Short-term post-mortality scavenging and longer term recovery after anoxia in the northern Adriatic Sea

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Abstract. The northern Adriatic Sea is one of nearly 500 areas worldwide suffering widespread mortalities due to anoxia. The present study documents post-anoxia macrofauna dynamics after experimentally inducing small-scale anoxia in 24 m depth (2 plots, each 50 cm x 50 cm). Time-lapse camera deployments examined short-term scavenging of the moribund and dead organisms (multi-species clumps consisting of sponges and ascidians) over two 3-day periods (August 2009: 71.5 h, September 2009: 67.5 h). Longer term recovery (days to 2 yr) in the same two plots was examined with an independent photo series. Scavengers arrived quickly and in a distinct sequence: demersal (Gobius niger, Serranus hepatus) and benthopelagic fishes (Diplodus vulgaris, Pagellus erythrinus), followed by hermit crabs (Paguristes eremita, showing a clear day/night rhythm in presence) and gastropods (Hexaplex trunculus). This sequence is attributed to the relative speeds and densities of the organisms. The sessile fauna was largely removed or consumed within seven (August plot) and 13 (September plot) days after anoxia, confirming our first hypothesis that decaying organisms are quickly utilised. The scavengers remained in dense aggregations (e.g. up to 33 P. eremita individuals at one time) as long as dead organisms were available. No recovery of sessile macrobenthos occurred in the experimental plots one and two years after anoxia, undermining our second hypothesis that small denuded areas are more rapidly recolonised. This study underlines the sensitivity of this soft-bottom community and supports calls for reducing additional anthropogenic disturbances such as fishing practices that further impede recolonisation and threaten benthic community structure and function over the long term.

1 Introduction

Dayton et al. (1995) established that coastal marine ecosystems are the world’s most endangered habitats, and the Mediterranean is no exception, with documented impacts on biodiversity (Danovaro and Pusceddu, 2007; Coll et al., 2010). One major disturbance, coastal hypoxia and anoxia, has been exacerbated by eutrophication. No other environmental variable than dissolved oxygen has changed in shallow coastal seas and estuaries so drastically in such a short time (Diaz and Rosenberg, 1995). Hypoxia adversely affects the community structure and trophodynamics of marine ecosystems (Gray et al., 2002), for example by eliminating sensitive species and promoting more tolerant species (Wu, 2002).

The northern Adriatic is one of nearly 500 eutrophication-associated dead zones worldwide, with a global area of more than 245,000 km² (Diaz and Rosenberg, 2008). It combines many of the features known to promote late summer hypoxia and anoxia (shallow depth, soft sediments, semi-enclosed water body, stratification, high riverine input, high productivity, long water-residence time; Stachowitsch and Avcin, 1988; Malej, 1995; Stachowitsch, 1991). Combined with anthropogenic eutrophication and massive marine snow events, the northern Adriatic Sea is a sensitive ecosystem (Stachowitsch, 1986) and also among the most heavily exploited and degraded seas worldwide (Lotze et al., 2011a). Bottom-water oxygen content has decreased from the mid-20th century (Justic, 1991) until the early 1990s, associated with extensive mortalities of epibenthic communities (Fedra et al., 1976; Stachowitsch, 1991; Hrs-Brenko et al., 1994).
Fedra et al. (1976) named the wide-ranging community studied here the *Opfiothrix-Reniera-Microcosmus* (ORM) community based on the dominant brittle stars *Opfiothrix quinquemaculata* (Delle Chiaje, 1828), sponges *Reniera* spp., and ascidians *Microcosmus* spp. It is largely composed of suspension-feeders aggregated into so-called multi-species clumps or bioherms (Zuschin and Stachowitsch, 2009). Filter- and suspension feeder communities here are an important stabilizing compartment that removes enormous amounts of suspended material from the waterbody (Ott and Fedra, 1977). Such communities have been termed a natural eutrophication control (Officer et al., 1982) and thus provide crucial ecological goods and services.

Damage to or loss of such communities has ecosystem-wide implications. During hypoxia and anoxia, the benthic organisms in the ORM community show a distinct sequence of behaviours and mortalities. This has been documented in situ during larger-scale anoxic episodes (Stachowitsch, 1984) and has been confirmed and analysed in great detail experimentally using an underwater chamber (0.25 m$^2$) on a small scale on the seafloor in 24 m water depth. These behaviours include, for example, emergence of infauna (Riedel et al., 2008a), unusual interactions including predation (Riedel et al. 2008b; Haselmair et al. 2010), altered locomotion and activity patterns (Prettereber et al., 2012), and a clear sequence of mortalities (Riedel et al., 2012). For an overview of these responses see Riedel et al. (this issue) and a sample 4-day film at http://phaidra.univie.ac.at/o:87923.

