CellTracker Green labelling vs. rose bengal staining: CTG wins by points in distinguishing living from dead anoxia-impacted copepods and nematodes

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Abstract. Hypoxia and anoxia have become a key threat to shallow coastal seas. Much is known about their impact on macrofauna, less on meiofauna. In an attempt to shed more light on the latter group, in particular from a process-oriented view, we experimentally induced short-term anoxia (1 week) in the northern Adriatic Sea (Mediterranean) and examined the two most abundant meiofauna taxa – harpacticoid copepods and nematodes. Both taxa also represent different ends of the tolerance spectrum, with copepods being the most sensitive and nematodes among the most tolerant. We compared two methods: CellTracker Green (CTG) – new labelling approach for meiofauna – with the traditional rose bengal (RB) staining method. CTG binds to active enzymes and therefore colours live organisms only. The two methods show considerable differences in the number of living and dead individuals of both meiofauna taxa. Generally, RB will stain dead but not yet decomposed copepods and nematodes equally as it does live ones. Specifically, RB significantly overestimated the number of living copepods in all sediment layers in anoxic samples, but not in any normoxic samples. In contrast, for nematodes, the methods did not show such a clear difference between anoxia and normoxia. RB overestimated the number of living nematodes in the top sediment layer of normoxic samples, which implies an overestimation of the overall live nematofauna. For monitoring and biodiversity studies, the RB method might be sufficient, but for more precise quantification of community degradation, especially after an oxygen depletion event, CTG labelling is a better tool. Moreover, it clearly highlights the surviving species within the copepod or nematode community. As already accepted for foraminiferal research, we demonstrate that the CTG labelling is also valid for other meiofauna groups.

1 Introduction

Detecting disturbance-induced changes in animal communities from a process-oriented perspective often requires short-term experiments and accurate detection techniques. This is especially true in examining stress-sensitive species and community threshold dynamics/responses. Coastal hypoxia (low dissolved oxygen DO < 2 mL L⁻¹; Diaz and Rosenberg, 2008) and anoxia have become recognized as a key emerging problem in the last decades, with an increasing number of affected sites coupled with increasing intensity, frequency and duration of events (Gooday et al., 2009; Rabalais et al., 2010; Zhang et al., 2010, see; also this issue and references therein). Adequate measurements to monitor the impacted fauna and identify sensitive and more tolerant species is therefore pivotal, especially when the course of oxygen decline is rapid, e.g. within several days to a week, as in the Adriatic Sea (Stachowitsch, 1984, 1991; Faganeli et al., 1985; Justić et al., 1993). While the mortality of macrofauna organisms can often be visually identified from changes in colour, behaviour and/or body shape of individuals (Riedel et al., 2012), the discrimination of dead from living meiofauna individuals requires more efficient methods and often a taxonomist’s expertise.
Beyond examining captured meiofauna organisms individually to see which are still moving or showing other signs of life (Steyaert et al., 2007), staining of freshly captured organisms is the traditional method used (Eleftheriou and McIntyre, 2005). This is the only feasible method to distinguish between living and dead organisms in the abundant and numerous replicated samples necessary to answer specific ecological questions. Recently, however, a number of foraminiferal studies showed that the traditional rose bengal (RB) staining method (Walton, 1952) is often inadequate to distinguish with certainty living from dead organisms.

The RB staining method is commonly used in meiofauna studies (Thiel, 1966; Tiemann and Betz, 1979; Higgins and Thiel, 1988; Eleftheriou and McIntyre, 2005; Giere, 2009). The RB is a typical bulk stain which adheres to (cytoplasmatic) proteins and is applied into formalin-fixed samples (Higgins and Thiel, 1988; Somerfield et al., 2005). As a stain it has various advantages (cheap, simple to apply, animals are easily visible under light microscope), but on the negative side it is a nonvital stain that stains proteins regardless of the cell/animal is dead or alive (Bernhard et al., 2006). It also colours different meiofauna organisms differently (Giere, 2009). Nonspecificity of RB stain explains the use of vital stains in foraminiferal studies (Bernhard et al., 1995, 2006; Bernhard, 2000; Pucci et al., 2009). The cytoplasm of Foraminifera stays in the shell for a long period after death, particularly in anoxic conditions, and it still stains well with RB (Bernhard et al., 1995, 2006; Bernhard, 2000; Pucci et al., 2009).

