

Microbiofacies analysis of Cambrian offshore carbonates from Sardinia (Italy): environment reconstruction and development of a drowning carbonate platform

Olaf ELICKI¹

Abstract: The Campo Pisano Formation of southwestern Sardinia is represented by an offshore carbonate succession spanning the latest Early to late Middle Cambrian. Paleogeographically, the fauna is characteristic of western Perigondwana, and indicates faunal relations to France, Spain, and Turkey. Microfaunal paleoecology reflects drowning of an isolated carbonate platform at tropical latitudes. Sessile epibenthic filter feeders dominate at the base, succeeded upward by sessile, epibenthic, suspension feeders. Upsection, a shift in the ratio of faunal groups indicates increasing replacement by mobile epibenthos. Autochthonous faunal elements decline near the top where allochthonous taxa become important. In the basal portion of the formation the faunal succession indicates relatively shallow neritic habitats with a moderate influx of suspended sediment, followed by a period of slightly deeper neritic conditions. Probably a shallow bathyal environment was established at the transition to the overlying siliciclastic Cabitza Formation. The bathymetric and ecofacies changes in the Campo Pisano Formation are interpreted as having been caused by a discontinuous rise in eustatic sea-level, probably accompanied by subsidence due to tensional tectonics. The depositional environment was that of a distal open-marine shelf or ramp without strong relief. This interpretation is supported by the lack of significant lateral changes in the fossil assemblages, and an upward deepening of facies. The applied micropaleoecological approach is proven a powerful tool in regional stratigraphy and in the reconstruction of sedimentary realms in the Mediterranean Cambrian, particularly when high-resolution biostratigraphic data and diagnostic sedimentary characteristics are sparse.

Key Words: Cambrian, Italy, Sardinia, micropaleontology, microfacies, paleoecology

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Résumé : *Analyse microbiologique de séries carbonatées marines du Cambrien de Sardaigne (Italie) : reconstitution paléoenvironnementale et développement d'une plate-forme carbonatée ennoyée.*- La Formation de Campo Pisano du SW de la Sardaigne est constituée par une série carbonatée - marine - couvrant un intervalle de temps qui s'étend du Cambrien inférieur terminal à Cambrien moyen élevé. Du point de vue de la paléogéographie, la faune y est de type périgondwanien et présente des affinités occidentales certaines (France, Espagne et Turquie). L'évolution de la paléoécologie des microfaunes retranscrit l'ennoisement de cette plate-forme carbonatée isolée en climat tropical. À la base, les organismes sessiles épibenthiques filtreurs dominant. En montant dans la série, leur succèdent des formes sessiles épibenthiques suspensivores. Enfin, vers le sommet, le glissement dans les proportions au sein des associations fauniques traduit le remplacement croissant des faunes autochtones par un épibenthos vagile, c'est-à-dire par des faunes allochtones. Dans la partie basale de la formation, la succession des faunes suggère qu'à une période initiale marquée par des environnements néritiques relativement peu profonds, avec de faibles apports détritiques en suspension, a succédé une période caractérisée par des environnements néritiques légèrement plus profonds. Le stade bathyal supérieur a vraisemblablement été atteint au toit de la série, avant le dépôt de la formation silicoclastique de Cabitza. Les variations écologiques et bathymétriques de la Formation de Campo Pisano semblent résulter d'une remontée irrégulière du niveau marin, probablement couplée à une subsidence d'origine tectonique. L'environnement de dépôt était comparable à celui de la partie distale d'une plate-forme ouverte ou d'une rampe avec des reliefs peu exprimés comme en atteste l'absence d'évolution latérale significative des associations fauniques reconnues. L'analyse micropaléocologique constitue un outil puissant pour les études stratigraphiques régionales ainsi que pour les reconstitutions des ceintures de faciès du Cambrien méditerranéen, en particulier quand figures sédimentaires caractéristiques et données biostratigraphiques de haute résolution sont rares ou absentes.

Mots-Clefs : Cambrien, Italie, Sardaigne, micropaléontologie, microfaciès, paléoécologie

Introduction

In Early- to Middle Cambrian times carbonate and mixed siliciclastic-carbonate environments were widely distributed on the European Shelf of the low-latitude Perigondwana realm (Morocco, Spain, France, Sardinia, Germany, and the Near- and Middle

East; Fig. 1). But sedimentary records of these environments are commonly incomplete, due to variations in bed thickness and shelf morphology, together with widespread erosion during the Late Cambrian to Early Ordovician, as in Sardinia (Sardic Phase movements), and possibly also in Germany. Thus, a full

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reconstruction of early Paleozoic development in this region is still problematic for biostratigraphic, paleogeographic and postdepositional geological processes are not yet completely defined.

Earliest Cambrian sediments on the Gondwana European Shelf are generally not exposed, either because they were not deposited or were removed through tectonism and subsequent erosion. However, from the later Early Cambrian to the late Middle Cambrian sedimentary successions up to 2000 m or more are well developed (see BENDER, 1974; PILLOLA, 1991; GEYER & LANDING, 1995; ELICKI, 1997; LIÑAN *et alii*, 2002; ÁLVARO *et alii*, 2003). These deposits show an overall trend upward from siliciclastic to relatively shallow carbonate and mixed carbonate-siliciclastic environments, followed by a siliciclastic cover denoting deepening. Younger Cambrian deposits are only sporadically developed in some areas of the European Shelf. Although the paleontology and sedimentary facies of the carbonate successions (mainly Early Cambrian) and the siliciclastic deposits (Early and Middle Cambrian) are well known, the transition from carbonate to siliciclastic deposition, reflecting the drowning of the carbonate platforms and consequently the reorganization of the Perigondwana realm, has been investigated only minimally in some areas.

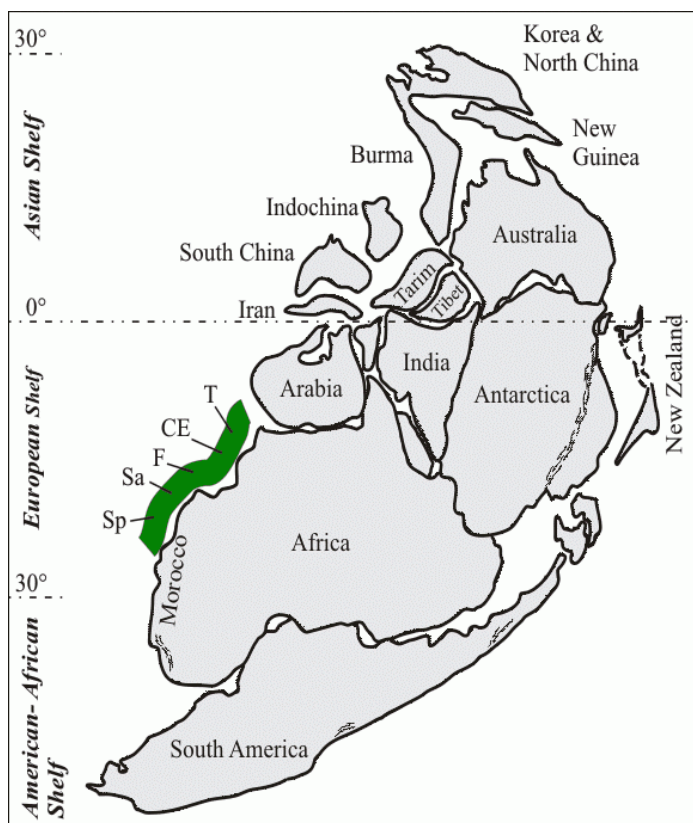


Figure 1: Paleogeographic map of Gondwana during late Early to early Middle Cambrian times with the tentative location of the Mediterranean province in dark gray (modified after COURJAULT-RADÉ *et alii*, 1992). T - Turkey, CE - Central Europe, F - France, Sa - Sardinia, Sp - Spain.

The main reason for the limited investigation of this critical interval in Sardinia may lie in the lithologic character of the transition, represented in general by rather monotonous, condensed, often nodular limestones, affected by a strong diagenetic overprint (COCOZZA & GANDIN, 1990; ELICKI, 2001). Nevertheless, the carbonates are commonly rich in shelly microfossils that usually are an important means of obtaining additional information on depositional conditions. However, here this microfossil wealth is frequently hard to bring to light. It is an extremely intricate, time-consuming and almost impossible task to extract these fossils from the host rocks if they are not diagenetically silicified or preserved as steinkerns. This problem exists for nearly all the important fossil groups other than organophosphatic brachiopods and problematic small shelly fossils that can be extracted by acidic dissolution. Furthermore, organic biota, such as palynomorphs are usually not preserved in the mostly light- and reddish-colored carbonates. These hindrances have led to a paleontological focus on archaeocyaths in Early Cambrian carbonate rocks and on trilobites in Early and Middle Cambrian siliciclastic beds. Therefore, the Middle Cambrian drowning stage of the Sardinian platform has not been properly investigated paleontologically and geological processes near the Early to Middle Cambrian boundary are not well known.

In southwestern Sardinia the drowning stage of the Cambrian carbonate platform is represented by the Campo Pisano Formation. A genetic interpretation of the depositional environment of this succession is problematic because of its limited sedimentological characteristics. Previous environmental interpretations ranged from shallow water, low-energy areas (COCOZZA, 1979) to lagoonal and partly euxinic tracts (GANDIN, 1979), to *ammonitico rosso*-type deposits (the typical Tethyan red-colored, nodular, deep water facies; SCHLEDDING, 1985; GANDIN, 1987b; BECHSTÄDT *et alii*, 1988), that are interpreted as being on tectonically unstable shelves with highs and troughs (COCOZZA & GANDIN, 1990). The fauna of the Campo Pisano Formation is mainly trilobites, echinoderms, poriferans, brachiopods, and cancelloriids. Nevertheless, this fauna has been investigated very scantily because of its extremely poor accessibility. Only some trilobites, a few foraminiferans, poriferan spicules, and hadimopanellids (dermal plates of palaeoscolecoid worms) have been the subject of sporadic work (CHERCHI & SCHROEDER, 1985; MOSTLER, 1985; PILLOLA, 1991). Systematic studies on mollusks, trilobites, brachiopods and some small shelly fossils were done recently by ELICKI (2002), ELICKI *et alii* (2003), ELICKI & PILLOLA (2004), and MERGL & ELICKI (2004).

This paper presents for the first time a regionally-scaled detailed micropaleontological and paleoecological analysis of the Campo Pisano Formation. Based on extensive field work that involved the measurement and sampling of 19 stratigraphic sections, together

with the study of about 300 thin-sections, some chemical preparations, and sedimentologic and diagenetic observations, a paleoecologic reconstruction of the drowning of the Sardinian Perigondwanan platform is feasible and offered here.