The longer term implications are less well known. The collapse of benthic communities and recurring disturbances – hypoxia/anoxia, marine snow events, fisheries damage – raise fundamental issues of ecosystem stability and resilience. One approach to addressing these issues is to examine post-disturbance events and successions. Two basic recolonisation strategies have been described: motile species immigrating into denuded areas (smaller scale damage) or larval settlement by pelagic recruits (Pearson and Rosenberg, 1978; Whitlatch et al., 1998; Norkko et al., 2010). The re-establishment of community structure on northern Adriatic soft bottoms requires biogenic structures (bivalve and gastropod shells, sea urchin tests) on which larvae can settle. Sedimentation, however, can cover such structures and hamper new epigrowth (Zuschin and Stachowitsch, 2009). Compared to mortality events, recovery is a much longer term process. For example, in a Swedish fjord it took about 3 yr after anoxia for the top sediment to recover to a state suitable for macrofauna recolonisation (Rosenberg, 1971). Recovery of benthic communities after experimental trawling in Scotland took more than 18 months (Tuck et al., 1998). Intertidal communities from muddy sand habitats also showed very slow biological and physical recovery rates (Dernie et al., 2003). In the northern Adriatic, Stachowitsch and Fuchs (1995) were unable to detect full recovery after an oxygen depletion event even after more than ten years, whereby recolonisation was repeatedly interrupted by commercial fishing damage (Kollmann and Stachowitsch, 2001). Elsewhere in the northern Adriatic, Rapido trawling significantly changed community structure and mean abundance of common taxa (Pranovi et al., 2000), and beam trawling is also known to alter the physical characteristics of the sea floor (Kaiser and Spencer, 1996). Such bottom sampling equipment can, in turn, increase oxygen consumption and nutrient concentrations and promote phytoplankton primary production, which changes overall community structures over the long term (Riemann and Hoffmann, 1991; Jennings and Kaiser, 1998).

When long-lived, sessile suspension feeders are reduced by disturbances, benthic communities often experience a shift to mobile detritus feeders; their activities can hinder recovery (Dayton et al., 1995). Such altered communities are often dominated by juvenile stages, mobile species and rapid colonisers (Pearson and Rosenberg, 1978). In the northern Adriatic Sea, for example, hermit crab numbers increased over a 5 yr period after anoxia, but the proportion of the keystone organisms such as brittle stars, sponges and ascidians remained low (Kollmann and Stachowitsch, 2001). Such long-term recovery processes make benthic communities the “memory” of disturbances (Stachowitsch, 1992).

The present work assesses both the immediate, short-term scavenging processes and the longer term recovery processes after artificially induced anoxia in the northern Adriatic Sea. Based on the rapid attraction of scavengers to baited traps as well as the rapid removal of discards and moribund benthos in the beam-trawl fishery for flatfish in the North Sea (Groenewold and Fonds, 2000), coupled with the high density of potential predators and scavengers in the Adriatic, we hypothesize that attracted scavengers quickly remove the damaged benthos. Although we expect full recovery (establishment of multi-species clumps) to be a longer term process, we also hypothesize that, based on the small scale of our experiments, recovery will be initiated relatively rapidly (i.e. weeks to months). We tested our hypotheses in two plots after experimentally induced anoxia using a time-lapse camera followed by longer term photographic series.

### 2 Material and methods

The activities of seven scavengers were examined. These encompass all the species that entered the experimental quadrats after the induced anoxia. The fact that six of the seven species were present in both experiments indicates that we have encompassed the typical set of organisms that potentially removes dead and moribund benthos in the study area. These scavengers included two invertebrates and four vertebrates: the murcid gastropod *Hexaplex trunculus* (Linnaeus, 1758), the hermit crab *Paguristes eremita* (Linnaeus, 1767), and four fish species: the demersal goby *Gobius niger* (Linnaeus, 1758) and comber *Serranus hepatus* (Linnaeus, 1758) as well as the benthopelagic bream *Diplodus vulgaris*

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Fig. 1: Study site; the northern Adriatic Sea (left), oceanographic buoy off Piran in the Gulf of Trieste (right).

(Geoffroy Saint-Hilaire, 1817) and pandora Pagellus erythrinus (Linnaeus, 1758).

2.1 Study site

The study site is located in the Gulf of Trieste, northern Adriatic Sea (45°32′55.68″ N, 13°33′1.89″ E) off Piran, Slovenia (Fig. 1). It is about 2 km offshore at a depth of 24 m, near the oceanographic buoy of the Marine Biology Station in Piran. This position was chosen to minimise disturbance and damage by commercial fisheries. No evidence of trawling was found at the experimental site (fishing boats maintain a safety distance from the buoy, no anchoring by pleasure boats or sport fishers is permitted within 50 m). The soft bottom is dominated by the ORM community described above. Directly under the buoy there is an accumulation of mussels that have fallen from the anchoring chains. This is associated with a higher density of gastropods, but our experimental plots were outside this immediate zone.

