value. bit score is a measure of similarity between the hit and the query that is derived from a raw score by normalization, the e-value revealing how many times an expected result could occur by chance. the higher the score, the better the alignment; the lower the e-value, the more significant the score. strong values for a bit score near 400 and the expectation below 1e-100 were preferred during alignments and superior querying. e-value threshold, larger than 1.0, was biologically relevant against environmental dna datasets in preliminary scans. t-coffee v5.68-7.71 servers were used for the automatic multiple sequence alignments and an annotation (notredame et al., 2000); the improved visualization was with boxshade v3.21 utility. empirical evaluation of the quality of multiple sequence alignment was prepared within regular parameters; residues with a yellow to red background are correctly aligned more than 80%. protein domain identification was carried out within interproscan (mulder et al., 2007) and pfam (bateman et al., 2000; bateman et al., 2002) databases. the comparative protein structure analysis was done with default parameters by three server machines which complemented each other – esyfpred3d (lambert et al., 2002), 3d-jigpsaw v2.0 (bates et al., 2001) and geno3d v2.2 (combet et al., 2002). protein visualization was made by an spdb viewer v4.0 (guev and petsch, 1997). links to the online servers are mentioned in the references. some techniques such as the normalization and correlation analysis are described in results.

| product name                        | start  | end    | strand | length | gene id          | locus  |
|-------------------------------------|--------|--------|--------|--------|------------------|-------|
| atp sulfurylase                      | 98093  | 99301  | +      | 402    | 5172354          | sfr   |
| adenyllysufate reductase membrane anchor | 99516  | 100365 | +      | 289    | 5171803          | apoA  |
| adenyllysufate reductase l-subunit   | 100417 | 100896 | +      | 159    | 5171218          | apoB  |
| adenyllysufate reductase             | 100896 | 102779 | +      | 627    | 5171222          | apoA  |
| sulfite oxidation protein SoxB       | 172596 | 174485 | +      | 629    | 5177231          | soxA  |
| sulfite oxidation protein SoxA       | 770792 | 771607 | -      | 271    | 5177240          | soxA  |
| sulfite oxidation protein SoxZ       | 771335 | 771937 | -      | 100    | 5179133          | soxA  |
| sulfite oxidation protein SoxX       | 771971 | 772414 | -      | 147    | 5179200          | soxA  |
| sulfite oxidation protein SoxX       | 772425 | 772772 | -      | 115    | 5179188          | soxA  |
| intracellular sulfite oxidation protein DsrR | 817196  | 817537 | -      | 113    | 5172350          | dsrR  |
| intracellular sulfite oxidation protein DsrR | 818938  | 820140 | +      | 400    | 5172402          | dsrR  |
| intracellular sulfite oxidation protein DsrR | 820691  | 820897 | +      | 281    | 5172402          | dsrR  |
| intracellular sulfite oxidation protein DsrJ | 820694  | 821277 | -      | 127    | 5172414          | dsrJ  |
| putative glutamate synthase (nadph) small subunit | 821307  | 823271 | -      | 654    | 5172337          | dsl   |
| intracellular sulfite oxidation protein DsrP | 823492  | 824926 | -      | 921    | 5177242          | dslP  |
| intracellular sulfite oxidation protein DsrM | 824894  | 825667 | -      | 257    | 5177239          | dslM  |
| intracellular sulfite oxidation protein DsrC | 825744  | 826067 | -      | 107    | 5177235          | dslC  |
| intracellular sulfite oxidation protein DsrC | 827204  | 829209 | -      | 327    | 5177232          | dslC  |
| intracellular sulfite oxidation protein DsrA | 828373  | 829674 | -      | 433    | 5177231          | dslA  |
| rhodanese family protein             | 950273 | 950752 | +      | 159    | 5179184          | cost  |
| sulfur-quinone reductase             | 955954 | 957240 | +      | 428    | 5172159          | sfr   |

Table 1: Genes coding sulfur metabolism in the bacterium vesicomyosocius okutanii/ha.
The resume of data available from CAMERA v1.3.2 used in experiments is as follows. The Hawaii Ocean Time Series (HOT) station collected genomes down to 4000 m (DeLong et al., 2006). The other metagenomics study was in the Ionian abyssal plain, a deep flat basin between Sicily and Greece in the Eastern Mediterranean, whose deep waters are free from an intrusion of cool polar waters that feed the bottom of the World Ocean. The next sample is the deep-sea hydrothermal vent polychaete worm, Alvinella pompejana, which maintains diverse microbial symbionts. The microbial community was isolated from the dorsal integument of A. pompejana collected from the East Pacific Rise at a depth of 2500 m. Another worm, Mediterranean gutless oligochaete Olavius algarvensis, lives with four microbial symbionts having different metabolic pathways. The symbiotic bacterial consortium demonstrates mutualistic relationships with their host and with each other (Woyke et al., 2006). The purpose of the Acid Mine Drainage biofilm sequencing project was to investigate the diversity of metabolic pathways in microorganisms in the sample obtained from the underground was localized within a pyrite (FeS2) ore body. oxidation, nitrogen fixation, and iron oxidation. The sample obtained from the underground was localized within a pyrite (FeS2) ore body. This biogeochemical system provided a coupling between microbial iron oxidation and acidification due to pyrite dissolution (Lo et al., 2007; Tringe et al., 2005). Waseca County is well-known for rich, black soil that produces record crops every year. Surface soil (0-10 cm) was collected from a farm in Waseca County, Minnesota. Microscopic analyses revealed the presence of various prokaryotic organisms in the sample. PCR-amplified 16S rRNA sequences from the sample confirmed numerous bacterial and archaeal lineages (Tringe et al., 2005). Additional information about samples is available from the CAMERA web page.

