INTRODUCTION

Supply-side ecology (e.g., Gaines and Roughgarden, 1985) stressed the importance of organism substitution for the persistence of marine benthic communities. Individuals die and are replaced, so that the future of a community depends on the success of larval settlement. In this framework, larval ecology (usually a plankton-based affair) becomes crucial in benthic ecology and life cycles acquire a central position in marine ecology (Boero et al., 1996; Marcus and Boero, 1998).

The efficacy of dispersal of benthic organisms is linked to propagule vagility, with the possibility of delaying settlement until proper substrates become available. As remarked by Vance (1973), the planktonic larvae of benthic invertebrates can be either lecithotrophic (with their own reserves) or planktotrophic (feeding on external food sources), this affecting their possibilities of dispersal and survival. Scheltema (1966, 1988) described long-lived larvae of benthic invertebrates able to cross oceans: teleplanic larvae. A severe constraint to such wide-range dispersal is the impossibility, for non-feeding larvae like those of most hydroidomedusae, to have the necessary reserve to lead a long larval life before settlement (see Cornelius, 1992).

Boero and Bouillon (1993) and Boero et al. (1997) reviewed the variety of hydroidomedusan cycles, describing a vast array of modifications of the classical polyp-medusa alternation: in many cases, either the medusa or the polyp can be suppressed. The primary larva in the hydroidomedusan life cycle is the planula, a stage with generally low vagility which, in some cases, even remains connected to the mother colony by mucous threads.
before settlement (Wasserthal and Wasserthal, 1973; Hughes, 1977). There are no records of feeding planulae and their settlement should occur soon after release, this limiting dispersal for species deprived of other planktonic stages, i.e. medusae. Boero and Bouillon (1993), however, remarked that the presence of a medusa in the life cycle is not correlated to wide distribution, so that other ways of dispersal are available to these animals (see also Cornelius, 1992).

In this paper we report on the life cycle features of two species, relating their developmental features to their strategies of dispersal while leading a larval or post-larval planktonic life.

**MATERIAL AND METHODS**

Colonies of *Clytia viridicans* and *Halecium nanum*, growing on algae of the genus *Cystoseira*, were collected by SCUBA diving from the rocky shore of the Ionian Sea (Porto Cesareo, Italy) during April-July (*C. viridicans*) and October (*H. nanum*) 1997. Fertile colonies were removed from the supporting algae and maintained in glass tanks with filtered seawater (FSW, 0.45 µm). Both temperature and photoperiod were controlled so to match field conditions. The medusae released in the laboratory were fed with *Artemia* nauplii and reared until maturity in FSW. Males and females were kept together so to have fertilisation.

The planulae, either released from the gonotheca (*H. nanum*) or produced in the water after fertilisation (*C. viridicans*), were maintained in glass tanks with FSW (0.22 µm). The presence of zooxanthellae was investigated *in vivo* by a light microscope with fluorescence apparatus.

**RESULTS**

*Clytia viridicans* (Leuckart, 1856)

Metschnikoff (1886a) described a medusa that he retained as identical with those described by Leuckart in 1856 as *Phialidium viridicans*. Later, Metschnikoff (1886b) reported and figured primary hydranths of *C. viridicans* with a gonotheca arising from a basal plate. Both hydroid and medusan features agree with the present material. Russell (1953) considered the possibility that *Clytia flavidula* and *C. viridicans* were referable to a single species.

Later, Cornelius (1982) referred *P. viridicans* to *C. hemisphaerica* because of their similarity. Finally, Calder (1991) described the hydrotheca of *C. hemisphaerica* with “U-shaped pleat extending inwards towards hydrothecal cavity”; he also reported an extensive synonymy list for *C. hemisphaerica*, including in also *P. viridicans* Leuckart, 1856.

The following is the first description of the life cycle of *C. viridicans*.