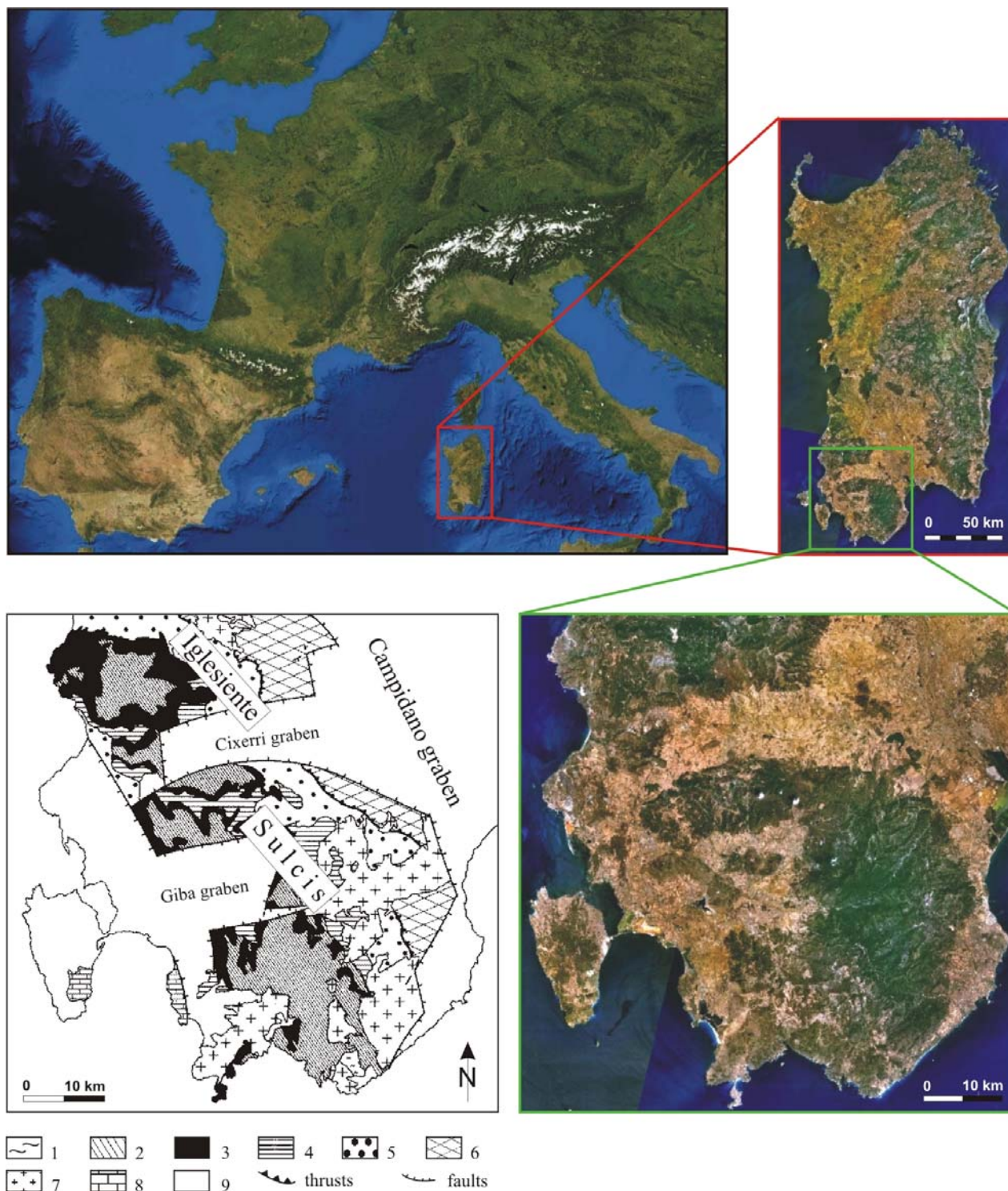


Figure 2: Location and simplified geological map of the work area in southwestern Sardinia. The two pictures at the bottom show the same segment and well correlation of main geological structures and those in satellite view (map modified after Bechstätt et alii, 1988; satellite photographs from NASA World Wind). 1 - Precambrian, 2 - Nebida Group (Early Cambrian), 3 - Gonnesa Group (Early Cambrian), 4 - Iglesias Group (late Early Cambrian to Early Ordovician), 5 - Early Ordovician to Devonian, 6 - allochthonous Paleozoic, 7 - Variscan granites, 8 - Mesozoic, 9 - Cenozoic.

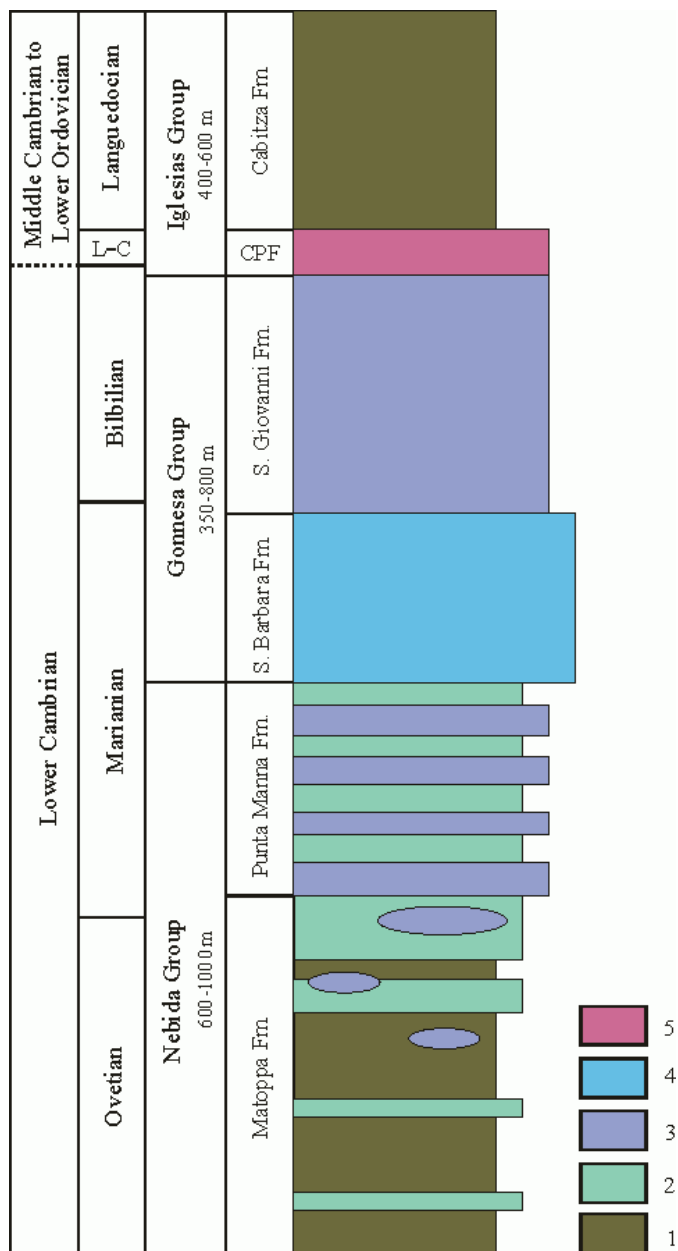


Figure 3: Stratigraphic succession of the Cambrian of SW Sardinia related to Spanish substages. 1 - claystone/siltstone, 2 - sandstone, 3 - limestone, 4 - microbial dolostone, 5 - nodular limestone, CPF - Campo Pisano Formation, L-C: Leonian and Caesaraugustan stages.

Geological framework

The Lower Paleozoic strata of southwestern Sardinia represent an autochthonous unit bordered to the east and north by Hercynian traps (Fig. 2). Prominent Cenozoic grabens separate this part of Sardinia from the remainder of the island and divide the area internally into several geographic regions. Locally, Hercynian granitoids have metamorphosed the Lower Paleozoic rocks.

Cambrian sediments are widely exposed in southwestern Sardinia (Fig. 2) and reach a thickness of 1500–2000 m (PILLOLA, 1991). Initial investigations were focused mainly on paleontology and biostratigraphy, but work has

now been done on sedimentology, litho- and biofacies, paleogeography and economic aspects (e.g., BORNEMANN, 1886, 1891; DEBRENNE, 1964; RASETTI, 1972; GANDIN, 1979; CHERCHI & SCHROEDER, 1984; MOSTLER, 1985; GANDIN, 1987a, 1987b; COCOZZA & GANDIN, 1990; PILLOLA, 1991; BECHSTÄDT & BONI, 1994; ELICKI, 2002; PILLOLA *et alii*, 2002; ELICKI *et alii*, 2003; ELICKI & PILLOLA, 2004; MERGL & ELICKI, 2004).

When the deposition of deposits of Cambrian age began in Sardinia is not precisely known: the only area where underlying rocks (Precambrian gneisses) are exposed is the southern part of SW Sardinia, but here the transition from gneisses to Cambrian sediments is represented by a nonconformity (JUNKER & SCHNEIDER, 1983). From bottom to top, the Cambrian succession is divided into the Nebida, Gonnese, and Iglesias groups (Fig. 3; PILLOLA, 1991). The middle to late Early Cambrian Nebida Group consists of clay-, silt- and sandstone in the lower part (Matoppa Formation), and a mixed carbonate-siliciclastic unit in the upper part (Punta Manna Formation). In southwesternmost Sardinia a discrete siliciclastic unit, the so-called "Bithia Formation", is described below the Nebida Group by JUNKER & SCHNEIDER (1983) and GANDIN *et alii* (1987). This placement was doubted by later authors who consider it as slightly heteropic facies of the lower Nebida Group (BECHSTÄDT & BONI, 1989). The Nebida Group is widely accepted as indicating a shallowing upward from a deep-water siliciclastic facies to a shallow mixed carbonate-siliciclastic ramp (e.g., BECHSTÄDT *et alii*, 1985; REUTER, 1987; BECHSTÄDT & BONI, 1994). Alternative interpretations initiate this evolution with a prodelta environment COCOZZA, 1979, GANDIN, 1979, and GANDIN *et alii*, 1987). The overlying Gonnese Group (late Early Cambrian) is characterized almost entirely by carbonate deposits with no siliciclastic influx; thus this group represents an isolated carbonate platform (*sensu* READ, 1985; e.g., BECHSTÄDT & BONI, 1994, and cited literature). Low in this group, higher salinity is denoted by the restricted, dolomitic, very shallow subtidal to intertidal laminites of the Santa Barbara Formation, overlain by shallow subtidal, open-marine limestones of a discrete and different facies (San Giovanni Formation). The youngest Cambrian unit is the Iglesias Group (latest Early Cambrian to Early Ordovician). At the base of this group are the limestones of the Campo Pisano Formation (at times nodular) deposited during platform drowning. It is succeeded by the deeper-water siliciclastics of the Cabitza Formation (Fig. 3). For detailed description of the origin and development of the early platform stages (Nebida and Gonnese groups), see the cited literature or the extended summary by BECHSTÄDT & BONI (1994), and the references cited therein.