**Results**

**Comparison of four thiotrophic bacterial genomes with CAMERA global databases**

Initially, I needed to have a suitable thiotrophic reference to scan global databases for sulfur genes. The CAMERA Fragment Requirement Viewer was used to compare complete genomes of two thiobacteria and two chemosynthetic endosymbionts with subsequent data-bases. GOS – All Metagenomic Sequence Reads (N), Metagenome of Marine NaCl-Saturated Brine and Microbial Community Genomics at the HOT/ALOHA (Figure 1). Free-living microorganisms such as Thiomicrospira crunogena XCL-2 and Thiobacillus denitrificans ATCC 25259 demonstrated multiple homologies with different stations (depicted in color). Spikes with more than 90% identity corresponded to 5S, 16S, and 23S ribosomal RNA, various tRNAs, and housekeeping genes that are common in many bacteria. T. crunogena with a genome of 2 427 674 nt in length presented a maximal number of matching sequences (63 134) which is difficult to study. A large set of T. denitrificans genes showed about 60-85% similarity to the microbiota at Sargasso Station 11 (Figure 1B, cyan). In contrast, symbiotic bacteria Candidatus Vesicomyosoccus okutanii HA and Candidatus Ruthia magnifica Cm represented less complex patterns of homology allowing analysis and reasonable interpretations (Figure 1C,D). Bacterium V. okutanii was chosen as a reference for future research.

**Gene collection**

Efficient browsing of NCBI, KEgg, and ERGO databases inferred metabolic pathways for a ‘minimal’ sulfur metabolism within bacterium Vesicomyosoccus okutanii HA. This symbiotic microorganism has a small single chromosome (1 022 154 nt) and is annotated (Kuwahara et al., 2007). As a result of the search, the 21 genes comprising 19 656 nt from V. okutanii were assigned to the hydrogen sulfide oxidation into sulfate (Table 1). The bacterium V. okutanii demonstrated a very compact organization of sox and

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**Figure 1: Fragment requirement plots for four thiobacteria in CAMERA Viewer.** Abscissa is the genome length, ordinate is the 50 to 100% homology comparisons with the databases, such as All Metagenomic Sequence Reads (N) – Global Ocean Sampling Expedition, Metagenome of Marine NaCl-Saturated Brine, and Microbial Community Genomics at the HOT/ALOHA; the visualization of stations is done by inverted colors.

**Figure 2: Fragments of V. okutanii HA genetic map with genes coding sulfur metabolism.**
Strong transport, \(k_2 = 0.1\); weak transport, \(k_2 = 0.01\); the sulfate output is 0. The same low ratios were assigned to cytoplasmic components, such as sulfide transporter, DsrAB, AprAB, and Sat transform the sulfide into sulfate in the cytoplasm. CT0714 permease releases the final product from the cell. The parameters were chosen as shown in Figure 3B. Initial values of sulfide and thiosulfite were high (\(=10\)) opposite the low range of sulfate (0). The same low rations were assigned to cytoplasmic components, such as sulfide transporter, DsrAB, AprAB, and Sat, and CT0714. Periplasmic proteins SoxAX and SoxB have intermediate initial values (1). All coefficients beyond \(k_2\) are equal to 0.1. The coefficient \(k_2\) is a feature of a flux between the sulfide transporter and DsrAB complex. Accordingly, the parameter \(k_2\) with values 0.1 and 0.01 describe the intensive and slow transport of hydrogen sulfide into the system.

The time course simulation revealed the role of sulfide transport in the sulfate production. The system reached a plateau for a new high-output stationary state at \(k_2 = 0.1\) very fast (Figure 4A). In contrast, the simulation of the weak transport at \(k_2 = 0.01\) showed a slow growth of output sulfate ratio, i.e. about 14 units versus 20 units at the end of the simulation (Figure 4B). The results support the assumption that the import of nutrition may be a limiting factor for the symbiotic metabolism. Multiple inputs would have led to a more reliable survival of organisms.

**Worldwide distribution of sox and dsr genes clusters**

Enzymes of sulfur metabolic network coded by sox cluster (5 genes) and dsr cluster (14 genes) from *V. okutanii* were selected for a future search in seven metagenomic databases; (1) “Sorcerer II” Global Ocean Sampling (GOS) collection, (2) bacterial consortium from the gutless worm *Olavius algarvensis*, (3) symbionts of polychaete *Alvinella pompejana*, (4) probes from the Deep Mediterranean, Ionian 3 km station, (5) data from Hawaii ocean station ALOHA, 130 m, (6) soil microbial community, Waseca County, and (7) samples from the Richmond acid mine. To compare polypeptide sequences with DNA targets, the TBLASTN program was used. Queries were formulated for each protein. CAMERA services looked for stations with the maximal number of hits and best matching sequences. Identities to sox and dsr strings were found at 41 stations (Figure 5A). The strong positive pair correlation \(R = 0.86\) between sox and dsr genes distribution in these samples was revealed. The absolute number of hits was higher for dsr cluster than for sox cluster due to the prevailing number of genes in the dsr cluster (14 vs. 5). This inconvenience was eliminated by normalization – the number of hits was divided in each sample by the deep-sea tube worm *Riftia pachyptila* is also an attractive subject for engineering because of the significant role of hydrogen sulfide transport through its body to symbiotic cells. Why would an artificial symbiotic bacterium not produce a secreted form of vestimentiferan hemoglobin in order to support itself in the host?