2.2 Experimental set-up

Two different benthic chamber configurations were used. The “closed configuration” creates anoxia by sealing a 50 cm × 50 cm × 50 cm volume off from the surrounding water with Plexiglass® walls. The “open configuration”, in which the Plexiglass® chamber is replaced by an open frame of the same size, is designed to observe behaviour under normoxic/post-anoxic conditions. A lid on top of the chamber/frame holds a digital time-lapse camera (Canon EOS 30D) with a zoom lens (Canon EFS 10–22 mm), two battery packs (9Ah Panasonic, Werner light power, Unterwassertechnik, Germany) and two flashes (“midi analog” series 11897; Subtronic, Germany). The battery packs, combined with a specially designed electronic control circuit, enable the equipment to be operated for about 72 h. Finally, a datalogger (PA3000UD, Unisense, Denmark) and a sensor array for measuring oxygen, temperature and hydrogen sulfide are also located on the lid (for a detailed description see Stachowitsch et al., 2007). Images were produced at 3 min intervals, sensor data logged every minute.

This configuration was the largest chamber (and instrument lid) size that could be handled by divers at 24 m depth and is among the largest benthic chambers deployed on a sublittoral soft bottom. A greater volume was also hindered by the generally poor visibility in this area and corresponding short camera distance from the sediment. The chamber/frame was large enough to enclose bioherms and the surrounding sediment, and such larger chambers are also preferable in benthic oxygen uptake experiments over smaller chambers (and extrapolations therefrom) (Glud and Blackburn, 2002).

The present two experiments involved the following deployment protocol. The underwater device was initially positioned in its closed configuration over a representative multi-species clump (with both sponges and ascidians). After max. 72 h, and creation of anoxia (visible in time-lapse films by mortality of the macrobenthos and blackened sediment – confirmed to be associated with anoxia by sensor data in a series of earlier experiments using this benthic chamber), the lid was lifted and brought to the surface to exchange the battery packs and download the images (the chamber remained sealed with a cover). Immediately thereafter (overall surface procedure < 1 h), the lid was again transported to the deployment site and the chamber replaced by the open frame,
marking the beginning of the post-anoxia recovery experiment. Post-mortality scavenging was documented for about 3 days. Combined, the two deployments yielded a dense data set comprising 139 h of autonomous time-lapse recordings with a total of 2814 post-disturbance images.

2.3 Short-term scavenging

Two experiments were performed. In the first, post-disturbance images (open configuration) were taken from 15 August 2009 (10:35 a.m. LT) to 18 August 2009 (10:05 a.m.) (“August” experiment). The second experiment extended from 14 September 2009 (03:25 p.m.) to 17 September 2009 (10:49 a.m.) (“September” experiment). The August experiment yielded a total of 1430 images, the September experiment 1384 images. This corresponds to a documentation time of 71.5 and 67.5 h, respectively. The images were also processed into time-lapse movies using the Adobe Premiere 6.5 program (August recovery time-lapse film available at http://phaidra.univie.ac.at/o:262380). The fishes were analysed by image: the data are summarised in 6 h steps. Because of the relatively slow movements of the hermit crabs and gastropods and the more gradual changes in their numbers, only every tenth image was evaluated for these groups. For every analysed image, the number of individuals of each scavenger species were recorded. The different multi-species clump substrates (sponges, ascidians, sediment) that the hermit crabs and gastropods were on were recorded. Normally, these two taxa are found only on the sediment surface. We therefore equate the dead or moribund organisms they chose with their preferred prey items. The time and sequence of arrival/departure and the maximum number of individuals were recorded. The hour-values in the Results refer to the times after removal of the benthic chamber (and not to the time after anoxia was generated inside the chamber). Day/night activity (based on sunrise and sunset, www.sun.exnatura.org) of hermit crabs and gastropods was analysed.

Individual specimens of the same species could not always be differentiated throughout a deployment because coloration and sizes were typically very similar and the perspectives often changed considerably. Accordingly, we use “observations” and “number of observed specimens” to refer to the number of fish, gastropod and hermit crab “individuals”, i.e. representatives of the species visible over the respective time period. Thus, a total of 12 gobies visible in 6 h, for example, could in theory represent between 1 and 12 separate individuals. In these territorial fish, however, we expect the actual number of specimens to be at the lower end of the range. Equally, many crab and gastropod individuals remained in the plots for long periods. While the number of “observed specimens” clearly overestimates the actual number of different individuals present, it does reflect the overall scavenging influence exerted by the respective species during that time.

2.4 Longer term recovery

These experiments were conducted on the same two 0.25 m² plots used for the time-lapse camera study. The pre-disturbance surface area occupied by living sponges and ascidians, as well as by other hard structures and lebensspuren, was calculated using CorelDraw9 and Excel 2010 based on the photos taken from the closed configuration. The corresponding surface areas were then measured again days to years after anoxia. The area occupied by recolonizers was calculated as a measure of recovery. Each plot was subdivided into four quadrants and an image taken of each quadrant with a hand-held camera (i.e. four macro-photos per plot per sampling date). The images of the August plot were taken on 18 (day 1 of long-term experiment), 22, 24 and 25 August 2009, those of the September plot on 19 (day 1 of long-term experiment), 20, 23, and 30 September 2009. Both plots were also photographed twice in each following year, namely, on 7 and 17 August 2010, and 6 and 7 August 2011. The objects were classified into four categories: (1) sessile fauna (living/dead) and anemones, (2) shells (bivalves and gastropods), (3) small pieces of shells and areas covered with many shell fragments (“coquina”) and (4) lebensspuren from vagile fauna and endofauna burrow openings (data not presented).