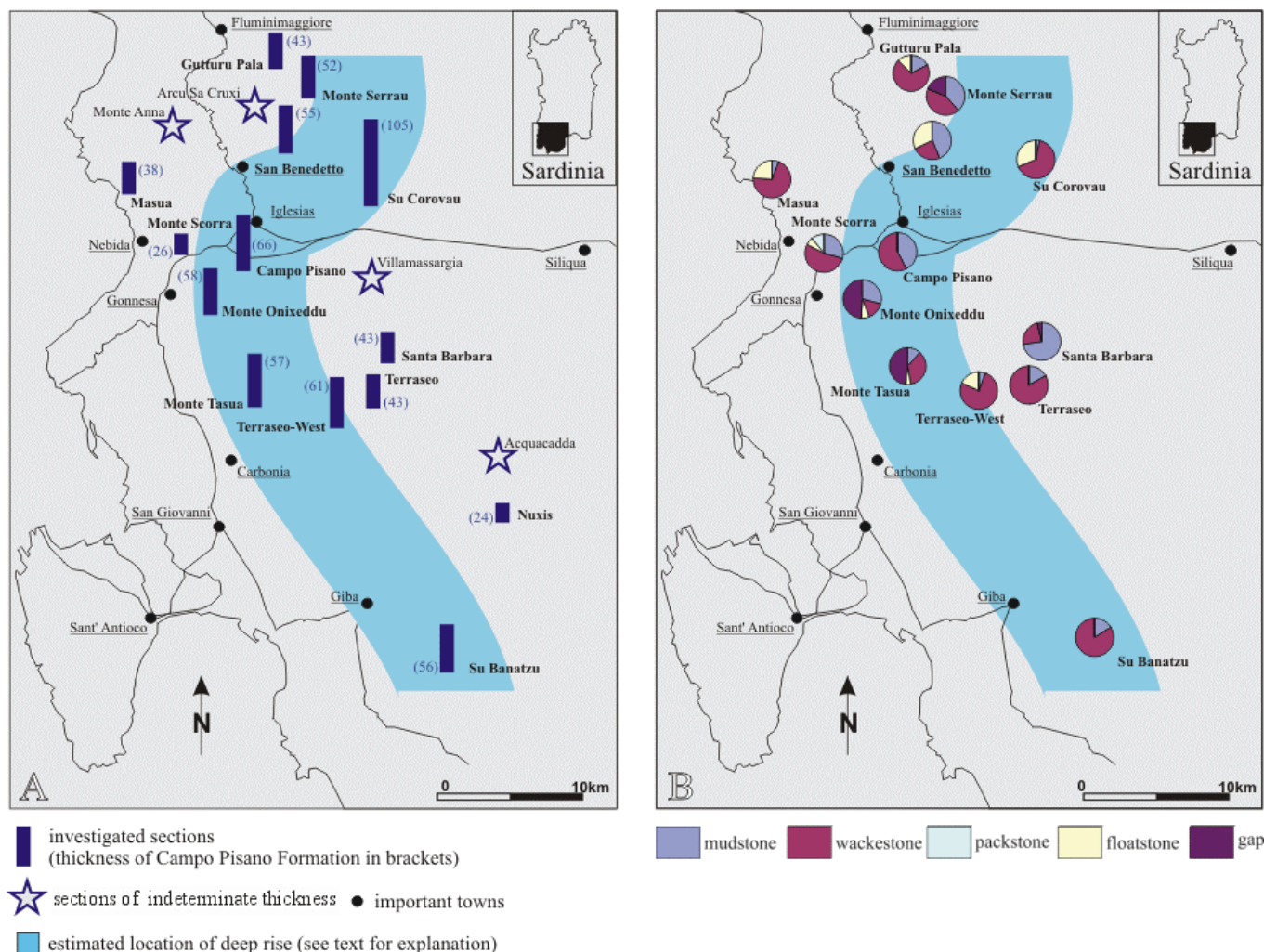


Figure 4: Plan of southwestern Sardinia showing the location of the measured sections of the Campo Pisano Formation, their thickness (A); and the relative amounts of the four carbonate lithotypes comprising them (B). The area occupied by suspected weak sea-floor high is in blue (see text for explanation). On the postulated high sections are thicker and the percentage of autochthonous carbonates greater, so the high is presumed to have favoured carbonate production.

The upper boundary of the Cambrian succession lies within the siliciclastic Cabitza Formation which continues to late Tremadocian times. The Cabitza Formation is truncated by a prominent erosional discordance caused by mid-Ordovician Sardinian tectonic movements on a regional scale (BECHSTÄDT *et alii*, 1994; LEONE *et alii*, 2002).

Lithology

The contact between the Campo Pisano Formation and the underlying massive limestones of the San Giovanni formation is abrupt, sharply defined (Fig. 3), and in many localities is marked by a breccia consisting mainly of San Giovanni clasts. But fragments from the basal Campo Pisano occur locally. (e.g., at San Benedetto, Mt. Marganai area, Mt. Tasua, Acquacadda, Terraseo, Fig. 4) (BONI *et alii*, 1981; SCHLEDDING, 1985; GANDIN, 1987a, 1987b). The Campo Pisano Formation is mainly wavy-bedded to nodular limestone with a variable clay content and a few quartz grains, although the siliciclastic content is greater locally (e.g., sections at Su Banatzu, Mt. Tasua,

Terraseo, Fig. 4) and continues farther East and South (Sulcis area), SCHLEDDING (1985). The most prominent features of the Campo Pisano Formation are: (1) the generally rich fossil content and (2) millimeter- to centimeter-thick bands of insoluble material, generated by diagenetic processes (GANDIN, 1987b; late diagenetic pressure dissolution: ELICKI, 2001). GANDIN (1979) and SCHLEDDING (1985) reported thin clay-rich and silty layers from the Campo Pisano Formation. The degree of diagenetic nodularity of the limestone varies both laterally and vertically. There is no regional or stratigraphic trend in the occurrence or intensity of nodularity, nor any correlation with a particular microfacies (see below).

The limestone of the Campo Pisano Formation is generally micrite-dominated. The most prominent lithofacies is light grey, yellowish, or reddish bioclastic wackestone (Figs. 4B, 5 & 6) with qualitatively and quantitatively distinct fossil content (see next section). Bioclastic floatstone is another



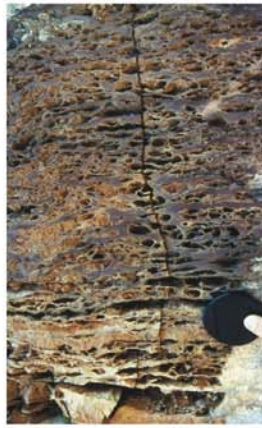
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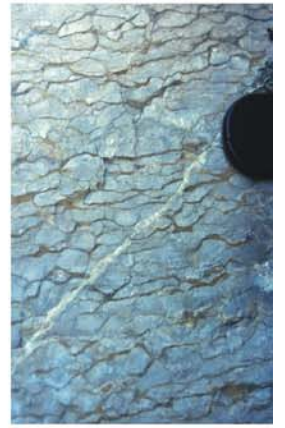
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◀ Figure 5:

1 – Field view of the San Benedetto section. Massive light limestones on the left represent the San Giovanni Fm (isolated platform, Gonnese Group); with sharp transition (red arrow), the Campo Pisano Fm follows (yellowish/reddish nodular banks of the drowned platform, thickness here about 45 m); to the right (yellow arrow) the Cabitza shales (deep siliciclastic cover of the platform; overgrown by vegetation) follow.

2 – Internal breccia at Arcu Sa Cruxi locality near San Benedetto. Note the big clasts imbedded in reddish matrix. Camera lid for scale.

3 – Macroscopic view of a typical Campo Pisano nodular limestone (Campo Pisano type section). Note the colour changes parallel to the bedding plane and the change in the degree of nodularity vertically. Hammer for scale.

4 & 5 – Nodular texture of the Campo Pisano Fm in mesoscopic view. If the carbonate is removed by erosion, only siliciclastic bands accumulated by diagenetic pressure solution remain (**4**, Gutturu Pala section); on fresh surfaces the carbonate nodules and stages of origin of pseudo-components are visible (**5**, Gutturu Pala section). Camera lid for scale.

6 – More or less isolated carbonate nodules enclosed in residual siliciclastic bands are rather rare, but, indicate the great role played by diagenetic pressure solution (Masua section). Hammer for scale.

7 – Wavy bedded to slightly nodular textures in the Campo Pisano Fm with nests of residual siliciclastics instead of bands (Gutturu Pala section). Scale bar is 10 cm.

8 – Massive texture of the Campo Pisano Fm indicated by a limestone bank (white arrows) within the wavy to slightly nodular succession at Monte Scorra section. Hammer for scale.

9 – "Shaly" texture of the Campo Pisano Fm at Su Banatzu section. Hammer for scale.

important lithofacies that occurs as intercalations at nearly all the sites investigated and in some sections occurs in significant quantities (e.g., Su Corovau, Mt. Serrau; Fig. 4). Fossil-bearing mudstone is intercalated in varying amounts (Fig. 4B). Bioclastic packstone was found as a small layer in only one section (Monte Scorra; Fig. 4B). More packstone occurrences were mentioned by SCHLEDDING (1985), GANDIN (1987b), BECHSTÄDT *et alii* (1988) and BECHSTÄDT & BONI (1994), but these authors did not report grainstone intercalations in the Campo Pisano Formation. Generally, wackestone and floatstone are more common in the western to northwestern areas ranging from the San Benedetto district to Su Corovau and Campo Pisano via Carbonia up to Su Banatzu, whereas mudstone is more abundant to the east and southeast (Fig. 4). Within the interval of transition to the overlying siliciclastic Cabitza Formation, limestones commonly alternate with thin greenish claystones.

Distinct (planar) bedding is not developed in the Campo Pisano Formation. Significant early compaction of the sediment was not observed. The fossil content is often random within the sediment (Figs. 6-1, 6-2 & 6-7), most probably, the result of shallow bioturbation (Fig. 6-7), though recognizable burrows were not seen.

The Campo Pisano Formation ranges in thickness from about 30 to 100 m, but most of the mapped sections are 40–60 m thick. From the "San Benedetto–Su Banatzu line" the thickness of the sections tends to decrease to the west–northwest as well as to the east–southeast (Fig. 4A). The occurrence of breccias at the base of the Campo Pisano Formation is reported from several sites by BONI *et alii* (1982), GANDIN (1987a, 1987b), and BECHSTÄDT *et alii* (1988). The distribution of these coarse clastics shows no distinct regional trend. SCHLEDDING (1985) and BECHSTÄDT & BONI (1989) described slumping and mass flow deposits from different locations in the area.

Fossils of the Campo Pisano Formation

Unlike the very pure limestone of the underlying San Giovanni Formation (isolated carbonate platform stage *sensu* READ, 1985) and the overlying siliciclastic Cabitza Formation (cover of the drowned platform), the deposits of the Campo Pisano Formation are characterized by a rich fauna of shelly fossils.

The most numerous are trilobites, brachiopods and echinoderms, with poriferans and cancelloriids locally common (Figs. 6-5, 6-6, 6-7, 6-8, 6-9, 6-12 & 7). The proportion of molluscs (pelagiellid gastropods), hyoliths (Fig. 6-13) and endolithic filamentous microbes is distinctly lower. Phosphatic small shelly fossils (palaeoscolecs, cambroclaves; Figs. 6-10 & 6-11) are present only sporadically in some beds of several sections.

The trilobite content is generally high but varies within sections (Fig. 7). The size of the identified trilobite remains ranges from a few millimeters up to one centimeter. Although most of them were recognized in thin sections (Fig. 6-7), some rocks yield silicified remains, so that dissolution to expose three-dimensional pseudomorphs made taxonomic investigation possible (Fig. 6-9) (ELICKI & PILLOLA, 2004). The chemical digestion procedure typically yields trilobite remains consisting mainly of disarticulated cephalons, pygidia, and a range of thoracic elements, but in some cases enrolled specimens and early ontogenetic protaspis and meraspis stages were obtained (ELICKI *et alii*, 2003; ELICKI & PILLOLA, 2004). Generally, agnostid and polymeran species are present together. Among agnostids, *Dawsonia bohemia* (Fig. 6-7) and *Condylopyge antiqua* are very common, whereas *Peronopsis* is rather rare. Polymerans are represented mainly by *Corynexochus* and less commonly by *Clavigellus*, *Dorypyge*, *Agraulos* and *Protolenus* cf. *pisidianus*. Taxonomic descriptions and discussions of this trilobite fauna are given in ELICKI & PILLOLA (2004).