**Modeling of hydrogen sulfide oxidation within JDesigner**

Even the small genome of bacterium *V. okutanii* demonstrates a great complexity of sulfur metabolism; chemical reactions run stepwise and parallel and involve many different enzymes. The JDesigner simulator was used to explore connections between main nodes in the biochemical network and to model metabolic fluxes via the ordinary differential equations. The nodes represented enzymes; the arcs denoted streams of substrates and their products (Figure 3A).

Now let us look at a simplified sulfur oxidation system where hydrogen sulfide and thiosulfite are inputs and the sulfate is the output. Enzymes SoxAX and SoxB convert thiosulfite into sulfate in the periplasm. A hypothetical sulfide transporter pumps HS\(^-\) ions into cells. Enzymes DsrAB, AprAB and Sat transform the sulfide into sulfate in the cytoplasm. CT0714 permease releases the final product from the cell. The parameters were chosen as shown in Figure 3B. Initial values of sulfide and thiosulfite were high (\(=10\)) opposite the low range of sulfate (0). The same low rations were assigned to cytoplasmic components, such as sulfide transporter, DsrAB, AprAB, and Sat, and CT0714. Periplasmic proteins SoxAX and SoxB have intermediate initial values (1). All coefficients beyond \(k_2\) are equal to 0.1. The coefficient \(k_2\) is a feature of a flux between the sulfide transporter and DsrAB complex. Accordingly, the parameter \(k_2\) with values 0.1 and 0.01 describe the intensive and slow transport of hydrogen sulfide into the system.
Figure 5: Distribution of sox and dsr genes by stations. A – all positives, B – best stations.
Figure 6: Location of experimental stations with a plenty of Dsr matching sequences. A – Gutless Worm, Bay of Capo di Sant' Andrea, Elba; B – Alvinella Pompejana, East Pacific Rise; C – Farm Soil, Waseca County; D – Deep Mediterranean, Ionian 3 Km station; E – Richmond Mine; F – Sargasso Station 13; G – Upwelling, Fernandina Island, Galapagos; H – Punta Cormorant, Hypersaline Lagoon, Floreana Island, Galapagos.
the number of genes in a cluster (Figure 5B). The normalization did not change the pattern of distribution on the whole. The number of hits in the examined data decreased exponentially, station by station, and only a small fraction of samples presented a sufficient content of genes for sulfur metabolism.

Then the eight best stations from the initial 41 stations were chosen for a more detailed analysis: (1) Gutless Worm, Bay of Capo di Sant’ Andrea, Elba, (2) Alvinella Pompejana, East Pacific Rise, (3) Farm Soil, Wasco County, (4) Deep Mediterranean, Ionian 3 Km station, (5) Richmond Acid Mine, (6) Sargasso See Station 11, 13, (7) Upwelling, Fernandina Galapagos Island, (8) Punta Cormorant Hypersaline Lagoon, Floreana Island, and Galapagos Archipelago. This final collection included microorganisms from different niches such as host environments (cases 1 and 2 with a maximal number of hits), rich farm soil, deep-sea anoxic water, acid mine drainage, epipelagial zone, upwelling zone, and hypersaline lagoon (Figure 6). Microbes from these diverse environments involve similar genes in sulfur metabolism. Nevertheless, the distribution of sox and dsr genes was not the same for all samples. The normalization revealed disparities in sox and dsr gene allocation for 8 preeminent stations (Figure 5B). Bacterial symbionts of Mediterranean gutless worm *Olivarius algarensis* from the Bay of Capo di Sant’ Andrea and microbiota from the deep eastern Mediterranean demonstrated a privilege for *dsr* genes in samples. In contrast, symbionts from the farm soil in Wasco County presented both *dsr* and *sox* metabolic pathways. The content of *dsr* and *sox* genes decreased sufficiently at the next stations. The *dsr* genes dominated in microbiota from Richmond acid mine, Sargasso See station 11, 13, and the upwelling at Fernandina Island. Amazingly, the microbes living in the Punta Cormorant hypersaline lagoon show an increased role of *sox* genes for sulfur metabolism in this extreme environment.

These facts show an obvious difference between *sox* and *dsr* genes clusters. Ecological distributions of those genes appeared depending on environmental conditions. The geographical location of genomes revealed an importance of the periplasmic sox component in the oxidative sulfur metabolism for particular environments like the hypersaline lagoon.

**Investigation of individual genes and proteins involved in sulfur metabolism**

In addition to Sox and Dsr proteins, I compared the S-quinone, AprA and Sat proteins from bacterium *V. okutanii* with environmental datasets by CAMERA BLAST tools. The Gutless Worm and GOS (P) databases were used in the experiments. Values of identities with the top score were chosen to represent results. Two heterologous proteins, such as CT0714 from *C. tepidum* and *Tcr_0602* (soxZ) from *T. crunogena*, were considered as a negative control in the correlation analysis Table 2. I observed a high positive linear relationship between gutless worm *O. algarensis* bacterial symbiotic consortium and GOS (P) microbial databases with the correlation coefficient *R* = 0.85 for Dsr proteins and *R* = 0.83 for Sox proteins. The correlation was smaller (*R* = 0.67) in the third group – S-quinone, AprA, and Sat proteins. The control group – CT0714, *Tcr_0602* – illustrated a negative correlation. Dsr proteins in the line from DsrA to DsrL came across with more than 55% identity whereas the proteins DsrB, DsrE, and DsrC demonstrated higher than 70% identity for both databases. The rest of the group, polypeptides from DsJ to DsrK, had low identities. The DsrR polypeptide displayed minimal identities 35% and 20% for both databases. Only SoxX protein from the *sox* gene cluster represented about 70% identity for two target datasets. Surprisingly, AprA polypeptide from the third protein group showed maximal identities 85% and 75% for the gutless worm and GOS (P), respectively. Control proteins CT0714 and *Tcr_0602* presented relatively small identities.