To better compare the two experiments, the times when the images were taken were categorised (e.g. 7–13 days is termed “10 days” in the following text). The data in the two sets of photographs taken in short succession at 1 yr and 2 yr at each plot are averaged, and referred to as “1 yr” and “2 yr”, respectively, unless stated otherwise.

Time series analyses and cross correlations were calculated to define the residence time of gastropods and hermit crabs in the plots. Graphs of time series analysis can be found in Blasnig (2012). For statistical analyses the program Past was used (Hammer et al. 2001).

3 Results

3.1 Short-term scavenging after anoxia

Moribund and dead sponges and ascidians attracted fishes, hermit crabs and gastropods in a rapid and distinct sequence. Numerous fishes arrived in the first hours. They were followed by hermit crabs (Paguristes eremita), which showed a rapid initial increase in the number of individuals. Finally, the gastropods (Hexaplex trunculus) appeared (Fig. 2), with a slow increase of individuals.

3.2 The fishes

In the August 2009 experiment (71.5 h), we observed three different species, Diplodus vulgaris, Serranus hepatus and Gobius niger, and additionally in the September 2009 experiment (67.5 h) we observed Pagellus erythrinus. In both...
3.3 The first 12 h: hermit crabs and gastropods

The first 12 h were evaluated separately to better determine the order in which the species and individuals arrived. In August, the first three *Paguristes eremita* were observed after 30 min and the number rapidly increased up to 19 individuals after 9 h. After 3 h the first *Hexaplex trunculus* arrived and the number slowly increased to three individuals after 8 h (Fig. 2a). In September, the first *P. eremita* appeared after 1 h, and numbers then increased up to 12 individuals after 6 h, before decreasing again to nine. Seven *H. trunculus* individuals survived the anoxia (Fig. 2b) and were present and visible from the start. After 30 min the first new gastropod entered the frame. Two hours later, three individuals again left the area (5 present), but thereafter the number increased steadily (max. 17).

3.4 Day/night activity

In both experiments the number of *Paguristes eremita* varied considerably over time. The values decreased in all three nights examined and increased conspicuously during daylight hours (Fig. 3). During the “day” the number of observed specimens in August increased to 24 (45.5 h after closed chamber removal) and 33 (after 71 h), while at night, values fell to six and to three (after 38 and 58 h, respectively). In September, the number peaked during daylight at 28 (17.5 h after closed chamber removal) and dropped to only two (after 37 h) and three observations (after 62.5 h). The time series analysis for *P. eremita* showed highly significant peaks at 21.2 h (August), 28.4 and 54 h (September), a significant peak at 12.4 h (August), and distinct but not significant peaks at 15.5 and 57.2 h (August) and at 10.8 h (September) (Chi-squared test, data not shown). These peaks demonstrate a semidiurnal and diurnal periodicity.

*Hexaplex trunculus* showed a relatively slow but constant increase, levelling off somewhat over the last day. The peak number was eight observations at three periods before 46.5 and 65.5 h in August, and 17 observations after 38.5 h in September. At the end of both experiments, 4 (August: 71.5 h) and 13 (September: 67.5 h) *H. trunculus* were still present in the plots. The time series analysis showed highly significant peaks (data not shown) at 18.5 and 38.1 h (August) as well as 13.5, 27.0 and 60.0 h (September). Visible but not significant peaks are present at 8.2, 9.7 and 11.2 h (August) as well as at 18 h (September). These peaks point to a periodicity of nine and 12 h as well as multiples thereof.

The fishes show highly significant (data not shown) peaks at 9.9 h (August) as well as 12.3 and 21.6 h (September) and visible but not significant peaks at 30.1 h (August) 6.4 and 7.8 h (September)(Fig. 2). This points to a semidiurnal and a roughly diurnal periodicity.

In the first 6 h of the August experiment, all three species were present: *G. niger* with 83, *D. vulgaris* 35 and *S. hepatus* 26 observations. The two latter species then decreased to very low numbers or zero in the successive 6 h periods. *G. niger* also decreased with time, but never fell below 22 observations per 6 h (i.e. total number visible over this time period, not necessarily different individuals, see Material and Methods).

In the September experiment, all four species were initially present and showed the highest abundance in the first 6 h time step: *G. niger* peaked with 80, *D. vulgaris* 18 and *S. hepatus* 13 observations. The number of observations decreased markedly by the end of the experiment.
3.5 Preferred substrates/prey items

The substrates that *Paguristes eremita* and *Hexaplex trunculus* chose were different and consistent in both experiments. Hermit crabs were observed on the sediment, on sponges and on ascidians, whereas *H. trunculus* occurred mainly on ascidians and to a lesser extent on the sediment (Fig. 4). All categories (i.e. compared species-experimental month pairs) are significantly different from each other. Importantly, there are bigger interspecific than intraspecific differences: the differences between the species (*P. eremita* in August and September versus *H. trunculus* in August and September) are bigger than between the months (*P. eremita* in August versus September and *H. trunculus* in August versus September) (Chi Square test, data not shown).