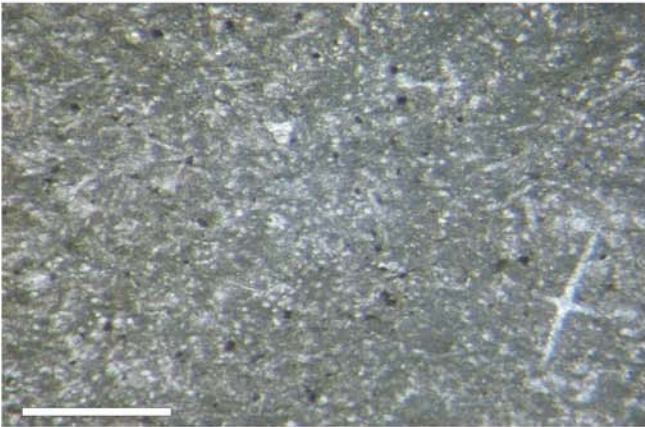
In large part brachiopods too are preserved as silicified pseudomorphs, but at one locality



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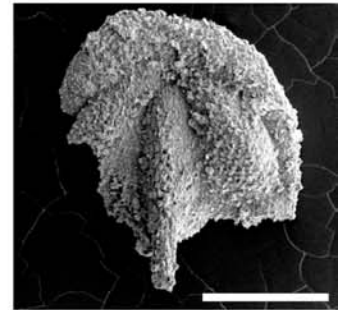
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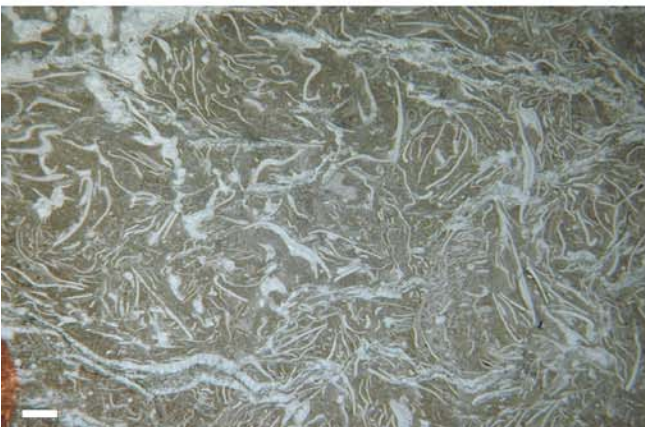
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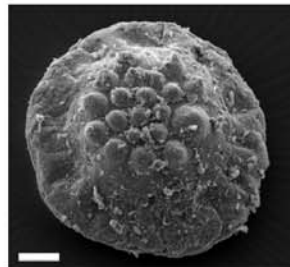
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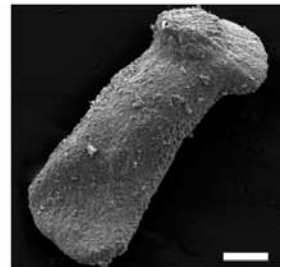
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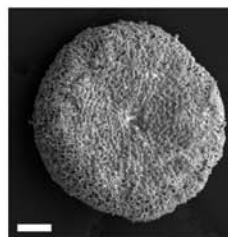
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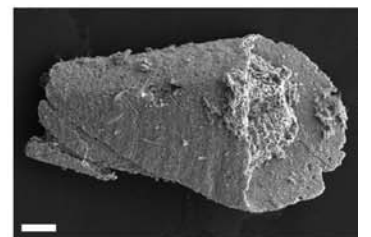
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◀ **Figure 6:**

1 - Origin of nodular texture by pressure solution (photomicrograph, Campo Pisano Fm, Monte Serravalle section). Note the snug fit of the pseudo-components. Scale bar is 1 mm.

2 - Photomicrograph of a siliciclastic band showing at the left a continuous lateral transition into limestone (the siliciclastic band seems to vanish). So, the band is not a primary sedimentary layer, but, a pressure solution suture. Nevertheless, the band can include carbonate relicts (pseudo-clasts). Masua section. Scale bar is 1 mm.

3 - Photomicrograph of platy texture in the Campo Pisano Fm described by previous authors as "calcareous slates" (Terraseo-West section). Scale bar is 1 mm.

4 - Photomicrograph of a rapid alternation of limestone and thin siliciclastic laminae typical of the transition from Campo Pisano Fm to the Cabitza Fm (Gutturu Pala section). Scale bar is 1 mm.

5 - Massive layers of sponge spicules (spiculite wackestone) common at the base of the Campo Pisano Fm (biofacies unit 1). Photomicrograph, Masua section. Scale bar is 1 mm.

6 - Echinoderm wackestone is characteristic for biofacies unit 2. Stereome preservation in the form of individual calcite crystals can be seen. Photomicrograph, San Benedetto section. Scale bar is 1 mm.

7 - Trilobites predominate in the upper portion of the sections (biofacies unit 3), sometimes they indicate bioturbation (see orientation of fossils). Trilobite wackestone. Photomicrograph, Gutturu Pala section. Scale bar is 1 mm.

8 - Sponge spicule from the Masua section (biofacies unit 1). SEM-micrograph. Scale bar is 0.1 mm.

9 - Silicified trilobite cephalon (*Dawsonia bohémica*) from biofacies unit 3 (Gutturu Pala section). SEM-micrograph. Scale bar is 1 mm.

10 - Phosphatic palaeoscolecid *Hadimopanella* sp. ("small shelly fossil") from biofacies unit 2 (Su Corovau section). SEM-micrograph. Scale bar is 0.03 mm.

11 - Phosphatic cambroclava ("small shelly fossil") from biofacies unit 2 (Su Corovau section). SEM-micrograph. Scale bar is 0.1 mm.

12 - Phosphatically preserved echinoderm ossicle from biofacies unit 2 (Su Corovau section). Note the stereome structure and the distinct lumen. SEM-micrograph. Scale bar is 0.1 mm.

13 - Hyolith *Microcornus* sp. from Su Corovau section (biofacies unit 2). SEM-micrograph. Scale bar is 0.1 mm.

(Gutturu Pala) there are phosphatic shells. Two groups of linguliformeans are recorded in great quantity: lingulides (*Lingulella*, *Schmidtites*) and acrotretides (*Opisthotreta*, *Vandalotreta*, *Linnarssonella*).

Echinoderm (Fig. 6-6) remains occur generally as disarticulated plates that often preserve the original netlike stereome structure. Phosphatized plates, segments and barrel-shaped elements are obtained by acid digestion (Fig. 6-12). The rarity of barrel-shaped elements is interpreted to indicate (1) a rather small proportion of stemmed forms and (2) that the stems were probably rather short. Much more widely distributed are polygonal thecal plates, commonly with sutural pores (epispines) for respiration (SPRINKLE, 1992).

Frequent annulate-segmented sclerites may represent brachiole or ambulacral parts. MOSTLER (1985) stated that most of the Campo Pisano Formation echinoderm remains represent eocrinoids, but a minor content of cincta (carpoideans) may be present too. Unfortunately, most were recognized in thin-sections so taxonomy is rendered difficult. But, epispines (typical in eocrinoids, cincta and some edrioasteroids) and the estimated ratios of sclerite morphologies: ([1] dominance of small probable brachiole/ambulacral elements, [2] many thecal plates, [3] rare stem segments, [4] a lack of typical stylocone segments *sensu* CLAUSEN & SMITH, 2005, and of [5] cinctan marginal plates and [6] axis channels *sensu* ZIEGLER, 1998), do indeed point to a probable dominance of eocrinoids. However, such a differentiation is not of key importance for basic paleoecological interpretations.

Disarticulated monaxons, tetractines, and pentactines are the most frequent sponge remains (Fig. 6-8). But, only the monaxons are occasionally abundant in discrete layers (Fig. 6-5). MOSTLER (1985) reported polyactinellid heteractinides from the Pisano Formation (*Phobetractina polymorpha*). Another heteractinide, *Eiffelia araniformis*, and the possible calcarean *Dodecaactinella triradiata* were identified.

Chancelloriids, a systematically problematic 'sponge-like' group, are lumped as coeloscleritophorans together with halkieriids, wiwaxiids, and siphonogonuchitids by BENGTSOHN & MISSARZHEVSKI (1981), but considered as a polyphyletic, non-poriferan group of unknown systematic affinity by MEHL-JANUSSEN (1999) and JANUSSEN *et alii* (2002). In the Sardinian samples most of the generally poorly preserved chancelloriids are represented by three- or four-rayed (*Allonnia*-type) and rosette-like (*Chancelloria*-type) sclerites. Many of the sclerites are single disarticulated rays. Identified forms include *Allonnia tripodophora*, *Chancelloria sardinica*, and *Archiasterella pentactina*. MOSTLER (1985) mentions *Chancelloria maroccana* and *Archiasterella* sp. as occurring in the Campo Pisano Formation.

Hyolithelminths (probable polychaetes), are nearly straight phosphatic open tubes with a more-or-less coiled proximal part and a round cross-section. The few specimens, up to 3 mm long, collected from near the base of the Campo Pisano Formation, are identified as *Hyolithellus* cf. *filiformis* (ELICKI *et alii*, 2003). A few specimens of the first known Sardinian bradoriid were found at the same stratigraphic level. Their subtriangular shape, phosphatic mineralogy, and other surface morphologic features identify them as *Hipponicharion ichnusum* (ELICKI & PILLOLA, 2004).

Molluscs are represented by pelagiellid and hyolith steinkerns with partially preserved