**Hydroid (Fig. 1a, b)**

Colony stolonal, growing on algae; pedicel short, annulated at base and below hydrotheca; hydrotheca conical, about 0.6 mm high, with 7-9 cusps, projecting inwards and with outwards perisarc projections in the bays between nearby cusps; hydranths about 0.5 mm high with a peduncled hypostome and 14-18 amphicoronate tentacles, the ones oriented downwards laying on the perisarc projections between adjacent cusps, the ones oriented upwards being sustained by cusps; gonothecae on hydrorhiza, either corrugated or smooth, about 0.8 mm high, containing a row of up to four developing green medusae.

**Newly released medusa (Fig. 1c)**

Umbrella hemispherical, about 0.5 mm high, bright green, with four radial canals, four perradial tentacled bulbs, four small interradial bulbs sometimes present; manubrium tubular, with four lips, reaching half of subumbrellar cavity; velum wide, with a small opening; eight statocysts along the circular canal, with one statolyth each. A row of nematocysts often present on exumbrella, parallel to umbrellar margin.

**Medusa development (Fig. 1d)**

Gonad rudiments developed on radial canals two days after release, along with other four tentacles on interradial marginal bulbs and other statocysts between adjacent tentacles. Marginal bulbs, tentacles and statocysts are gradually added and umbrella tends to flatten until, 30-45 days after release, medusae reach maturity.

**Mature medusa (Fig. 1 e)**

Umbrella flattened, about 6 mm wide and 1.5-2 mm high, with 14/16 tentacles; manubrium, gonads and tentacular bulbs bright green, exumbrella trans-
parent; manubrium short, on a short peduncle, mouth with four corrugated lips; gonads on distal part of radial canals, not reaching umbrellar margin.

**Planula production (Fig. 2)**

Spawning took place at night; the eggs, about 200 µm in diameter, were immediately fertilised, and the embryos reached a gastrula stage within 10 hours, to differentiate into free-swimming hollow planulae within 20 hours. Settlement took place within 48 hours and 6-10 days after zygote formation.

**Settlement on the bottom (Fig. 2)**

Most planulae settled on the glass of the rearing jars, forming a wide basal disc from which a primary polyp arose within 12 hours followed, after 24 hours, by a gonotheca. Two days later, 3 or 4 medusae were released. After medusa liberation, some polyps degenerated, but most developed a hydrorhiza and new polyps, producing large colonies that started to develop gonothecae and to release medusae. Several generations were easily obtained under laboratory conditions. No differences were observed in development and growth.
pattern between medusae from natural colonies and medusae from laboratory-reared ones.

Settlement on air-water interface (Fig. 2)

Some planulae settled on the underside of the surface film of water and gave rise to floating colonies that developed for weeks in that position. Accidental break off of the interface induced colony sinking to the bottom. Sunken colonies settled on the bottom of the rearing jars, becoming indistinguishable from those that immediately developed there.

Remarks

The development of gonothecae from the base of primary hydranths confirms Metschnikoff’s observations. Such developmental pattern was never reported for other Clytia species, but recalls what Bouillon et al. (1991) described for Laodicea indica. The bright green colour of the medusa, furthermore, is clearly suggested by the specific name viridicans, from the Latin viridis: green. The adult medusa is similar to Clytia hemisphaerica but the green colour of gonads, manubrium and marginal bulbs is a diagnostic character in living material. The shape of the hydrothecal cusps, with the perisarc projections between adjacent cusps, is a diagnostic character for the hydroid.

The origin of floating colonies directly from a planula is rare for usually benthic hydroids; Billard (1917) reported on three free-living species, two of which perhaps represent fragments of detached colonies, while the third species, Campanularia pelagica Van Breeman can be derived from an unsettled planula or from a planula settled on a sand grain (see also Cornelius 1995). Cornelius (1982) considered Campanularia pelagica as identical with Clytia hemisphaerica.
**Halecium nanum** Alder, 1859

Together with *H. conicum*, *H. pusillum* and *H. tenellum*, *Halecium nanum* is one of the small *Halecium* of the Mediterranean. Motz-Kossowska (1911) provided a good description of this species which, in spite of being rather common, has been rarely treated in hydroid literature.