The duration of the stay of nine *Paguristes eremita* and nine *Hexaplex trunculus* was calculated. The individuals were chosen based on their recognizability (e.g. shell epigrowth). This duration averaged 5 h and 19 min for *P. eremita* and 12 h and 9 min for *H. trunculus*. Thus, on average, the gastropods stayed more than twice as long as the crabs. In the August experiment, several *P. eremita* dragged off the ascidian *Phallusia mammillata*. Within 53 h, they dragged it (along with an attached anemone *Cereus pedunculatus* and the ascidian *Microcosmus* sp.) 8 cm in one direction, then 21 cm in the opposite direction, just outside the frame. The ascidian, which had become discoloured, was then partially consumed by the end of the film, with pieces being visibly removed.

3.6 Sea anemones

The August experiment contained three *Cereus pedunculatus*: two attached to ascidians (*Microcosmus* sp. and *P. mammillata*), one next to a large sponge. All three survived the anoxia but showed extreme elongation and rotations. One was carried outside the frame by hermit crabs (attached to *P. mammillata*, see above), the second was also flipped out of the frame (attached to *Microcosmus*), probably by hermit crabs.

The third individual, immediately adjacent to the sponge, fully emerged from the substrate and began to crawl away after 66 h and 15 min of re-oxygenation. At the end of the short-term post-disturbance evaluation (71.5 h, 18 August) the specimen was still inside the frame. During the subsequent longer term evaluation this anemone was still alive and visible at that position on the image of 22 August. Two days later, it disappeared from the images.

3.7 Longer term recovery

At the start of the August experiment, 1939 cm$^2$ (77.5 %) were covered by sessile organisms and 540 cm$^2$ (21.5 %) by vagile fauna (mostly hermit crabs and gastropods;
Fig. 5. Coverage of sessile (dead and living) and vagile fauna after anoxia in the August 2009 (above) and September 2009 (below) plot. Note different scales on y axis. Start of experiment defined as 0.1; values measured inside closed chamber (original coverage).

Fig. 6. Coverage of shells after anoxia of the August and September 2009 plot (above) and of coquina (below). Note different scales on y axis. Start of experiment defined as 0.1 and initial values measured inside closed chamber (original coverage).

total values can approach or exceed 100% because animals on living substrates were counted separately). After the initial increase in scavengers (see above), the vagile fauna decreased markedly to 155 cm$^2$ coverage (6%) by day six of the long-term experiment (Fig. 5a). The sessile fauna dropped to 12 cm$^2$ (0.5%). In September, the sessile fauna initially covered 631 cm$^2$ (25%), the vagile fauna 663 cm$^2$ (26.5%). As in August, both groups decreased drastically, with closely parallel trends (Fig. 5b). After six days, the sessile fauna fell to 286 cm$^2$ (11.5%), the vagile fauna to 154 cm$^2$ (6%). In the “10 day” category (see Material and Methods), nearly the whole fauna was consumed (sessile) or had moved away (vagile). Both 1 and 2 yr after the start of the experiment, no macroscopic sessile epifauna was observed on either plot. The exception was one sea anemone, *Ceramium pedunculatus*, in the September experiment; it survived anoxia and was still present in both following years.

Shells (> 1 cm) and smaller shelly material (“coquina”: smaller shells and shell fragments forming loose aggregations on top of the sediment) were visible in both plots (Fig. 6a, b). Overall, in both experiments shell coverage (Fig. 6a) fluctuated but showed similar values at the beginning and end (2 yr). In August, cover increased from an initial 126 cm$^2$ to 185 cm$^2$ after 1 yr, with a drop after 2 yr to 62 cm$^2$. In September, coverage decreased 1 and 2 yr post-anoxia, with a minimum of 46 cm$^2$ after “2 yr”. Interestingly, short-term fluctuations occurred, for example between the 1 yr and the 1 yr + 10-day sampling (67 versus 139 cm$^2$).

Coquina coverage (Fig. 6b) fluctuated more widely and clearly increased over time. The trend in the two experiments
was less similar, with values in the September experiment always exceeding those of August.

In August, no coquina was initially observed, but after 3–4 days the coverage increased to 204 cm$^2$. After “10 days” it fell again to 57 cm$^2$, followed by a peak of 842 cm$^2$ after 1 yr+10-days. Note, however, that 10 days earlier, the value was zero, showing the potential for major fluctuations even over short time periods. In September, values initially decreased from 428 cm$^2$ to 257 cm$^2$, but then steadily increased to 904 cm$^2$, dropping after 1 year to 261 cm$^2$ but peaking again at 1499 cm$^2$ after an additional 10 days (i.e. 1 yr+10-day sampling). Thus, values changed considerably within even a 10 day period. With few exceptions (early August), coquina always covered more surface than shells.