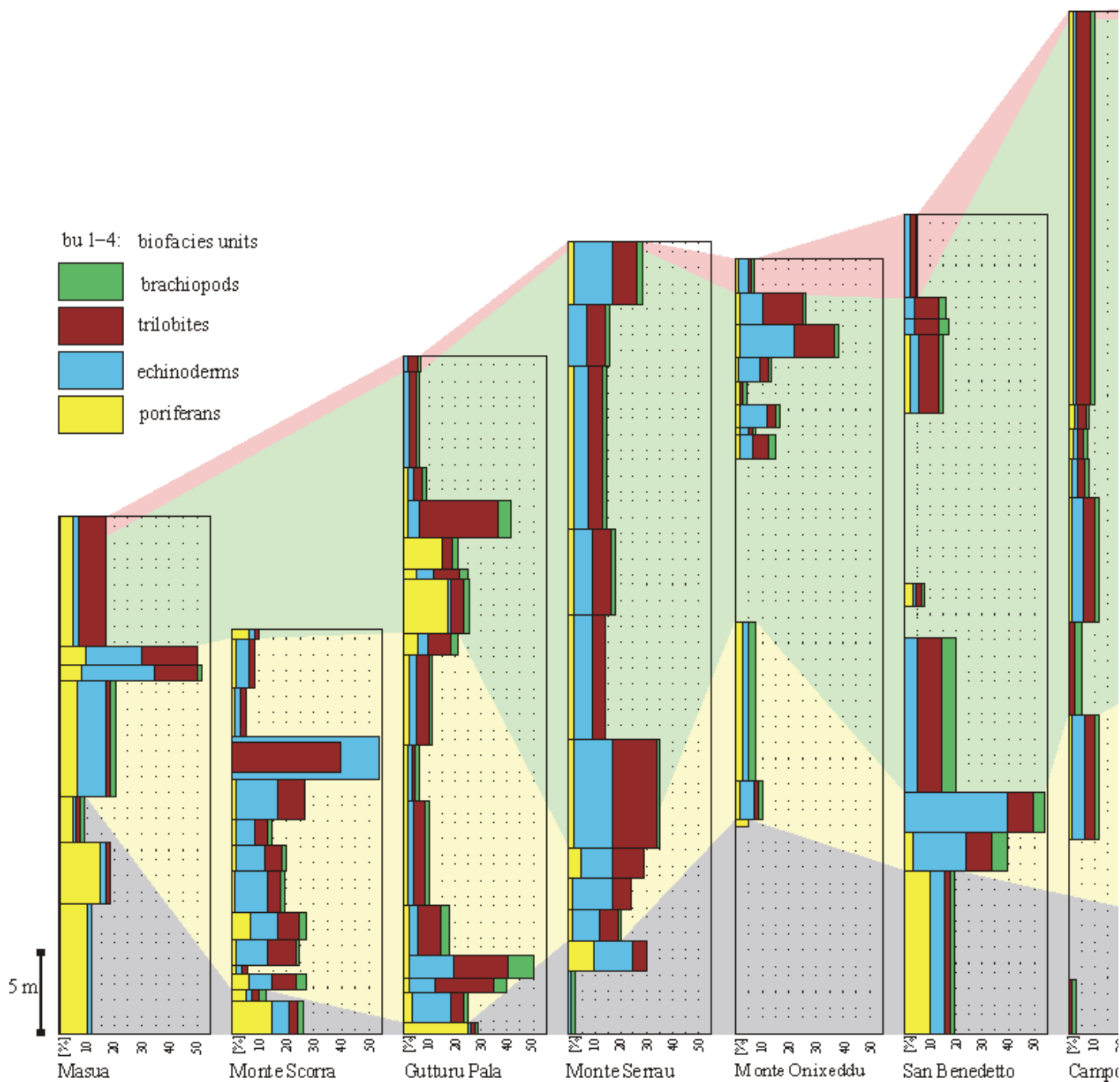
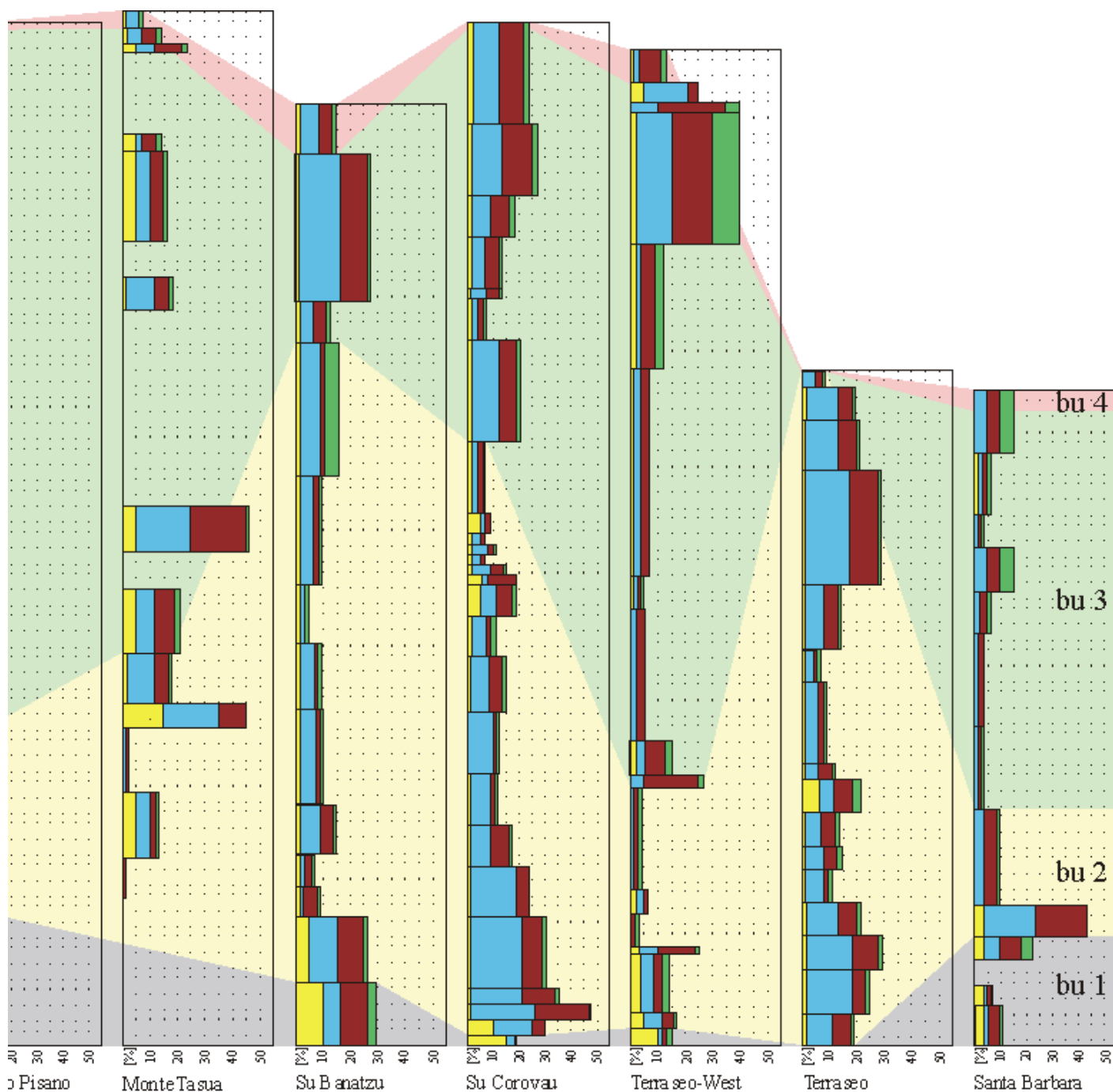


Figure 7: Fossil content (from thin sections) of stratigraphic sections of the Campo Pisano Formation arranged from West (Masua section) to East (Santa Barbara). Some sections shown on Fig. 4 are not included because of a stronger metamorphic overprint and/or local tectonics. Some tectonic events cause an interruption in the continuity of the fossil record. The units so affected are designated.

pseudomorph shells. Though the systematic affinity of pelagiellids is debated (GUBANOV & PEEL, 2000; PARKHAEV, 2001; LANDING *et alii*, 2002), the Sardinian specimens all belong to *Pelagiella subangulata* (for detailed description see ELICKI, 2002). Both hyolithomorphs and orthothecimorphs are present in the Campo Pisano Formation, but the only genus identified thus far is *Microcornus* (Fig. 6-13).

So called "small shelly fossils" (phosphatic microproblematica) are represented by hadimopanellids and cambroclaves. The

hadimopanellids (considered to be palaeoscolecids; HINZ *et alii*, 1990; MÜLLER & HINZ-SCHALLREUTER, 1993) are represented by very well preserved specimens of *Hadimopanella* (Fig. 6-10), abundant in discrete horizons of the lower portion of the Su Corovau section (Fig. 4). In addition, some poorly preserved sclerites are known from the Campo Pisano Formation type section (*Hadimopanella* cf. *oezgueli*; see CHERCHI & SCHROEDER, 1984). MOSTLER (1985) also mention the occurrence of hadimopanellids in the Gutturu Pala section (Fig. 4), but gives no detailed information or



figures. Cambroclaves (Fig. 6-11) are represented by minute sclerites of probable wormlike animals. They came from the same locality and levels as the hadimopanellids (ELICKI & WOTTE, 2003).

Microbes occur in the Campo Pisano Formation too. The endolithic cyanobacterium *Endoconchia angusta* was observed between the steinkerns and shell pseudomorphs of molluscs, and some single specimens of the possible oscillatoriacean *Girvanella problematica* are reported from the Campo Pisano Formation by CHERCHI & SCHROEDER (1984). The lack of other microbial fossils may be to the result of

taphonomic and diagenetic phenomena.

Biofacies and paleoecology

Taphonomic remarks

The microfauna of the Campo Pisano Formation is chaotically oriented (Fig. 6-7). There is no parallel "bedding", distinct layering, or size gradation. Even though no clear indication of burrows exists, the chaotic orientation of the biota in this succession, which was deposited at a low rate of sedimentation (LOI *et alii*, 1995), might be interpreted to be a result of bioturbation. Nevertheless, because of the lack of infaunal burrows or other deep traces and the absence of fecal pellets, a rather

shallow bioturbation near the sediment surface can be assumed. The lack of surface traces of bioturbation may have been caused by intensive bioerosion due to low sedimentation rates, or by late diagenetic pressure dissolution (ELICKI, 2001). Most of the fossils are preserved as calcite pseudomorphs.

Mechanical abrasion of the fossil remains was not seen. Even very fine and fragile structures on cambroclaves and microbrachiopods are perfectly preserved (Figs. 6-10 & 6-11). That portion of the fauna that had scleritomes covering or embedded in soft tissue is represented exclusively by disarticulated sclerites (poriferans, cancelloriids, echinoderms, etc.). Postmortem microbial decomposition of organic matter may have led to disarticulation *in situ* rather than as an accumulation caused by currents or slides. Too, trilobites in thin sections commonly show articulations (if the orientation of the section permits their recognition). These observations, together with the absence of lithoclasts and the near-total absence of coarse siliciclastics suggest that taphocoenosis was autochthonous. Cone-shaped fossils such as hyoliths occasionally show cone-in-cone stacking, thus indicating some bottom currents with low to moderate energy. Generally, bioclasts become fewer and smaller immediately below the transition to the overlying Cabitza Formation.

Biofacies units

There is no clear correlation between the occurrence of certain fossil groups and carbonate textural types (mudstone, floatstone, wackestone, packstone). Sections in the southeastern part of the study area tend to have fewer fossils than sections in the northwest. Generally, the various fossil groups show a similar pattern of stratigraphic distribution everywhere: at most localities fossil content is highest at the base of the Campo Pisano Formation. In nearly every section, the Campo Pisano Formation begins with a poriferan spiculite (locally as mass accumulations; e.g., at Su Banatzu, Terraseo West, Santa Barbara, Monte Scorra, Su Corovau, Monte Serrau; Figs. 4 & 7), whereas other fossil groups are clearly under-represented (rare brachiopods, small trilobites or echinoderms and single hyolithelminths and bradoriids), or totally absent. This remarkable initial biofacies is referred to here as 'biofacies unit 1' (bu1; Figs. 6-5, 7 & 8).

The overlying 'biofacies unit 2' (bu2; Figs. 7 & 8) is characterized by an abrupt onset of echinoderm remains (probably mostly eocrinoids, MOSTLER, 1985; Figs. 6-6 & 7) which are dominant here. Trilobites and brachiopods are conspicuous accompanying faunal elements while sponge spicules are of minor importance. New (in comparison to bu1), but minor elements of the biota are cancelloriids,

pelagiellids, and (rare) hyoliths. Of particular paleoecologic and biostratigraphic interest is the restriction in the occurrence of phosphatic small shelly fossils to this biofacies and stratigraphic level (Figs. 6-10 & 6-11).

Transition into 'biofacies unit 3' (bu3; Figs. 7 & 8) is gradual. Bu3 is characterized by about the same fossil groups as bu2, but trilobites dominate the fauna (Fig. 6-7). Echinoderms are also important constituents, and brachiopods, poriferans, hyoliths, and cancelloriids are occasionally present. This faunal composition continues upward to the base of the Cabitza Formation, but fewer and smaller fossils are present in the last few decimeters of the Campo Pisano Formation, designated here as 'biofacies unit 4' (bu4; Figs. 7 & 8).