**Planula production**

The colonies of *Halecium nanum*, as well as the planula, are packed with zooxanthellae. The solid planulae, with a ciliate epidermis, moved slowly by ciliary propulsion. They also crawled on the bottom with peristaltic movements, continuously changing from an elongated to a spherical shape.

The majority of the about 100 planulae released in the laboratory died within one week, but two survived and went through the following states:
- 0-30 days: they continuously crawled on the jar bottom;
- 31-70 days: they stopped, and became nearly spherical, as if they were encysted;
- 71-75 days: they became active again, acquiring a pear shape with a pike from which, after three days, a polyp was born;
- 76-100 days: a colony with five hydranths was built.

**Remarks**

The colonies produced by the two planulae had zooxanthellae in their tissues and regressed to hydroid within a week after settlement; they were not kept further to see if they were able to regenerate after a period of rest.

**DISCUSSION**

Planula dispersal is generally limited because planulae are lecitothrophic: in most hydrozoan species, in fact, planula settlement occurs within few hours or days (Sommer, 1992), so their contribution to dispersal should be little. Planula types reflect the vagility of species with and without medusa: those produced by medusae are usually hollow and able to swim, so to reach the bottom from the water column, whereas those produced by hydroids are usually solid, do not swim much and just crawl on the bottom (see Bouillon, 1994). These different properties should lead to higher endemicity in species without medusae and wider distribution in species with medusa. This is not what Boero and Bouillon (1993) found in the distribution of Mediterranean species, hypothesising a dispersive role also for the hydroid. The specialised asexual propagules of *Halecium pusillum* are a paradigm of hydroid dispersal (Huvé, 1955), even though many species simply disperse by colony fragments.

Medusae are obvious dispersal agents, but they can become ripe while distant from proper settling substrates for the planulae, so that sexual reproduction might result in a failure. Such inconvenient can be overcome by prolonging the life of medusae by fission, by multiple gonad ripening, or by production of gonothecae and fertile polyps on the medusa, but in *Clytia viridicans* this risk is faced with the possibility for the planula to produce hydroid colonies attached to the air-water interface that can become secondarily benthic.

Cornelius (1992) discussed the possibility for hydroids to settle on floating organisms and raft on them so to become widely dispersed. The efficacy of this way of dispersal has not been tested experimentally but, however, is probably very high for the hydroids that settle on algae, such as the ones we investigated. That of *Clytia viridicans*, with planulae producing floating hydroids, is a particular case of rafting, because the hydroid itself is the raft, contrary to what happens for the species settling on other organisms, labelled as rafters. Floating hydroids are usually originated by fragmentation of benthic colonies whereas, for *C. viridicans*, the rafting colonies derive from planulae. It might be questionable whether this planula behaviour can really occur in the field. The study of many hydrozoan species, to our knowledge, never recorded such planula behaviour whereas, in *C. viridicans*, it occurs quite often. Particularly calm conditions are common in the Mediterranean during summer months, so allowing planula settlement on the interface. Once the colony is established, then, it can even become detached from the surface and be transported by the currents.

As described for dozens of species by Boero and Fresi (1986), many hydroids deprived of medusa paradoxically, after sexual reproduction, “disappear” for several months instead of becoming more abundant. The zooxanthellate planulae of *Halecium nanum*, remaining viable for at least three months, are one of the keys of this lapse between planula production and appearance of new colonies. For
species deprived of zooxanthellae, another possibility might be that planulae settle, become wrapped by perisarc and remain dormant until the return of favourable conditions.

These planula behaviours were observed in the laboratory and we do not know their importance in the field. The contribution of planulae to species dispersal, however, might be greater than usually thought and it is possible that further types of planula development will be found in the future, showing still unexpected aspects of the renewed developmental plasticity of hydrozoans.

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