4 Discussion

In the northern Adriatic, hypoxia/anoxia cause rapid mortalities of macroepibenthic communities (Stachowitsch, 1984; Kollmann and Stachowitsch, 2001), but little is known about post-anoxia processes other than that recolonisation is apparently slow and additionally hampered by bottom trawling (Stachowitsch, 1991). The present study extends prior investigations on the behavioural responses and mortality sequences (Riedel et al., this issue, and references therein) using a specially designed underwater chamber and time-lapse system. It helps fill in the gap between community collapse and recolonisation by examining short-term scavenging and longer term recovery processes.

4.1 Short-term post-anoxia scavenging

After anoxia, the moribund and dead organisms quickly attracted scavengers. This confirms our first hypothesis that the attracted scavengers quickly remove the damaged benthos. The process resembles that described after simulated benthic fisheries damage to scallops in the north Irish Sea (Jenkins et al., 2004); while baited, heavily damaged scallops were eaten within 24 h, lightly damaged individuals were inaccessible until large crustaceans (Cancer pagurus) made the body tissue available, causing protracted aggregation and feeding for up to 96 h. Our dead organisms were shell-less, but the tunicas of ascidians, for example, probably also delayed access by less powerful hermit crabs or small gobiids, explaining the presence of scavengers at the end of the experiments (67.5 and 71.5 h).

The images revealed a clear sequence of scavengers. We attribute the quick arrival of fishes mainly to their swimming speed (e.g. Diplodus vulgaris: 12.3 cm s$^{-1}$; Planes et al., 1997), which is orders of magnitude faster than the next arrival, Paguristes eremita (21.6 m d$^{-1}$; average speed 2.1 m h$^{-1}$ = 3.5 cm min$^{-1}$; Stachowitsch, 1979). Other factors include distance from the disturbance and the size of the disturbed area. Gobius niger has an estimated density of about 1 individual 10 m$^{-2}$, Serranus hepatus 1 individual 22 m$^{-2}$ (Wurzian, 1982), so that we expected individuals to be close by. These densities would also help explain the relative proportions we recorded of these two species: Gobius is more abundant and its numbers in the images were more than twice as high as those of Serranus. Based on the speeds of other, smaller gastropods (e.g. Littorina littorea, 2.88–4.47 cm min$^{-1}$; Erlandsson and Kostylev, 1995) and on the similar size of large P. eremita and adult H. trunculus, we believe the speeds of the hermit crab and gastropod are similar. In such cases, arrival times would depend on density, with more abundant species having a greater probability of having individuals close by. In this benthic community, P. eremita has a density between 1.9 individuals m$^{-2}$ (quadrat sampling; Stachowitsch, 1977) and 2.4 individuals m$^{-2}$ (transect method: Pretterebner et al., 2012). H. trunculus, in turn, has a density of 0.2 individuals m$^{-2}$ in the ORM community (Wurzian, 1982), although the value may be slightly higher here due to the proximity of the oceanographic buoy (see Material and Methods, “study site”). This explains why the gastropods arrived last.

Fishes were conspicuously abundant in the first six hours after switching to the open frame. This is the time when the most food items were available, including smaller soft-bodied infaunal organisms freshly emerged from the sediment and cryptic fauna from multi-species clumps. The fishes were dominated by Gobius niger, which feeds on polychaetes, amphipods, mysids and decapods (Richards and Lindeman, 1987), Pagellus erythrinus mainly preys on benthic organisms such as polychaetes, brachyuran crabs and benthic crustaceans (Fanelli et al., 2011), Serranus hepatus on invertebrates, mainly decapods (Labropoulou and Eleftheriou, 1997), and Diplodus vulgaris on benthic echnoids such as Echinocyamus pusillus and Psammechinus micrurus, but also on decapods and bivalves (Pallaoro et al., 2006). In the present study, P. eremita and H. trunculus mainly fed on material that remained after the fishes had already been present for several hours: based on their positions, mostly ascidians (Phallusia mammilata, Microcosmus spp.) and sponges. The time-lapse camera approach was unable to provide direct evidence that the fishes also fed on sponges or ascidians, although the fish were often positioned in an oblique angle with the mouth facing down. Many ascidians and sponges rely on chemical antipredator defences (e.g. Pawlik, 1993). Ascidians also exhibit physical defences (e.g. tunic toughness: Koplovitz and Mc Clintock, 2011). The latter mechanism may still be effective in freshly dead individuals, making them unattractive for small fishes as the quickest arrivals. Young (1989), for example, observed ascidians being eaten by gastropods, which insert their proboscis into their siphon. Our evaluation showed some H. trunculus on the siphon of Microcosmus spp., but other individuals were positioned on other parts of the ascidians.