All sections of the Campo Pisano Formation show more-or-less the same general trends of biofacies described above, albeit with local minor variations in fossil content and distribution (Fig. 7). In several regions, e.g., Su Banatzu, the Terraseo region, the Mt. Scorra-Masua region, and at Mt. Serrau the content of fossils increases in the upper levels of bu3 (Fig. 7). At Gutturu Pala, sponge spicules are prominent and reach a second maximum at these same levels (Fig. 7). Throughout the entire succession at Masua, poriferans are much more abundant than they are at other localities.

Paleoecological implications

The fossil groups of the Campo Pisano Formation occupied a variety of habitats and display a number of feeding strategies. Completely infaunal forms are represented by hadimopanellids (Fig. 6-10), usually interpreted as palaeoscolecids, for which a deposit-feeding strategy is assumed (e.g., WRONA & HAMDI, 2001). Whether or not some of the agnostid trilobites were also deposit feeders cannot be determined with certainty. The lack of eyes and small size alone do not validate such an interpretation. Unquestionable infaunal trace fossils have not been found in the formation. Semi-infaunal forms, represented by hyoliths with a rounded cross-section are found only rarely in the Campo Pisano Formation (ELICKI, 2000).

Most of the fauna in the Campo Pisano Formation is epibenthic. Sessile and semi-sessile epibenthic forms are represented by poriferans, cancelloriids, most of the echinoderms, and brachiopods (Figs. 6-8 & 6-12). For hyolithelminths a secondary epibenthic mode of life is probable. Whereas the filter-feeding sponges ingested mainly very small food particles and bacteria, the other sessile epibenthos lived on suspended detritus and microorganisms. The infaunal and sessile-epifaunal organisms noted above all suggest a soft firm- to mud ground habitat. There is no evidence for hardgrounds.

Mobile epibenthic biota are also common (most of the trilobites, some hyoliths, pelagiellids, bradoriids and some small shelly fossils such as cambroclaves; Figs. 6-9, 6-11 & 6-13). Some of the echinoderms too may have had only limited movement (cinctans, perhaps stylophorans, some edrioasteroids; SPRINKLE, 1992; FRIEDRICH, 1993; ZIEGLER, 1998; GUENSBURG & SPRINKLE, 2001; LEVEBVRE & FATKA, 2003; CLAUSEN & SMITH, 2005). A few of the epibenthos could have adopted a sporadic or nektobenthic life-style (some trilobites and bradoriids). All these groups are commonly viewed as low-level suspension feeders, but deposit feeding (e.g., some trilobites and possibly some carpoid echinoderms) or grazing (some molluscs) might have been feasible.

Direct indication of predation was not found. However, indirect evidence for this mode of

feeding may be provided by the cambroclaves, of which the skeletal elements have been interpreted as the protective external armor of a mobile epibenthic animal (CONWAY MORRIS & CHEN, 1991). Exclusively nektic elements of the Campo Pisano Formation fauna are unknown (even though some animals may have had such behavior during their early ontogeny). Whether or not some of the biota were scavengers is not apparent. So far, the only representative of the primary producers identified is *Girvanella* (oscillatorian cyanobacterium; CHERCHI & SCHROEDER, 1984).

The geographic (lateral) and stratigraphic (vertical) distribution of the biofacies units (bu1 to bu4) and the trophic and habitat characteristics of the fauna are fundamental to an interpretation of the depositional environment of the Campo Pisano Formation.

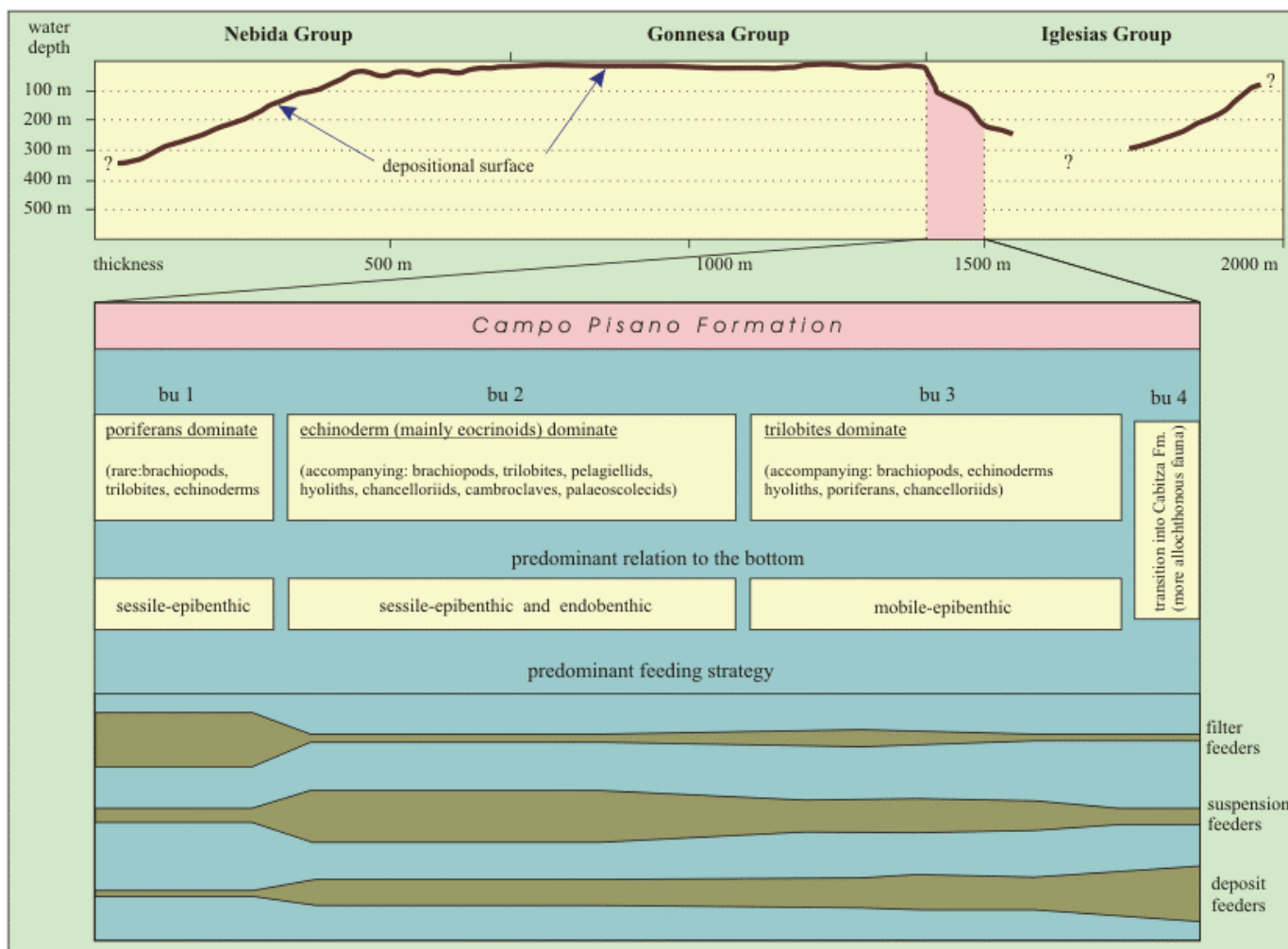


Figure 8: Bathymetry and thickness of the Sardinian Cambrian deposits and paleoecological succession (character of habitats and the dominant feeding strategy) in the Campo Pisano Formation that together indicate an intermittent drowning process.

To summarize: in the 19 sections investigated the basal unit of the Campo Pisano Formation is overwhelmingly dominated by sessile epibenthic filter feeders (spiculite biofacies, bu1; Figs. 7 & 8). Sessile and mobile suspension feeders are rarely present in bu1.

Upsection, a low-level suspension-feeder fauna becomes important (echinoderm biofacies, bu2; Figs. 7 & 8). The dominant fossils of bu2 are sessile forms (most echinoderms, brachiopods), but mobile elements are also common (trilobites). The proportion of poriferans

decreases significantly. In the accompanying fauna, distinctive elements include a rare infaunal biota (hadimopanellids) and other fossils, which suggest adaptations for protection against predators (cambroclaves). The upper Campo Pisano Formation (bu3; Figs. 7 & 8) is characterized by mobile epibenthic suspension feeders and by an increase of probable nektonic or planktic elements (greater proportion of agnostids). The proportion of sessile organisms decreases significantly. In the uppermost Campo Pisano Formation (bu4) an increase of transported elements (indicated by small broken bioclasts; Figs. 7 & 8) is interpreted here as indicative of a more pelagic origin. Figure 8 depicts this paleoecological succession of biofacies along with habitat characteristics, feeding strategies and bathymetry.

Biostratigraphy

The age of the Campo Pisano Formation was determined primarily by trilobites, but, because of their scarcity, the stratigraphic resolution obtained is low.

The assignment of the trilobites from the Campo Pisano Formation to the Middle Cambrian (RASETTI, 1972) was refined by PILLOLA (1991) and LOI *et alii* (1995) who identified two stratigraphically sequential assemblages. The oldest trilobite fauna of the Campo Pisano Formation (CP1 fauna of LOI *et alii*, 1995), located several meters above the base of the formation, includes *Protolenus* (*Protolenus*) cf. *pisidianus* that indicates the Early to Middle Cambrian boundary interval (for discussion see LOI *et alii*, 1995). The base of the Campo Pisano Formation therefore is in the uppermost Early Cambrian in the Mediterranean sense (for discussion of the problematic interregional correlation of the Early–Middle Cambrian boundary see GEYER & LANDING, 2004), whereas the immediately overlying portion of the formation (early CP2 fauna of LOI *et alii*, 1995) correlates with the *Paradoxides mureoensis* Zone (Leonian, early Middle Cambrian; early to middle Agdzian of the western Gondwana scale of GEYER & LANDING, 2004; Fig. 3). The middle portion of the Campo Pisano Formation (middle CP2 fauna of LOI *et alii*, 1995) yields *Calodiscus foveolatus*, some oryctocephalids, dorypygids and others and corresponds to the *Badulesia* Substage of the mid-Middle Cambrian (early Caesaraugustan; LOI *et alii*, 1995; Fig. 3). The upper part of the Campo Pisano Formation (late CP2 fauna of LOI *et alii*, 1995) contains, among others trilobites, *Pardailhania hispida* and *Ptychagnostus* sp., relating to the *Pardailhania* to basal *Solenopleuropsis* Substages (middle Caesaraugustan) of the mid- Middle Cambrian. The transition from the Campo Pisano Formation to the Cabitza Formation should thus be in the late Caesaraugustan (late *Solenopleuropsis* Substage of the late Middle Cambrian; Fig. 3).

Non-trilobite age determinations are

consistent with the trilobite biostratigraphy. Lingulid and acrotretid brachiopods from the upper Campo Pisano Formation (especially *Linnarssonella* sp.) imply a late Middle Cambrian age (MERGL & ELICKI, 2004). Poriferans, chancelloriids, hyoliths (*Microcornus*), pelagiellids, and small shelly fossils (e.g., *Hipponicharion*) are in accordance with a late Early to Middle Cambrian age (BENGTSON *et alii*, 1990; PARKHAEV, 2001; ELICKI, 2002; GOZALO & HINZ-SCHALLREUTER, 2002; ELICKI *et alii*, 2003).