The present study is based on the paper from Bernhard et al. (2006), who compared CellTracker Green (CTG) labelling with rose bengal staining to distinguish live from dead benthic foraminifera. The principal idea behind the approach is to determine the most accurate method that exclusively labels surviving cells/organisms (Bernhard et al., 2006; Peperzak and Brussaard, 2011) in short-term disturbance studies such as hypoxia/anoxia experiments. When living cells are incubated with fluorogenic probes such as CTG, the probe passes through the cellular membrane and reaches the cytoplasm, where hydrolysis with nonspecific esterase causes the fluorogenic reaction (Bernhard et al., 2006; Pucci et al., 2009; Morigi and Geslin, 2009; Heinz and Geslin, 2012). Once in the cell, the CTG probe is converted to cell-impermeant reaction products (Peperzak and Brussaard, 2011). CTG is applied in many fields, such as medicine e.g. human tissue cultures (Boleti et al., 2000); parasitology e.g. drug–multicellular parasites relation (Trejo-Chávez et al., 2011), phytoplankton ecology (Peperzak and Brussaard, 2011); and microbiology – i.e. benthic microalgae, ciliates, flagellates and foraminifera (Bernhard et al., 2003, 2006; Pucci et al., 2009; First and Hollibaugh, 2010; Figueira et al., 2012, Langlet et al., 2013a,b). The use of CTG for metazoan organisms, however, has so far been limited to dysoxic laminated sediments in 400 to 600 m water depth (oxygen minimum zone in the Santa Barbara Basin; Bernhard et al., 2003), and deep-sea (3300 to 3600 m) anoxic sediments (Mediterranean; Danovaro et al., 2010). Here, we specifically test its usefulness to quantify live meiofauna (harpacticoid copepods, nematodes) in coastal sublittoral anoxic vs. oxic sediments.

As in most marine sediments, harpacticoid copepods (Crustacea, Copepoda) and nematodes (Nematoda) are the two most abundant meiofauna taxa in the soft sublittoral sediment in the Gulf of Trieste, northern Adriatic Sea (Vrišer, 1984; Travisi, 2000; Grego et al., 2009). The two groups are morphologically very different and are known to have very different tolerances to low dissolved oxygen conditions. Copepods are in general more sensitive; according to Vernberg and Coull (1975) they tolerate anoxia from 1 h (sand dwellers) to a maximum of 5 days (mud dwellers). They show a drop in abundance and diversity with progressing hypoxia/anoxia (Hicks and Coull, 1983; Moore and Bett, 1989; Murrell and Fleeger, 1989; Moodley et al., 1997; Modig and Ölafsson, 1998; Wetzel et al., 2001; Levin, 2003). Grego et al. (2013) describes the change in copepod community composition due to anoxia in more detail. Nematodes, in contrast, can survive up to several months of extended periods of hypoxia/anoxia and show little decrease in biodiversity before the system reaches longer term anoxic conditions (Josefson and Vidbom, 1988; Murrell and Fleeger, 1989; Austen and Vidbom, 1991; Hendelberg and Jensen, 1993; Wetzel et al., 2001; Levin, 2003; Van Colen et al., 2009). In oxygen minimum zones, for example, nematodes are found in higher densities (Vitt-Köhler et al., 2009) compared to surrounding well-oxygenated sediments, presumably benefitting from high food supply and low predation and competition pressure (Neira et al., 2001). Certain species, however, are exceptions: copepods that tolerate anoxia (Vopel et al., 1998) or surface-dwelling species of nematodes that die when oxic conditions become hypoxic (Modig and Ölafsson, 1998). It is difficult to define the impact of oxygen concentration alone on benthic communities because, with the progress of hypoxia to anoxia, the concentration of sulfide increases. Sulfide is toxic for most of the fauna, although some species have developed strategies to detoxify it (Vaquer-Sunyer and Duarte, 2010). Some nematode species inhabit thiobiotic environments (Wetzel et al., 2001; Ott et al., 2004, 2005), but no such evidence has been reported for copepods.