Data revealed a high similarity between *V. okutanii* and queried genomes.
One should remember that the thioautotrophic bacterium Vescomyosocius okutanii is an intracellular symbiont harboring in the gill epithelial cells of the deep-sea clam, Calyptogena okutanii (Kuwahara et al., 2007). In my experiments, the adenylylsulfate reductase AprA query sequence from V.okutanii bacterial genome demonstrated a high identity to the readings from the symbiotic microbial community of gutless marine oligochaete Olavius algarvensis that inhabits the shallow sub littoral sediments on the northwest coast of Elba Island, Italy (Figure 6A). Endosymbiotic sulfate-reducing bacteria, living within this worm produce sulfide that can serve as an energy source for complementary sulfide-oxidizing symbionts (Dubilier et al., 2001). I found 18 subject DNA fragments with an E-value less than 1.35e-70, a bit score higher than 267.7, and an alignment length of 185 to 336 aa in the Gutless Worm database. The best search result is the sequence GUTLS WRM ELBA READ 1433546210 with an 85% identity. An additional search in the Gene Bank using the BLASTP program revealed a significant degree of identity (92%) between the read 1433546210 and the adenylylsulfate reductase (Gene ID: 3673410) from the β-proteobacterium Thiobacillus denitrificans ATCC 25259. A narrow subtree of AprA protein family with a close distance from AprA-like protein (read 1433546210) to homologous members of β- and γ-proteobacteria (data not shown) can indicate that the read 1433546210 belongs to a proteobacterium living in the gutless worm O.algarvensis.

Each of the 18 hypothetical AprA protein sequences was compared with each other and with the query – AprA protein from bacterium V.okutanii, Gene ID: 148244258; numbers – selected reads from the Gutless Worm database; yellow – sufficient insertions, deletions and exchanges for the AprA protein, as well as other cytoplasmic polypeptides, such as DsrB, DsrE and DsrC whereas DsrR chaperon-like protein exhibited low identity and a potential diversity in ecology.

**Discovery of aprA-gene family in thiotrophic symbionts of gutless worm O.algarvensis**

- **Figure 7: Alignment of the protein reads from microorganisms living in the gutless worm O.algarvensis to the AprA polypeptide from bacterium V.okutanii**

ApR – protein from V.okutanii, Gene ID: 148244258; numbers – selected reads from the Gutless Worm database; yellow – sufficient insertions, deletions and exchanges.
V. okutanii. The T-COFFEE automatic server found a perfect matching between short experimental reads (about 300 aa) and the large AprA polypeptide (627 aa) covering nearly all of the AprA sequence. After the manual elimination of similar fragments, I finished with the seven final sequences demonstrating numerous exchanges, insertions, and deletions (Figure 7). The read 1433708223 (327 aa long) was similar to N-part of AprA protein; and the read 1433547459 (321 aa) matched C-terminus with some variations. The most irregular read 1433688352 (314 aa) demonstrated multiple exchanges, one deletion and 2 insertions with flanking rearrangements. In addition, read 1432856135 (301 aa) showed an insertion/exchange with possible neighboring compensations in its own N-terminus. The read 1433547459 (321 aa) is very interesting because it has a long region including 28 amino residues with 2 deletions, 1
sulfur-oxidizing bacterial symbionts from gutless marine oligochaeta *Inanidrilus makropetalos* and from the giant tube worm *Riftia pachyptila*. The last data confirmed a vast diversity of AprA-like proteins discovered in bacterial symbionts of the gutless worm *O.algarvensis*. The heterogeneity might be a result of the large and dissimilar microbial community utilizing different APS reductases. This diversity provoked intriguing questions about their origin, complementary functions, and the speed of microevolution.

**Multiple alignments of AprA-, DsrA-, DsrB- and SoxX-protein families**

AprA, DsrA, DsrB, and SoxX proteins are objects for future research because of their importance in sulfur metabolism and their high degree of identity in previous experiments (Table 2, bold characters). Top matching reads for the entire proteins were selected from the best samples, such as *O.algarvensis* symbionts, *A.pompejana* symbionts, Waseca farm soil, deep Mediterranean, Richmond acid mine, Sargasso station, upwelling at Fernandina Island, and hypersaline lagoon on Floreana Island. These reads were used for comparison within each protein family by T-COFFEE multiple sequence alignment program. The highest homology (71%) was found within AprA-protein family. The DsrA and DsrB families demonstrated 61% and 66% identity respectively. The SoxX-protein family had a low matching score (48%). However, these results were affected by the different quality of databases; many hypothetical proteins were represented by their fragments. A good example is shown in Figure 8, namely the multiple alignment for reduced DsrB-protein family (identity 88%). The high identity of the core alignment to the annotated DsrB protein from *V.okutani* supposed a biological significance of this conserved sequence.