Precise correlation of sections in the study area is hindered by a lack of trilobite material. However, on a regional scale this problem can be somewhat mitigated through the use of ecostratigraphic data. For example, in the Campo Pisano type section, the abrupt appearance of 'biofacies unit 2' (bu2) is the equivalent of the top of the *Protolenus* (*Protolenus*) cf. *pisidianus* level. This distinct change is interpreted as a regional environmental reorganization. For the study area, this means that the base of bu2 coincides with the base of the Middle Cambrian (in the Mediterranean sense) and that bu1 is latest Early Cambrian in age. So the bu1–bu2 transition represents an ecostratigraphic level permitting recognition of the Early–Middle Cambrian boundary in southwestern Sardinia. Using this model, a correlation of the basal portions of most of the sections is possible although trilobites are absent. The validity of this technique has been confirmed recently by finding the *Protolenus* (*Protolenus*) cf. *pisidianus* index taxon in the Su Corovau section (PILLOLA, pers. comm., 2004; Fig. 4) at precisely the level predicted by ecostratigraphy.

Paleobiogeography

The Campo Pisano trilobites have a western Gondwanan affinity (Fig. 1). Geographical relationships exist with southern France (Montagne Noire), northern Spain and southern Turkey, *i.e.* the Mediterranean region of the so-called "Atlantic Faunal Province" *sensu* SDZUY (1972) and PILLOLA (1994). Associations with the northern region of the "Atlantic Faunal Province" seem to be more distant. Biogeographic relations with the Far East as interpreted by SDZUY (1972), PILLOLA (1991) and ÁLVARO *et alii* (2003) for some Early Cambrian faunas have not yet been found among the trilobites of the Campo Pisano Formation. But, some small shelly fossils (cambroclaves) may indicate biogeographic relationships with the Asiatic realm. Until recently, cambroclaves were known only from Kazakhstan, China, and Australia. But, the latest finds from the late Early Cambrian of Germany (ELICKI, 1994) and early Middle Cambrian of Sardinia (ELICKI & WOTTE, 2003), both on the western Gondwana shelf, represent the oldest and youngest representatives of this group known thus far and may have some paleogeographic

significance. The mollusc *Pelagiella subangulata* shows a similar paleobiogeographic distribution along this "Palaeotethyan belt" (BRASIER, 1989; PARCHAEV, 2001; ELICKI, 2002). Small shelly fossils offer further support for close connections between the Mediterranean Cambrian areas: e.g., the bradoriid *Hipponicharion* is a good index taxon for the

western Gondwanan late Early and early Middle Cambrian beds of Germany, Spain, Morocco, and Sardinia (ELICKI, 1994; GOZALO & HINZ-SCHALLREUTER, 2002). In summary, a good faunal exchange among and connections between the western Gondwanan paleogeographic regions is clearly established.

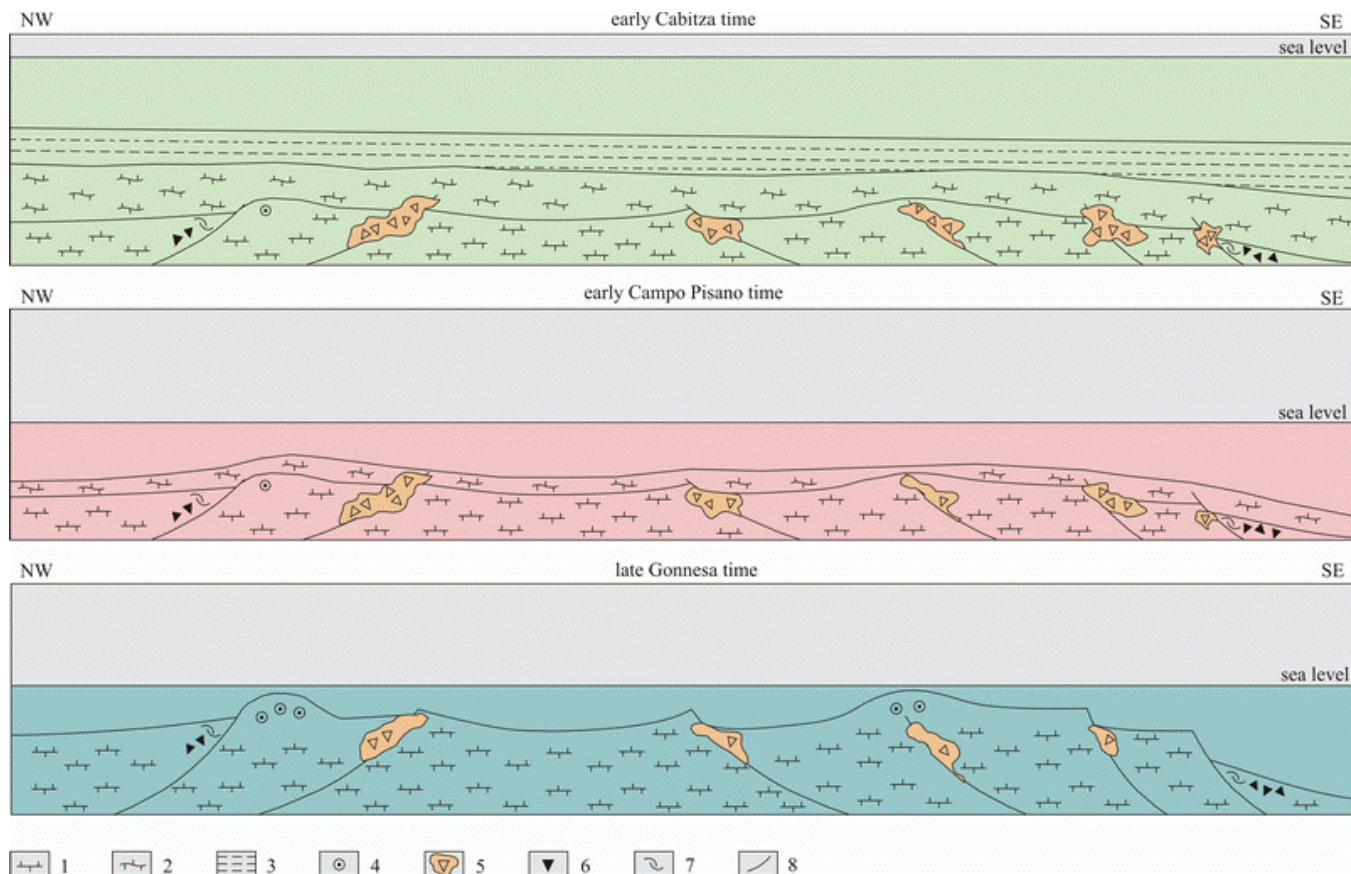


Figure 9: Interpretation of development stages at different time slices: During late Gonnesa time, the carbonate platform was already segmented by tensional tectonics (origin of internal breccias) resulting in a significant bottom topography; at steep flanks mass flows occurred. Early in Campo Pisano times the sea level rose and tensional-related faults were still active locally. This led to the continued development of internal breccias and bottom topography was only moderately accidented. Early in Cabitza times the rise in sea level continued and deeper marine clastics covered the drowned carbonate series. 1 - San Giovanni Formation, 2 - Campo Pisano Formation, 3 - Cabitza Formation, 4 - oolites, 5 - internal breccia, 6 - external breccia, 7 - mass flow, 8 - fault.

Discussion

Distribution of facies patterns

Microfacies analysis of the Campo Pisano Formation demonstrates that many macroscopic features, e.g., color and nodularity, are of little value in such studies. After extensive field and microfacies investigations, the following characteristics were found to be essential for environmental interpretation: (1) microfacies, (2) geographic and vertical distribution of biota, (3) taxonomic variety, (4) non-biogenous components, (5) ichnology, (6) regional variation in stratigraphic thickness, and (7) the characteristics of underlying and overlying strata and of the formational limits.

Within the study area, the thickness of the Campo Pisano Formation increases slightly from southeast to northwest as far as the San

Benedetto to Su Banatzu line (Figs. 4 & 7). Thickness decreases northwest of this line. Because deposition was almost entirely autochthonous (see above), this region is assumed to be a carbonate depocenter - a weakly elevated, low-energy area - where conditions for carbonate precipitation were favorable (Fig. 4). The range in number and quantity of the autochthonous biota shows a similar distribution (Fig. 7), suggesting that this area was favorable to life and hence a preferred habitat. This view agrees well with the inference that it was a weak high. For a part of the Early Cambrian, a similar, but high-energy shallow area was proposed by BECHSTÄDT *et alii* (1985), but it was sited some 5-10 km to the west, coinciding approximately with the present southwestern Sardinian coastline. The weak rise during the Middle Cambrian may represent a

later stage of this topographic elevation, relocated by a rise in relative sea-level (see below).

The abrupt arrival of the silty Campo Pisano Formation over the shallow, isolated carbonate platform of the Gonnese Group (Figs. 3 & 5-1) is indicated by its initial spiculite biofacies (bu1; Figs. 6-5 & 7) in nearly every section, sometimes as a gradual transition, in other places associated with breccias (see GANDIN, 1987b; and the discussion of these breccias below). The sudden influx of siliciclastic material into the depositional area with the arrival of the Campo Pisano Formation is best interpreted as the aftermath of the completion of the fill of a sediment-trapping basin between an erosional area and the isolated carbonate platform. This is the interpretation of BECHSTÄDT *et alii* (1985) ("Eastern Sulcis basin"), that accounts for fine siliciclastic sediment reaching a formerly isolated depositional area. Suspension currents were presumably the primary agent for the transport of the very fine-grained siliciclastics of the Campo Pisano Formation. Infrequent storms may have occurred, but the hummocky cross-bedding, allochthonous shell layers, reworked and transported fauna, and extraclasts commonly produced by storms were not observed. These facts support the suggestion by previous authors that extensional tectonics combined with a relative rise in sea-level affected the area and led to the "drowning" of the platform (GANDIN, 1987a, 1987b; BECHSTÄDT *et alii*, 1988; BECHSTÄDT & BONI, 1989; COCOZZA & GANDIN, 1990). The brief dominance of filter feeders (bu1; Figs. 7 & 8) at the base of the Campo Pisano Formation reflects the successful adaptation of one group of organisms to the newly increased depth and to the sudden rise in the quantity of suspended material along with more influential water currents, and in addition the advantages gained by a rapid adaptation to new ecological niches.

The transition from bu1 (spiculite biofacies) to bu2 (echinoderm biofacies) is a conspicuous change of milieu in the lower Campo Pisano Formation affecting the entire depositional area (Figs. 7 & 8). It is interpreted as reflecting an increase in water depth. The persistence and strong dominance of sessile epibenthic suspension feeders in this biofacies (Fig. 8) suggest a permanent increase in the influx of suspended nutrients. Reasons for this change might include a reorganization in current patterns or a wetter climate that would intensify chemical erosion and sediment transport in the hinterland and thus an increase in the input of siliciclastic sediment. These changes may have been caused by a shift of this region to a higher paleogeographic latitude (COURJAULT-RADÉ *et alii*, 1992; MCKERROW *et alii*, 1992; SESLAVINSKY & MAIDANSKAYA, 2001).

The next change in the biofacies of the

stratigraphic succession, from bu2 to bu3 (Figs. 7 & 8), took place without a conspicuous change in lithology; trilobites (especially agnostids) became more numerous, while the sessile epibenthos declined (Fig. 8). This change is best attributed to an ongoing, steady rise in relative sea-level.