Due to the often rapid onset of anoxia and its sometimes short duration, well-designed experiments are needed to distinguish between still living and recently dead meiofauna at the time of sampling. This is even more important for differentiation between sensitive and tolerant species within the respective taxa. Based on initial successes with the CTG labelling method in foraminiferal studies (Bernhard et al., 2006; Pucci et al., 2009), we applied this method to another set of meiofauna organisms, namely copepods and nematodes. Here we test whether the CTG labelling is a more accurate method than the widely used RB staining for quantifying living meiofauna in hypoxia/anoxia studies. The primary
goal of the work described here was to compare harpacticoid copepod and nematode density in normoxic sediment samples with those in anoxic samples. Anoxia was experimentally induced by means of an underwater chamber (Stachowitsch et al., 2007, Riedel et al., 2013). A significant difference in meiofauna density between the two staining techniques could point to a major influence of the staining used, and thus on the final interpretation of ecological studies on meiofauna. Therefore this methodological test on both major meiofauna taxa (copepods, nematodes) will yield well-grounded advice for future staining procedures in meiofauna research.

2 Material and methods
2.1 Experimental set-up
The experiment was performed in the northern Adriatic Sea (Mediterranean) under the oceanographic buoy of the Marine Biology Station Piran (45º32.90’ N, 13º33.00’ E) in 24 m depth on a poorly sorted silty sand bottom. Artificial hypoxia and anoxia was created with a underwater chamber (50 × 50 × 50 cm), originally designed to document the macrofauna epifauna and infauna behaviour during oxygen decline. The separate lid houses a time-lapse camera (images taken in 6 min intervals), 2 flashes, battery packs, a microsensor array (dissolved oxygen, hydrogen sulfide, temperature) plus datalogger (Unisense©). The sensors were positioned 2 cm above the sediment; measurements were taken in 1 min intervals. pH was recorded at the beginning and end of the deployment with a WTW TA 197 pH sensor. For more details on the method see Stachowitsch et al. (2007). For this experiment, the plexiglass chamber plus lid was positioned at the bottom without abundant macroepifauna or without major traces and structures such as mounds or pits that would indicate the presence of larger infauna species. One ascidian and 2–3 brittle stars were placed on the sediment inside the chamber to better visually follow and verify the oxygen decline based on their behaviour (Riedel et al., 2008). To promote the oxygen decline (consumption), 2 additional ascidians were put into mesh bags hanging from the lid.

The chamber was deployed for 5 days from 8–12 August 2009. Dissolved oxygen concentration (initial value ca. 5.5 mL L⁻¹) steadily dropped and reached beginning hypoxia (2 mL L⁻¹) in 12 h, and anoxia in 48 h after chamber closure. At the transition from severe hypoxia (0.5 mL L⁻¹) to anoxia, almost all epifaunal brittle stars were dead and the sediment started to turn greyish black due to the development of hydrogen sulfide (H₂S; final value ca. 30 µmol L⁻¹) reacting with reduced Fe to form FeS. During ongoing anoxia, one sipunculid and two infaunal sea urchins, Ova canaliferus (Fig. 1), emerged from the sediment. The temperature within the chamber remained constant (20.1 °C); the bottom water salinity was 38 PSU. The average pH dropped from 8.1 to 7.6 by the end of the experiment.