To better determine food preferences, we evaluated the substrates on which the scavengers positioned themselves,
equating the chosen dead or moribund species with the preferred prey. In some cases, these substrates were clearly eaten and reduced in size. While both species fed on sponges and ascidians, H. trunculus was much more frequently positioned on *Microcosmus*. The August images also showed *P. eremita* dragging off and consuming an ascidian (*Phallusia mammillata*). *H. trunculus*, in contrast, crawled up the ascidians and typically remained there for several hours: on average, the gastropods stayed within the frame about 7 h longer than the crabs. This agrees with Sawyer et al. (2009), who documented 10.5, 5.2 and 2.9 h for feeding and manipulating the mussel prey in three selected *H. trunculus* individuals in this community. Under normoxic/undisturbed conditions, hermit crabs and gastropods are found almost exclusively on the sediment, not on bioherms, so that we have never observed this type or this level of scavenging before.

The semi-diurnal and diurnal rhythm of *P. eremita* adds a new facet to the hermit crab’s activity behaviour. Stachowitsch (1979) recognised two types of movement interruptions based on time-lapse films: short stops related to feeding and investigating structures on the sediment and longer pauses during the night hours reflecting a resting period, i.e. fewer individuals therefore moving on or across the quadrats at night. Our explanation for why fewer individuals remained in the plot near their prey items at night in the present experiments is that, in their dormant night phase, crabs might move away from larger prey because staying would increase the risk that they themselves would be consumed, along with the prey, by other, larger (fish) predators/scavengers. The same may hold true for *H. trunculus* based on the 9 and 12 h periodicity indicated by the time series analysis (Blasnig, 2012).

As a scavenger, the gastropod *H. trunculus* has one advantage over the hermit crab *P. eremita*: it is among the most anoxia-tolerant species in this soft-bottom community (Riedel et al., 2012). All individuals survived anoxia in our September experiment and were therefore immediately present upon re-oxygenation. Molluscs, for example, are generally considered to be more tolerant to hypoxia than many other invertebrate groups (Vaquer-Sunyer and Duarte, 2008). Such tolerance not only improves survival but may also enable more successful predation during and after hypoxia events (Pihl et al., 1992). Although no *P. eremita* survived anoxia in this study, this species is among the most tolerant crustaceans in this community (Riedel et al., 2012), making them potential predators and scavengers during and after shorter mild to moderate hypoxia. In our experiments, the abundant presence of both species in the same plots, and their preference for different substrates/prey, suggest no mobile species interactions (i.e. no competitive interactions for food). Depending on the severity and duration of oxygen depletion, both species may be able to take advantage of more vulnerable prey. Such altered predator–prey relationships may affect community structure. Moreover, the size of the disturbed area is important: in larger areas in which gastropods have survived but hermit crabs did not, the gastropods would have a lengthy advantage before the crabs from outlying regions can repopulate the affected area. Our experimental set-up is unable to mimic that scenario, which could only be tested in future natural collapses. Scaling up presents numerous practical problems (Zajac et al. 1998): simply capturing the short post-anoxia scavenging phase is a challenge in itself, probably explaining the lack of information on such scavenging elsewhere.

The sea anemone *Cereus pedunculatus* – as the only surviving macrofaunal organism beyond *H. trunculus* – showed an unexpected post-anoxia behaviour. It crept away from its original position 66 h and 15 min after the chamber was opened. Anemones, although normally firmly attached, can move, and some hermit crabs for example can detach and re-attach symbiotic anemones to their shells. The movement of *C. pedunculatus* in our experiment is intriguing because it occurred after re-oxygenation rather than during anoxia. One explanation is that the high density of post-anoxia feeding hermit crabs and gastropods disturbed the anemone. Alternatively, the adjoining dead sponge in that plot may have caused unfavourable conditions. Accordingly, the post-anoxia disturbance by aggregating scavengers was apparently more severe than the effect of anoxia itself: in none of the many other behavioural experiments we conducted did anemones ever move away during severe hypoxia or anoxia (with or without H2S conditions), although they showed severe stress reactions (body contractions, rotation, extension; Riedel et al., this issue).

### 4.2 Longer term recovery

Even 2 yr after anoxia, no macroepibenthic recovery occurred in either of our plots. This contradicted our second hypothesis that recovery will be initiated rapidly at the relatively small scale of our experiments. Our conclusions are not a function of image resolution (macro-photos), and sample collection would not have yielded recruit epigrowth. Benthic recolonisation is, no doubt, scale dependent and can involve larval settlement (larger scale, longer term) and immigration by vagile organisms (smaller scale, shorter term) (Güntner, 1992; Whittlatch et al., 1998). Past mortalities in the northern Adriatic Sea ranged from several km² (Stachowitsch, 1992) to thousands of square kilometres (Stefanon and Boldrin, 1982). In our small-scale experiments, both recolonisation strategies are conceivable, but we would expect immigration to play the more immediate role. However, replacement of the sponges and ascidians that form the characteristic multispecies clumps may be more difficult than thought, despite adjoining bioherms (as larval producers) and vagile organisms (as immigrants).