Near the lithological transition to the overlying siliciclastic Cabitza Formation (Fig. 7), the environment changed again. At this level the fossil content as a whole decreases significantly and among the trilobites and brachiopods, only small, mostly broken parts remain (bu4); this fragmentation is assumed to have been caused by the action of currents that transported the fossils. The accompanying lithological change is interpreted as evidence of a second distinct eustatic pulse that caused the depositional environment to deepen (Fig. 8). However, it is improbable that the depth of water exceeded that of an outer shelf because of the general persistence of similar lithofacies in overlying strata. The existence of deeper conditions may not be taken as clearly expressed lithologically until a fully siliciclastic regime was established in the Cabitza Formation. LOI *et alii* (1995) have demonstrated that younger portions of the Cabitza Formation were deposited in relatively shallow waters near storm wave base. This supports the view that the uppermost Campo Pisano and the lower Cabitza Formations were not much deeper than the overlying strata, because extreme fluctuations in bathymetry in such a short time are unrealistic. Hence, the rather rapid lithological change from the carbonates of the Campo Pisano Formation to the siliciclastic sediment of the Cabitza Formation probably does not indicate a dramatic increase in water depth, but rather reflects a drift to a higher latitude (cooling and increased precipitation).

Morphology and bathymetry of the depositional area

The interpretation of the depositional environments of the Campo Pisano Formation used in previous models was based on facies and sedimentological studies (GANDIN, 1979, 1987a, 1987b; BECHSTÄDT *et alii*, 1988; COCOZZA & GANDIN, 1990; and others). The resulting model for the Campo Pisano limestone was a type of nodular deep water facies ("*ammonitico rosso*" lithofacies) deposited on pelagic topographic highs on a broken, drowned platform, while troughs between these highs would serve as areas of deposition for calcareous clays and silt. Nevertheless, judging from the microfacies and paleoecological results presented here, this model must be modified. In the 19 studied sections (Fig. 4A), nodular, wavy-bedded, silty limestones, and massive carbonate banks alternate repeatedly and irregularly (Figs. 5-3, 5-4, 5-5, 5-6, 5-7, 5-8 & 5-9). Closely adjacent sections do not show the same patterns of vertical distribution for these

lithotypes, so that their sequence cannot be used for correlation. Furthermore, biofacies do not correlate with these lithofacies. In the discrete more or less widely separated sections although there is no difference in the sequential position of the zones established by the autochthonous fossils, this fossil zonation has no relationship to the lithology of the approximately corresponding stratigraphic levels: that is whether or not the same fossils (qualitatively and quantitatively) are enclosed in nodular, wavy-bedded, silty or massive limestones. The origin of these textural types is related to diagenetic processes (GANDIN, 1987b; ELICKI, 2001) that have accentuated minor differences in the lithology of the primary sediment (e.g., siliciclastic content, permeability), and early cementation; the direction and velocity of minor currents may also be involved. The relative uniformity of biofacies over the entire area of study demonstrates that during deposition of the Campo Pisano Formation bottom conditions that could have caused ecological differences were almost completely absent. However, a slight difference in bottom topography may have led to small regional differences in siliciclastic content (an increase to the East) as reported by BECHSTÄDT *et alii* (1988). Semi-quantitative analysis of sections from several localities found quartz content to be below 2 % that HAMANN (2003) regarded as probably aeolian. Nevertheless, this weak eastward increase in siliciclastic content in the Campo Pisano limestones may support the interpretation of BECHSTÄDT *et alii* (1988), that a hinterland lay to the East, in sharp contrast to the views of GANDIN *et alii* (1987), who postulates a basin that direction.

Breccias

Breccias exist in the beds marking the changeover from the Gonnesea platform to the Campo Pisano offshore carbonates (BONI *et alii*, 1981; GANDIN, 1987a, 1987b; BECHSTÄDT *et alii*, 1988). These breccias (Fig. 5-2) have usually been interpreted as due to tensional tectonics and/or karstification. GANDIN (1987b) described two discrete types: (a) breccias consisting of well sorted clasts (up to 20 cm), and (b) unsorted megabreccias. The first type (a – Mt. Agruxiau type) GANDIN interpreted as karst-related with a red hematitic and/or marly matrix sometimes yielding Campo Pisano fossils. The second (b – Mt. San Giovanni type) was called a collapse breccia with a marly matrix, and lacking Campo Pisano Formation clasts. A karstic origin for these breccias was doubted by BECHSTÄDT *et alii* (1988). For some authors these breccias indicate a profound relief on the sea floor (CARANNANTE *et alii*, 1984; SCHLEDDING, 1985; GANDIN, 1987b; BECHSTÄDT *et alii*, 1988; COCOZZA & GANDIN, 1990). However, significant relief with highs and lows would promote distinct differentiation in the biofacies of the autochthonous fauna. But such

differentiation does not exist. Biotic assemblages occur at the same stratigraphic levels over the whole area, and there are no marked lateral differences in their composition. The characteristics of the breccias mentioned above (clast content and shape, matrix, thickness, stratigraphic position) are easily explained as typical of fault-related internal brecciation (Fig. 9). Breccias of this origin are typically monomictic (or nearly monomictic) and part of their marl matrix may be derived from overlying strata, so can contain fossils from overlying beds injected with the marls into the still unconsolidated internal breccias below (FÜCHTBAUER, 1988). Breccias of this kind originate beneath a sedimentary cover that is under a tensional tectonic regime (FÜCHTBAUER & RICHTER, 1983; FÜCHTBAUER, 1988). ARRIBAS *et alii* (1990) observed transitions from the Sardinian breccias in the upper levels of the Gonnesea platform into unbrecciated rocks below, a phenomenon characteristic of such an origin (FÜCHTBAUER, 1988). ARRIBAS *et alii* (1990) were the first to interpret the breccias at the Gonnesea–Campo Pisano transition as flexure-related rocks caused by tension. A reinterpretation of lithological features, taking into consideration regional tectonics (e.g., GANDIN, 1987b; BECHSTÄDT & BONI, 1989; COCOZZA & GANDIN, 1990), and paleoecologic data presented here indicate that when these breccias formed the morphologic gradient was rather low (in contrast to the gradient inferred for other types of breccias and debris flows that developed on the platform slopes during older Gonnesea time, BECHSTÄDT *et alii*, 1988). If the gradient was low, a horst-and-graben environment for the breccias of Campo Pisano time cannot be sustained (Fig. 9).

There are two opinions regarding the depth of water under which the Campo Pisano Formation was deposited. The fossil content has been interpreted as a neritic shelf fauna, but the nodular texture of the carbonates has been thought to indicate depth up to several hundreds of meters (e.g., MOSTLER, 1985; SCHLEDDING, 1985). Shallower conditions were assumed by COCOZZA (1979, "shallow water"). The Campo Pisano carbonates are typical Cambrian offshore sediments, deposited in the photic zone below storm wave base (Figs. 8 & 9). No indicators of wave and tidal activity, erosion, transport of clasts from shallower areas, or gradation, or hummocky bedding was observed. Rarely, thin allochthonous shell layers occur (ÁLVARO, GANDIN, pers. comm.); possibly they are indicative of storms. All these features are typical in distal ramp or shelf environments, which, in the Cambrian are usually characterized paleoecologically by a dominance of echinoderms and sponges and by intensive bio-retexturing (READING, 2002). The source of carbonate mud in such environments could be (1) adjacent shallow platforms (if allochthonous), or (2) biotic precipitation by

planktic microbial organisms (if autochthonous) or (3) a mixture of both (HANDFORD, 1986; AURELL *et alii*, 1995; READING, 2002). An autochthonous microbial origin is favoured for the Campo Pisano carbonates. No shallow platform remnants "able to deliver carbonate ooze are known from that time" BECHSTÄDT & BONI, 1989). Furthermore, the high mud content points to a rather low slope on the sea floor, a conclusion compatible with the interpretation from fossils of a shelf/ramp environment.

Although "neritic" conditions are strongly indicated by the fauna, by the lithology (LOI *et alii*, 1995), and by the context of syntectonic evolution (BECHSTÄDT & BONI, 1989; GANDIN *et alii*, 1987), absolute water depths are hard to calculate because of limited knowledge concerning rates of subsidence, sediment budgets and the range of rises in sea-level. Nevertheless, because of (1) the shallow subtidal to intertidal conditions indicated for the underlying upper Gonnese Group (BECHSTÄDT *et alii*, 1985; GANDIN *et alii*, 1987; FRÖHLER, 1994), (2) the conclusions of COCOZZA & GANDIN (1990) and LOI *et alii* (1995), that the overlying lower Cabitza Formation indicates a deeper shelf, and (3) the two regional well-defined paleoecologic changes in the Campo Pisano Formation indicate a two-step deepening of the whole area during Campo Pisano time (Fig. 8). Consequently, it is with considerable confidence that the siliciclastic lower Cabitza Formation is assigned a depth below the photic zone and the Campo Pisano Formation itself is considered to have recorded an intermittent deepening of the sea within the photic zone (Figs. 8 & 9).

Conclusions

The late Early to late Middle Cambrian Campo Pisano Formation of southwestern Sardinia is represented by a succession of nodular, wavy-bedded, silty, and massive offshore limestones (Fig. 5) deposited during the drowning of the isolated Sardinian Early Cambrian carbonate platform. After a distinct rise in relative sea-level that caused the isolated carbonate platform conditions of the underlying Gonnese Group to end (Fig. 9), but with water-depth still relatively shallow, the initial Campo Pisano fauna developed with epibenthic filter feeders (sponges) dominant and with a moderate amount of suspended input from the hinterland (Fig. 7). This fossil community predominated in the lower beds of the Campo Pisano Formation in nearly all of the 19 sections mapped.

Higher in the succession poriferans persist as an important element of the fauna only in some westernmost areas. Upsection, echinoderms come to dominate the biofacies (mainly sessile epibenthic eocrinoids and other low-level echinoderm suspension feeders) and they in turn are succeeded by a trilobite-

dominated biofacies in which the mobile epibenthos becomes increasingly important upward as the depth of water increases (Figs. 7 & 8). Near the boundary with the deeper water Cabitza Formation an increase in the strength of currents is suggested by the fact that the fossils are all fragmentary and have been transported. This trend to less proximal and deeper-water sedimentation is apparent over the entire working area for it is expressed in both stratigraphy and paleoecology. Regional paleoecologic trends (Figs. 4 & 7) are areally extensive and not pronounced, so they are thought to have been caused by a rise in sea level rather than by tensional block-tectonics and fracturing on the sea floor. The data are interpreted as indicating the existence of a weak rise (low-energy swell) trending SE-NW that served as a carbonate depocenter and is thought to have been a preferred habitat for the Campo Pisano biota (Fig. 4B). There is no evidence of a high-energy shallow area anywhere. The breccias are classed as "internal" that is formed below a cover of sediments while tensional tectonics were active (Fig. 9).

The biofacies and the sedimentary criteria: thickness, mud-, fossil-, and clastic content indicate that the area gradually drowned as one unit; it was undifferentiated bathymetrically (Figs. 8 & 9). It is classed as a neritic open-marine shelf or ramp. An intermittent rise of relative sea-level, accompanied by a tensional tectonic regime, along with a move into higher paleogeographic latitudes, led to a regional uniformity in facies. The sudden rise of relative sea-level at the transition from the Gonnese platform to the Campo Pisano offshore carbonates caused the entire area to be under about the same depth of water (Fig. 8). Great morphological differences within the area, caused by tectonism during the isolated platform stage (Gonnese Group), are interpreted here as having been leveled by Campo Pisano times through relief-filling redeposition during late Gonnese times and by offshore sedimentation during early Campo Pisano times (Fig. 9). So the model presented here resolves seeming contradictions between sedimentologic and paleontologic criteria.

The techniques of micropaleoecological analysis presented here are a very useful tool for modeling the configuration and evolution of Perigondwanan depositional areas during the Cambrian. The method can be successfully applied to the reconstruction of regional geological processes for that time, in particular where biostratigraphic data and sedimentary manifestations are fragmentary or unavailable.

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