Meiofauna cores (inner diameter 4.6 cm) were taken by scuba divers. Eight cores were randomly taken outside the chamber (normoxia control treatments) and another eight cores were taken inside the chamber before the experiment was terminated (anoxia) (see Fig. 1). All cores were immediately transported to the laboratory in cooling boxes and

Fig. 1. Image of sediment surface enclosed by the benthic chamber at the beginning of the deployment (a) and after 5 days of oxygen decline (b), before taking core samples. Note dark colour of sediment compared to outside; sensors visible in corners. Centre: the ascidian Phallusia mammilata, dead and partly overturned brittle stars Ophiothrix quinquemaculata plus datalogger (Unisense©). The sensors were positioned 2 cm above the sediment; measurements were taken in 1 min intervals. pH was recorded at the beginning and end of the deployment with a WTW TA 197 pH sensor. For more details on the method see Stachowitsch et al. (2007). For this experiment, the plexiglass chamber plus lid was positioned at the bottom without abundant macroepifauna or without major traces and structures such as mounds or pits that would indicate the presence of larger infauna species. One ascidian and 2–3 brittle stars were placed on the sediment inside the chamber to better visually follow and verify the oxygen decline based on their behaviour (Riedel et al., 2008). To promote the oxygen decline (consumption), 2 additional ascidians were put into mesh bags hanging from the lid.

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Fig. 2. Representative sample of CTG-labelled harpacticoid copepods. Left: strongly fluorescing (living) specimens, right: weakly stained (dead) individuals.
transferred into a thermostatic room with in situ temperature. The cores were sliced at 0.5 cm intervals until 2 cm depth (and below this in 1 cm steps down to 5 cm depth). The sliced sediment was placed into separate 250 mL containers. For this study, 6 anoxia cores (3 stained with RB and 3 with CTG) and 6 normoxia cores (3 stained with RB and 3 with CTG) were analysed down to 2 cm depth. The two remaining cores were taken as backup cores.

2.2 Labelling/staining protocol

CellTracker Green 5-chloromethylfluorescein diacacetate (CellTracker™ Green CMFDA; Molecular Probes, Invitrogen Detection Technologies) is a fluorescent probe. When living cells are incubated in CTG, the probe passes through the cellular membrane and reaches the cytoplasm, where hydrolysis with nonspecific esterase produces the fluorogenic compound which is observed with the accurate excitation (492 nm) and emission wavelength (517 nm). Unlike other fluorogenic substances, Cell-Tracker Green CMFDA does not leak out of the cell via ion channels in the cell membrane once it is incorporated (Bernhard et al., 2006; Pucci et al., 2009).

Rose bengal (4,5,6,7-tetrachloro-2′,4′,5′,7′-tetraiodofluorescein disodium salt) is a widely used bulk stain (Higgins and Thiel, 1988; Somerfield et al., 2005) that adheres to (cytoplasmatic) proteins, regardless of whether the cell/animal is dead or alive (Bernhard et al., 2006). It colours different meiofauna organisms differently (Giere, 2009) and has the advantage of being cheap and easy to apply, and only a light microscope is required; on the negative side it is a nonvital stain.

The overlying seawater from the cores (10 mL) was added into each 250 mL container including the sediment slices. For CTG labelling, 1 mg of CTG (stored at −20 °C) was dissolved in 1 mL dimethyl sulfoxide (DMSO). The final concentration of CTG/DMSO solution was ≈ 1 µM, corresponding roughly to ≈ 5 µL CTG/DMSO per 10 mL of sediment and liquid together. Overall, 10 µL of the CTG/DMSO solution was added to the sediment-overlying water with a micropipette. The containers were incubated at in situ temperature in the dark for 12 h. Afterwards, samples were fixed in 4 % borax-buffered (5 g L⁻¹) formalin. For RB staining, the RB was applied as 1 g L⁻¹ solution (in 10 % formalin) into fixed samples (4 % borax-buffered formalin) to yield a final 1 % solution (Higgins and Thiel, 1988; Somerfield et al., 2005).