**Identification of domains within the hypothetical DsrB protein**

The DNA sequence GUTLS WRM ELBA READ 1433974420, with positions 35-973 nt coding a hypothetical DsrB protein 313 aa in length (Figure 8, upper bold string), was chosen for an annotation. This poly peptide sequence was submitted to InterProScan and PFAM databases. The InterProScan server identified the query as a sulfite reductase, dissimilatory type β-subunit encoded by dsrB-type gene. The poly peptide comprises a ferredoxin-like hemoprotein β-component in the N-terminus typical for sulfite/nitrite reductases and a 4Fe-4S region in the C-terminus also usual for sulfite and nitrite reductases. More detailed information about the protein domains in the query sequence was retrieved from the PFAM server (see annotated data in Figure 9). In addition to sulfite/nitrite reductase ferredoxin-like half domain (blue), I uncovered two 4Fe-4S binding sites (yellow) and a region for the repressor of phase-1 flagellin (red) within the large sulfite reductase 4Fe-4S domain (green). The recapitulated active center of the hypothetical DsrB sulfite reductase explains the conservatism of the area obtained in multiple alignments of fragments from different sources (Figure 8, yellow characters). An unexpected insertion, the repressor for flagellin synthesis, with a very conservative Arg residue and less conservative Asp and Ile residues, was also found in other members of the DsrB family (Figures 8, 9, aqua characters). This finding may be evidence of an autoregulation that couples energy metabolism and bacterial movement.

**3D reconstruction of the protein encoded in GUTLS WRM ELBA READ 1433974420**

Several comparative modeling servers, e.g. ESyPred3D, 3D-JIGSAW, and Geno3D were used to predict 3D structure of the poly peptide encoded in the read 1433974420. Each individual server...
retrieved only a part of the submitted sequence. The job of ESyPred3D server is depicted in Figure 9 as a bold cursive, the message from 3D-JIGSAW is shown in bold characters, and Geno3D result is represented as a cursive string. These server machines build three-dimensional models for proteins based on homologues of known polypeptide structures. The first 3D model was built by ESyPred3D program using the 2akj chain A with 2.8 Angstroms resolution. This template shared 14.2% identity with the query sequence. The 2akj protein is a spinach ferredoxin nitrate oxido-reuctase (EC 1.7.7.1) from chloroplasts whose enzyme catalyzes a limiting step in siroheme biosynthesis (Swamy et al., 2005). The other server, the Geno3D offered a large set of potential templates with a top hit for the sulfite reductase, dissimilatory-type β subunit from Halorhodospira halophila DSM 244/SLI (Gene ID: 4710578, score 323, expectation 4e−77, identities 73%). Finally, using the protein fragment retrieved by ESyPred3D server, I was able to reconstruct the C-part of the sulfite reductase ferredoxin-like half domain (blue) and the N-terminus of sulfite reductase 4Fe-4S domain (green). See Figures 9 and 10 for details. The 3D-JIGSAW server predicted two internal 4Fe-4S binding sites (yellow) within the large sulfite reductase domain (green). Geno3D server built the C-terminus of the sulfite reductase 4Fe-4S domain. The modeling started with the last Asn residue at the second 4Fe-4S binding site and finished after a repressor of phase-1 flagellin (red). Summarizing the jobs of all three servers, it was possible to reconstruct a 3-dimensional structure of the whole polypeptide encoded in the read 1433974420 except for 10 aa in the N-terminus and 14 aa in the C-terminus.

I identified two parallel α-helices on the surface of the protein core and two antiparallel β-sheets that possibly interact with the siroheme molecule inside the sulfite reductase ferredoxin-like complex (Figure 10A, blue). Remarkably, the two loops of 4Fe-4S binding site (Figure 10B, yellow) are close enough to permit the possible interaction with an iron-sulfur cluster. The repressor of phase-1 flagellin represents a small α-helix outside the core protein (Figure 10C, red). The assemblage of those three polypeptide chains into a complete protein molecule by SPDBV viewer was not easy because of an intricacy of the sulfite reductase 4Fe-4S domain (green) and a chaotic character of polypeptide terminus within each of 3D models (Figure 10D). However, the result demonstrated the possibility of reconstructing the protein structure from an environmental DNA read. The 3-dimensional model obtained is in close proximity to the intact DsrB-like protein sequence from the symbiont of amazing gutless worm Olavius algarvensis found in the Bay of Capo di Sant’ Andrea on the Elba Island in Italy.

Discussion

A minimal gene set

Starting with a minimal cell, it is difficult to define a minimal set of genes that support life (Koonin, 2000) because the complexity of life is entirely dependent on the specific environment. The minimalist approach leads to an evident controversy concerning the definition of life. For instance, the virus Porcine circovirus comprises 1 768 nt and consists of only 2 genes (Table 3). The minimal known symbiotic bacterium Buchnera aphidicola is much bigger (Perez-Brocal et al., 2006) comprising 416 380 nt and consisting of 397 genes which might confirm a difference between viruses and bacteria, as well as a distinction between nonliving and living objects. However, a huge virus like Acanthamoeba polyphaga comprises 1 181 404 nt and consists of 1 258 genes which is more than in some bacteria, such as Nanoarchaeum equitans, Mycoplasma genitalium, Mesoplasma florum, Vesicomyosocius okutanii, and Aquifex aeolicus. In this case, early ideas about the reconstruction of an ancestor genome by comparing complete modern bacterial genomes (Mushgheian and Koonin, 1996) may not be enough to make an artificial cell. An alternative could be a more realistic community metabolism in a special niche.