Based on the abundant *lebensspuren* (data not shown), vagile species did criss-cross the denuded surface, but this does not lead to recolonisation with bioherms. Nonetheless, vagile fauna plays the key post-anoxia scavenging role. The rapid removal of moribund or dead sponges and ascidians

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within the first six days (August) and 13 days (September) after anoxia is attributed to the fauna that entered the experimental plots (or by the *H. trunculus* that survived inside the plots). The subsequent decline in food items was correlated with a corresponding drop in scavengers. The vagile fauna photographed in the plots after 1 and 2 yr consisted mainly of *P. eremita*, *H. trunculus*, the sea urchin *Psammechinus microtuberculatus* and the brittle star *Ophiothrix quinquelaculata*. Kollmann and Stachowitsch (2001) consider *lebensspuren* to be helpful in quantifying community status and activity of mobile forms. The *lebensspuren* coverage in our plots (data not shown) points to abundant vagile fauna here. Such high densities, however, may also keep larval recruits low: grazing on and manipulation of the shells that the multi-species clumps initially require to grow on (Zuschin et al., 1999) could also help explain the lack of recovery. This could be interpreted as a disturbance agent although, as opposed to fishing gear, no physical damage occurs.

We explain the unexpected lack of longer term recolonisation with the fact that settlement and growth of sessile organisms depend on the presence of adequate substrates. In the northern Adriatic, these are typically empty gastropod and bivalve shells or echinoderm tests (Zuschin and Stachowitsch, 2009 and references therein). Although we recorded shells in our plots, they were apparently not large enough, sufficiently exposed, or available at larval settlement times. Thus, the percent coverage by shells fluctuated and ultimately declined after 2 yr. Small-scale factors that affect availability include manipulation by vagile species or removal (use as camouflage) by the sea urchins. One shortcut to the establishment of multi-species clumps in this community – the deposition of heavily encrusted gastropod shells by hermit crabs (Stachowitsch, 1980) – did not take place within our two plots in the two years examined.

Beyond biological factors, anthropogenic factors play a role which we could not mimic with our plots. Nonetheless, outside the protected zone around the oceanographic buoy, fishery gear such as dredges and bottom trawls leaves deep tracks, crushes multi-species clumps, shears off structures projecting from the bottom (e.g. pen shells), and overturns shells, impacting invertebrates growing on them (Stachowitsch and Fuchs, 1995). Fisheries and the passage of large ships also re-suspend enormous amounts of sediment, creating major turbidity and sedimentation events. Such anthropogenic resuspension is at work in our experimental site, where water currents can transport such suspended matter from afar. This sedimentation may also hinder larval-based recolonisation and epigrowth by covering available hard substrates. Such activities or events may help explain the considerable variation in the visible coverage by shells and the sudden changes in visible coquina within only a few days.

Anoxia-related disturbance and recovery in the northern Adriatic has been described as “rapid death – slow recovery” (Stachowitsch, 1991): most organisms die shortly after anoxia is attained, but recovery takes years. The ORM community showed little recovery 10 years after the collapse in 1983; although certain vagile fauna such as hermit crabs increased, larger multi-species clumps did not develop. This was attributed to repeated anoxia and fisheries damage (Stachowitsch and Fuchs, 1995; Kollmann and Stachowitsch, 2001). Recovery is therefore also determined by the frequency of disturbance. In wide areas of the Gulf of Trieste, northern Adriatic, oxygen depletion and the above combination of factors have altered the community structure observed by Fedra et al. (1976) in the early 1970s. This points to the difficulty in restoring community structure and function in the northern Adriatic. It also supports the notion that recovery from human disturbances takes longer than from natural disturbances and, in complex marine ecosystems with slow successions, recovery could take decades or longer (Jones and Schmitz, 2009; Lotze et al., 2011b). It is therefore not surprising that we failed to observe any recovery of sessile organisms during our experiments. Despite prior knowledge about lengthy recovery based on diver observations, random scuba-diver-taken samples and phototransect work (Kollmann and Stachowitsch, 2001), this study is the first to confirm the lack of recovery in specific quadrats with known pre-anoxia macroepifauna composition.

Using small-scale experiments to predict larger-scale recolonisation and succession is difficult because the combined factors controlling these processes at different spatial scales may differ fundamentally (Zajac et al. 1998). Despite our relatively small plot size and lack of replication, we nonetheless argue that our results are a step forward in distinguishing potential post-anoxia scenarios after differently sized collapses. Thus, the sequence of arrivals (fish, hermit crabs, gastropods) – related to their relative speeds and densities – and the roles the organisms played shed light on potential larger-scale phenomena. In extensive anoxic events, immediate immigration would be slower due to the greater distances involved, but the sequence of arrivals would presumably be the same. Surviving species (gastropods) would have a long-term scavenging advantage to the exclusion of more sensitive immigrants. Despite the inherent problems of scale, our results point to alarming long-term effects and call for intensive management measures to reduce further pollution and physical destruction of shallow-marine environments.

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References