2.3 Meiofauna extraction and counting

The samples were processed following the common meiofauna protocol (De Jonge and Bouwman, 1977; McIntyre and Warwick, 1984). Formalin-fixed sediment samples were washed with tap water to eliminate formalin and clay by pouring them on a 38 µm sieve. The sediment recovered on the 38 µm sieve was transferred into 1 dL centrifuge tubes and a Levasil® (distilled-water) solution (specific density = 1.17 g cm⁻³) was added and gently mixed through the sediment prior to centrifugation. After centrifugation, the soft-bodied meiofauna was retained in the floating phase. Copepods and nematodes were counted using a Nikon SMZ 800 binocular microscopes equipped with a Nikon INTENSILIGHT C-HGFI for UV production and an A 488 filter. CTG samples containing a large number of animals (surface layers of normoxic sediment) were split into several petri dishes to avoid long exposure of animals to UV light in order not to lose fluorescence.

2.4 Data treatment

Analyses were performed using the R statistical software package (Team, 2010). The nematode and copepod data were first tested for normality with Lilliefors (Kolmogorov–Smirnov) test using the “nortest” package. The nematode abundance data followed the normal distribution, whereas copepod abundance data were significantly different from normal and showed a positively skewed distribution; therefore the data were log-transformed for the calculations of three-way factorial ANOVA (Dytham, 2003; Sokal and Rohlf, 1995). The three-way factorial ANOVA was calculated for copepods and for nematodes separately to test for significant differences (p < 0.05) among three factors: method (CTG vs. RB), oxygen (anoxia vs. normoxia) and sediment layer (0–0.5 cm, 0.5–1 cm, 1–1.5 cm, 1.5–2 cm vertical sediment depth). The relationships of first-order interactions (oxygen X method, oxygen X depth and depth X method) were plotted on the graphs, where the third factor (not shown) was averaged (Sokal and Rohlf, 1995).

3 Results and discussion

The experimental design (Stachowitsch et al., 2007) successfully mimicked the hypoxic and anoxic conditions and associated meiofaunal responses in the expected time frames. For the first time, the CTG technique was applied specifically to label two main groups of meiofauna, copepods

<table>
<thead>
<tr>
<th>Copepods</th>
<th>DF</th>
<th>Sum Sq</th>
<th>Mean Sq</th>
<th>F value</th>
<th>Pr (&gt; F)</th>
</tr>
</thead>
<tbody>
<tr>
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<td>95.09</td>
<td>31.7</td>
<td>298.302</td>
<td>0.000000</td>
</tr>
<tr>
<td>Oxygen</td>
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<td>5.5</td>
<td>5.5</td>
<td>51.805</td>
<td>0.000000</td>
</tr>
<tr>
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<td>4.9</td>
<td>4.9</td>
<td>46.089</td>
<td>0.000000</td>
</tr>
<tr>
<td>Layer: oxygen</td>
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<td>1.83</td>
<td>0.61</td>
<td>5.734</td>
<td>0.005680</td>
</tr>
<tr>
<td>Layer: method</td>
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<td>0.31</td>
<td>2.88</td>
<td>0.050860</td>
</tr>
<tr>
<td>Oxygen: method</td>
<td>1</td>
<td>2.52</td>
<td>2.52</td>
<td>23.732</td>
<td>0.000027</td>
</tr>
<tr>
<td>Layer: oxygen: method</td>
<td>3</td>
<td>0.29</td>
<td>0.1</td>
<td>0.895</td>
<td>0.453810</td>
</tr>
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<td>Residuals</td>
<td>33</td>
<td>3.51</td>
<td>0.11</td>
<td></td>
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</tr>
</tbody>
</table>
and nematodes, in an attempt to document the course of an oxygen deficiency event more precisely. Especially in modern, short-term, process-oriented studies, definitive determinations of living and dead components are essential. This approach is much more satisfactory and fine scaled than comparing samples from different times and distinguishing communities based on presence/absence.