Recent generalizations say that biological systems are modular; and life plays with interdependent modules (Watson, 2006; Woese, 2002). Evolution acts on a combination of individual genes, gene clusters, and functional genetic systems including genetic and protein networks, as well as cellular pathways that are called here ‘modules’ (Bork et al., 1998; Marcotte et al., 1999; Overbeek et al., 1999). Metagenomic data and their annotation and analysis have revealed that the full record of proteins is far from being complete (Yooseph et al., 2007). Proteins may have appeared very early parallel to RNA and lipids worlds (Guimaraes, 1994; Segre et al., 2001; Takahashi and Mihara, 2004; Weismann, 2005). Perhaps, even the functional modules evolved very early and contributed to cellularization much earlier than previously imagined (discussed below) (Lindahl, 2004). My state-of-the-art research assumes a relatively complex kernel, a gene set comprising basic ‘living’ functions, such as core replication, repair, transcription, translation, several central metabolic pathways, possibly cytoskeleton and membrane forming, as well as cell division (Gabaldon et al., 2007; Gibson et al., 2008; Gil et al., 2004; Glass et al., 2006; Koonin, 2003). This approach includes the establishment of modules, and their combination and hierarchical design. For instance, sulfur metabolism might be organized within a single module. Modules perform special functions related to the particular environment, e.g. the hydrogen sulfide conversion (Figure 11).

Synthetic symbiont

An autonomous agent which exploits a reach environment and complements certain functions in the host organism, e.g. sulfur energetic metabolism was considered as a synthetic symbiont. It assumed that the host provided a nutrition transport (H2S) and removed wastes (SO42−). In agreement with the ‘modular logics’, the 21 genes from the genome of the symbiotic bacterium Vesicomyosocius okutanii HA were inferred (Table 1, Figure 2) and seven important enzyme activities were considered in the simplified model of food transport and processing (Figure 3). The simulation of hydrogen sulfide oxidation showed that the import of nutrition may be a limiting factor for symbiotic metabolism (Figure 4). Following dynamic modeling, the effective transport system and multiple diets might have provided a more reliable survival of the symbiotic system. That is why the sulfide- and sulfate-transporters have been proposed for use in Engineering. In some cases, a contact of host tissues with hydrogen sulfide can be reduced by the sulfide binding to hemoglobin in the blood (Arp and Childress, 1983). The artificial symbiotic bacterium would produce a secreted H2S transmitter similar to the hemoglobin from the giant tube worm R. pachyptila to protect the host organism from toxic sulfide.

The set of coupled genes (Table 1, Figure 2) and the metabolic model Figure 3 were similar to the sulfur metabolism in Chlorobium tepidum TLS. Bacterium C. tepidum can utilize thiosulfate and sulfide as a source of sulfur. The genes involved in sulfur oxidation are organized in clusters. For example, one cluster includes most of the sox genes to oxidize thiosulfate. Three flavoprotein reductases may oxidize sulfide to form polysulfides as a reserve. Further oxidation to sulfate takes place in the cytoplasm by using dissimilatory siroheme sulfite reductase (DsrABC), adenosine-5′-phosphosulfate reductase
Bacterium \textit{V. okutanii} HA was chosen as a preferred reference from other thiobacterial microorganisms, such as \textit{Ruthia magnifica} Cm, \textit{Thiomicrospira crunogena} XCL-2, and \textit{Thiobacillus denitirificans} ATCC 25259. After that, reads from 41 positive stations around the world were carefully investigated by \textsc{camera} BLAST tools for an identity to \textit{sox} and \textit{dsr} genes from bacterium \textit{V. okutanii}. Instead of evenly distributed \textit{sox} and \textit{dsr} genes by stations, numerous \textit{sox} and \textit{dsr} sequences were found mostly in the specific niches, such as the host’s environment; farm soil, deep-seep anoxic water, acid mine drainage, epipelagial zones, upwelling zones and hypersaline lagoon (Figures 5, 6). Although, \textit{sox} and \textit{dsr} genes demonstrated high correlation in experimental set ($R = 0.86$), samples with disproportions of \textit{sox} and \textit{dsr} genes were also found. One possible reason for this difference might be the heterogeneity of populations as a result of bacterial specialization and compartmentalization of reactions. Data obtained exhibited the complementarity of \textit{sox} and \textit{dsr} genes in various niches depending on the specific environment.

Current investigation has revealed a homology of \textit{Apr} protein, as well as \textit{DsrB}, \textit{DsrE}, and \textit{DsrC} proteins from \textit{V. okutanii} with the corresponding reads in \textsc{camera} databases (Table 2). In contrast, \textit{DsrR} polypeptide represented a low identity that may be a consequence of variability in chaperone genes. The 18 subject reads from gutless worm \textit{O.algarvensis} bacterial symbionts demonstrated a high similarity to \textit{Apr} query polypeptide from \textit{V. okutanii}. The protein family that was discovered, adenosine-5-phosphosulfate reductases, showed an immense diversity following different lineages. I supposed that these unknown symbionts belonged to \textit{β}- and \textit{γ}-proteobacteria. Other authors described chemoaotrophic sulfur-oxidizing \textit{β}-proteobacteria and \textit{γ}-proteobacterial sulfate reducing symbionts in \textit{O.algarvensis} (Dubilier et al., 2001; Woyce et al., 2006).