3.1 Meiofauna densities

The copepods, as the most sensitive meiofauna group to oxygen depletion (Hicks and Coull, 1983; Moore and Bett, 1989; Moodley et al., 1997), were of primary interest here. We found a significant drop in copepod density from normoxia to anoxia (factor oxygen, Table 1) regardless of the staining/labelling technique (Fig. 3). Moreover, the density was significantly higher in the surface layers with both techniques, in any sediment depth layers. The density of RB stained copepods was nearly 2 times higher in the top layer (56 ± 16.5 vs. 102 ± 8.2 for CTG and RB, respectively) and 5 times in the lower layers (0.5–1 cm, 1–1.5 cm, 1.5–2 cm) of anoxia (Fig. 3). This explains the significant interaction of the factors oxygen and method as the method was relevant only in anoxic circumstances (Table 1). In normoxia the copepod abundance was very similar for CTG and RB (Fig. 6a). In anoxia, however, there is the trend of higher copepod abundance in RB versus CTG (Fig. 6a). The three-way ANOVA (Table 1) indicates a significant interaction also for the factors oxygen and layer (demonstrated on Fig. 6b) because there is a huge difference in the copepod abundance between anoxia and normoxia especially in the top sediment layer, but not in deeper layers. The interaction of layer and method was not significant, as the same trend of decreasing copepod abundance with sediment depth was observed for both methods (see parallel lines in Fig. 6c).

Although we found significant differences in the density of copepods between normoxia and anoxia already with RB staining (De Troch et al., 2013), the density is greatly overestimated with the RB method. Importantly, the CTG labelled only the animals that were still alive at the time of fixation. Accordingly, with CTG we can extract from the samples only the true surviving species of short-term anoxia (more information in Grego et al., 2013). This provides highly relevant information in connection with impact studies.

The nematode density, as the copepod density, was also significantly affected by each factor separately (method, oxygen and sediment layer, Table 2, Fig. 5). When analysing the impact of oxygen (normoxia/anoxia) solely on the RB-stained nematodes, De Troch et al. (2013) did not find a significant effect, mainly due to high variation of nematode abundance, when stained with RB. When the CTG-labelled nematode densities are integrated in three-way ANOVA calculations (in the deeper layers of anoxia the CTG densities are much lower than RB) the factor oxygen becomes significant. The interaction of factors oxygen and method was not significant (Table 2, Fig. 7a), as regardless of staining method, the density of nematodes slightly dropped from normoxia to anoxia. The interaction of factors oxygen and depth is significant as in normoxia the peak of nematode density is in the second layer (0.5–1 cm), while in anoxia the density decreases linearly with depth (Table 2, Fig. 7b). The application of different staining methods to various sediment layers (layer and method) did not affect the trend of nematode

![Fig. 3. CTG-labelled vs. RB-stained copepod density in normoxia and after experimental anoxia in four sediment depth layers. Average values (±STD) are calculated from three replicates.](image-url)

**Table 2.** Three-way ANOVA for factors method, oxygen and layer (depth of the sediment). The calculation was done on nematode abundances. Significant factors or interactions are bolded ($p < 0.05$).

<table>
<thead>
<tr>
<th>Nematodes</th>
<th>Df</th>
<th>Sum Sq</th>
<th>Mean Sq</th>
<th>F value</th>
<th>Pr (&gt; F)</th>
</tr>
</thead>
<tbody>
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<td>30 103</td>
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<tr>
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<td>29 277</td>
<td>5.235</td>
<td>0.028670</td>
</tr>
<tr>
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<td>216 395</td>
<td>38.697</td>
<td>0.000001</td>
</tr>
<tr>
<td>Layer: oxygen</td>
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<td>27 222</td>
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<td>0.006520</td>
</tr>
<tr>
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<td>0.124</td>
<td>0.945170</td>
</tr>
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<td>0.437560</td>
</tr>
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<td>1778</td>
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<td>0.812280</td>
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<tr>
<td>Residuals</td>
<td>33</td>
<td>184 535</td>
<td>5592</td>
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density, which is generally decreasing with depth (parallel lines—Table 2, Fig. 7c).