My results may not reflect phylogeny because the \textsc{blast} local sequence alignment removes the most divergent regions from the sequences, thus obscuring the relationships between sequences. Due to the low quality of databases (random and partial DNA reads), there were some difficulties during the multiple sequence alignments with the T-COFFEE program. It could be that many hypothetical proteins were represented only by their fragments. Nevertheless, I was able to reveal different patterns of variability for \textit{Apr} and \textit{DsrB} proteins. Numerous insertions and deletions in \textit{Apr} polypeptides were found, as well as prevailed amino acid exchanges in \textit{DsrB} proteins (compare Figure 7 and 8).

Kuever and co-workers (Meyer et al., 2007; Meyer and Kuever, 2007; Meyer and Kuever 2008) supposed that \textit{apr} genes are distributed between photo- and chemotrophs including invertebrate symbionts with anaerobic or facultative anaerobic lifestyles. The sulfur-oxidizing prokaryotes (SOP) turn into two phylogenetic lineages. The proteins of \textit{Arl} lineage, e.g. from \textit{Chlorobiaceae}, are similar to the enzymes from the sulfate-reducing prokaryotes (SRP). This clustering is in contradiction to the dissimilatory sulfate reductase phylogeny (Dsr) and specifies the possible horizontal \textit{aprA} gene transfer from SRP to SOB. Another paper (Klein et al., 2001) illustrated that \textit{DssrB} sequences generated a tree incompatible with the corresponding 16S rRNA phylogeny. Evidence for \textit{dsr} genes horizontal transfer was found within \textit{β}-proteobacteria, affecting \textit{Desulfobacula toluolica}. The \textit{dsrAB} genes in \textit{Archaeoglobus species} are also the result of an ancient horizontal gene transfer from a bacterial donor. Maybe, the horizontal gene transfer played a significant role in the distribution of sulfur metabolism between populations (Boucher et al., 2003). Perhaps, sulfur-oxidation (\textit{sox}) and dissimilatory sulfite reductase (\textit{dsr}) clusters, which represent different functional submodules for the energetic sulfur metabolism, were developed independently and accepted many times by various inhabitants of our planet.
Horizontal gene transfer

HGT is widespread (Andersson, 2005; Boucher et al., 2003). Communal planetary genes persist in bacterial genomes and have a tendency to cluster. Groups of travelling genes are involved in the basic molecular processes and are engaged in adaptation. It could be these genes have existed since the transition of nonliving to living matter and provided a mechanism for acquiring new functions during the exploration of various environments. The horizontal gene transfer could be a very old mechanism supporting a compatibility of modules (Brown, 2003). The ancient virus world should also be mentioned because of the essential role of viruses in evolution. About 10^{20} bp of DNA are transduced by phages every year in the World Ocean (Paul et al., 2002). Viruses can contribute to global horizontal gene transfer by moving DNA between species and biomes (Sano et al., 2004). Microbial biodiversity may change due to migration of viruses and cross-species genetic exchanges. In a recent metagenomics project, the photosystem I gene cluster was identified in genomes of marine cyanophages and is thought to be a function in photosynthesis of their hosts (Sharon et al., 2009).

Additionally, the species of lower metazoans can be an attractive target for gene transfer, e.g. a massive HGT in bdelloid rotifers was well documented recently (Gladyshev et al., 2008). The horizontal gene transfer mediated by sperm was revealed in the bivalve mollusk *Mytilus galloprovincialis* Lam. (Guerra et al., 2005; Kuznetsov et al., 2001). HGT involves particular molecular mechanisms both for prokaryotes (Smith et al., 1981) and eukaryotes (Pittoggi et al., 2002). Viruses can contribute to global horizontal gene transfer by moving DNA between species and biomes (Sano et al., 2004). Microbial biodiversity may change due to migration of viruses and cross-species genetic exchanges. In a recent metagenomics project, the photosystem I gene cluster was identified in genomes of marine cyanophages and is thought to be a function in photosynthesis of their hosts (Sharon et al., 2009).

Reconstruction of DsrB protein

The read 1433974420 from GUTLS WRG ELBA dataset exhibited a conserved protein core (Figure 8) that helped me detect domain structures within the polypeptide, such as (1) sulfite reductase ferredoxin-like half domain, (2) large sulfite reductase 4Fe-4S domain including two 4Fe-4S binding sites, and (3) a repressor for flagellin synthesis (Figure 9). Moreover, the comparative structural analysis by ESyPred3D, 3D-JIGSAW, and Geno3D servers allowed the 3D reconstruction of an almost complete hypothetical sulfite reductase, DsrB protein (Figure 10). Just compare the discovered domains side by side with known proteins.

As has been considered by other scientists, sulfite reductase is a main enzyme for both dissimilation of oxidized anions and biosynthetic assimilation of sulfur (Hansen, 1994). Sulfite reductase is a multisubunit enzyme composed of dimers of either α/β or α/βγ subunits, each containing a siroheme and iron-sulfur cluster prosthetic center (Pierik et al., 1992). For instance, the well-known Escherichia coli sulfite reductase (SIR) is a complex enzyme composed of two proteins; a flavoprotein alpha-component (SIR-FP) plus a hemoprotein beta-component (SIR-HP), demonstrating 8x4-β quaternary structure (Zeghouf et al., 2000). SIR-FP contains both FAD and FMN, while SIR-HP includes 4Fe-4S cluster coupled to a siroheme through a cysteine bridge. SIR-HP has a two-fold symmetry which causes a unique three-domain α/β fold that controls assembly and enzymatic activity (Crane et al., 1995).

4Fe-4S domain is a superfAMILY whose members bind to iron-sulfur clusters. Structure of the domain is antiparallel β-sheets. Members include bacterial-type ferredoxins and various reductases and dehydrogenases. Ferredoxins are iron-sulfur proteins mediating electron transfer in a variety of metabolic processes. They are comprehensive from bacteria to mammals and fall into several groups (Eck and Dayhoff, 1966; George et al., 1985). Most ferredoxins contain at least one conserved domain including 4 Cys residues that bind to the 4Fe-4S cluster. During the evolution of bacterial ferredoxins, gene duplications, transpositions, and fusion occurred resulting in the appearance of proteins with multiple iron-sulfur centers, e.g. bi-cluster [2Fe-4S] and polycferredoxins, iron-sulfur subunits of bacterial succinate dehydrogenase/fumarate reductase, pyruvate-flavodoxin oxidoreductase, formate hydrogenlyase and dehydrogenase complexes, NADH:ubiquinone reductase, and others. In some bacterial ferredoxins, one of the duplicated domains lost some conserved Cys residues. These domains lacked their iron-sulfur binding property. 3D structures are known both for mono- and bi-cluster 4Fe-4S ferredoxins (Duce et al., 1994; Fukuyama et al., 1989).

Flagellin subunits assemble into filaments of bacterial flagella. 1 speculated that the insertion of a repressor for flagellin synthesis into the 4Fe-4S domain might be an example of unusual regulation coupling the energy metabolism and bacterial movement. It may also be interesting evidence of the hierarchical modularity, where the flagellum-inhibiting helix is a small protein building block of low hierarchy, which is able to interconnect two larger functional modules, such as mobility and sulfur metabolism on a higher hierarchical state. DNA-binding repressor of phase-1 flagellin encoded by βflA gene was identified in the genomes of Salmonella abony and Salmonella typhimurium (McClelland et al., 2001). The peptide sequence Phe-Asp-Trp-Val-Ser-Arg-Ile at the position 166-172 from Salmonella proteins is similar to the sequence Asn-Asp-Trp-Val-Ser-Arg-Ile at the position 166-172 from Salmonella proteins is similar to the sequence Asn-Asp-Trp-Val-Ser-Arg-Ile at the position 166-172 from Salmonella proteins is similar to the sequence Asn-Asp-Trp-Val-Ser-Arg-Ile from the discovered polypeptide (Figures 9, 10, red). The virulence regulation by nutrient limitation is known for Legionella.
Conclusion, the complexity of biological systems

Unicellular life emerged about 3.5-3.8 billion years ago and another 2.5 billion years passed before multicellular organisms made their appearance. Nevertheless, it took less than one billion years for architecturally remarkable species to evolve (Fenchel, 2002; Wacey, 2009). The speed and complexity of evolution has increased visibly. It seems that gradual evolution might not explain the major evolutionary transitions indicated by fossil records (e.g. Opabinia regalis, Burgess Shale fauna, Cambrian). On the other hand, HGT can alter the speed of evolution and design by a propagation of ‘constructive mutations’ across geographical populations (Beiko and Charlebois, 2007; Hao et al., 2004; Jain et al., 2003; Meyer and Kuever, 2007). Functional combination of genes by HGT could check the fitness of interaction between genes and other components in the whole system (Watson, 2006). If life stores genetic information in a modular form, then the modular nature of genetic information makes it possible to accept useful units from other species to swap the global gene set (Beiko et al., 2005; Simonson et al., 2005). It supposes an idea regarding the dramatic level of complexity that geochemical, biochemical, and genetic systems should attain before they come alive as a result of modularization. The lithotrophic biosphere extends many kilometres into the deep sea and Earth crust (Gold, 1992). Our restricted knowledge of this kind of life stems from sampling and drilling. Deep sea and subterranean habitats may point out that lithotrophy contributes much more to the biomass of Earth than thought (Falkowski et al., 2008). It can even give insight into possible life on dwarf planets and satellites with a thermal silicate core and oceans beneath an ice surface, such as Pluto, Enceladus, Callisto, Ganymede, and Europa (Garzón and Garzón, 2001; Gold, 1992; Vance et al., 2007).

Nature has always been a source of metaphors and inspiration. In recent years, it has provoked many successful techniques. When nature is complex, some problems can be solved through design by analogy and the development of novel systems. The results of this research show that anaerobic sulfur metabolism of bacteria-invertebrate symbioses within the giant tube worm Riftia pachyptila, the deep-sea clam Calyptogena okutamii, hydrothermal vent polychaete Alvinella pompejana and the gutless oligochaete Olavius algarvensis, offers a great potential for future exploration of anoxic zones. Data recovered from official databases by NCBI and CAMERA, offers a great potential for future exploration of anoxic zones. Data recovered from official databases by NCBI and CAMERA, offers a great potential for future exploration of anoxic zones. Data recovered from official databases by NCBI and CAMERA, offers a great potential for future exploration of anoxic zones. Data recovered from official databases by NCBI and CAMERA, offers a great potential for future exploration of anoxic zones. Data recovered from official databases by NCBI and CAMERA, offers a great potential for future exploration of anoxic zones.

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