In contrast to what we found for copepods and what Bernhard et al. (2006) found for foraminiferans, where the RB significantly overestimated the abundances in anoxia, for nematodes it was not so, at least not in all sediment layers (Fig. 5). There were no differences in nematode abundance if labelled with CTG or stained with RB in top anoxic layer. The RB overestimated the abundances in the top sediment layer of normoxia (139.3 ± 24.0 vs. 300.8 ± 101.1 for CTG and RB, respectively) (Fig. 5). Also in the 1–1.5 cm layer of normoxia the abundances were significantly higher with RB. The presence of lower numbers of live – CTG-labelled – nematodes in top normoxic layer might reflect the short generation time of some species, namely days to weeks (Platt et al., 1985; Martinez et al., 2012). Even though the decomposition is relatively fast (Kammenga et al., 1996), nematodes do not disintegrate within hours/days; therefore the density of living nematodes may be overestimated in ecological/monitoring studies based solely on RB staining. As RB stains all material containing proteins, the presence of different material on, for example, the cuticula of nematodes can contribute to an overestimate of living organisms. This finding has important implications for future counts based on RB-stained samples. Also the variation of the nematode abundance between replicates is much higher for nematodes stained with RB, especially in deeper sediment layers of anoxic treatment (Fig. 5). This implies that the CTG method is more accurate. This high variation with the RB method could be due to the presence of many already dead but not yet decomposed nematodes in RB anoxic samples. In contrast, the abundance of survivors is similar in all CTG-labelled cores.

Oxygen is the most important abiotic variable to determine the vertical distribution of meiofaunal organisms (Gray and Elliot, 2009); the same holds true in oxygen minimum zones, where the copepod density was positively correlated with oxygen concentration (Levin et al., 2002). Nematodes, in contrast, were shown to be most tolerant to anoxic stress (Josefson and Widbom, 1988; Murrell and Fleeger, 1989; Hendelberg and Jensen, 1993; Moodley et al., 1997; Modig and Olafsson, 1998; Levin, 2003). The mortality and depth distribution of nematodes depend mainly on the length of anoxia, and can drop markedly, even by a third already after a 14-day exposure to anoxia (Steyaert et al., 2007). Some nematode species, however, are known to inhabit anoxic layers (Soetaert and Heip, 1995; Steyaert et al., 2007) and show a negative correlation with oxygen concentrations (Levin et al., 2002). In our shorter-term experiment the overall nematode density did not show a major decrease at anoxia. Rather, we detected an accumulation of individuals in the top layer and lower densities in the deeper layers with both staining techniques (Figs. 5, 7b). We attribute this to an upward migration of nematodes towards the sediment surface under anoxic conditions, as reported elsewhere (Hendelberg and Jensen, 1993).

Sutherland et al. (2007) pointed to an important effect of free sulfide that is formed in anoxic conditions. They found weak correlations between free sulfide concentration and the number of nematodes, while clear trends occurred between free sulfide and numbers of crustaceans. The crustaceans were represented mainly by copepods, and with increasing concentration of free sulfide the number of copepods dropped. This is in accordance with our study. The free sulfide concentrations rose up to 30 μmol L⁻¹ in
anoxia (in the water column), which is above the level causing significant negative effects on marine benthic organisms (> 14 µmol L\(^{-1}\); Vaquer-Sunyer and Duarte, 2010). The small influence of sulfide on nematodes can be explained by adaptive strategies. The nematode community inhabiting the deeper sediment layers (0.5–2 cm) is therefore already adapted to sulfide (Mirto et al., 2000), while copepods are not adapted, as they are mainly abundant in the top layer (0–0.5 cm), which is better oxygenated. The muddy sediment of the northern Adriatic Sea often lacks oxygen and contains high concentrations of free sulfide during warm periods with strong water stratification (Faganeli et al., 1991; Hines et al., 1997).

Finally, the mortality of nematodes during sample handling may play a role. Such mortality is indicated by a significant difference between RB and CTG techniques in nematode numbers in normoxic samples, but no such difference was found for copepod numbers in normoxic samples. Nematodes may be more sensitive to physical disturbance during sediment slicing and sediment mixing while applying the CTG label. Also, some species from the northern Adriatic muddy community can be sensitive to high oxygen concentrations during sample processing. Wetzel et al. (2001) and references therein state that nematodes from poorly oxygenated muddy sites can be intolerant to well-oxygenated interstitial waters.

### 3.2 Labelling/staining implications

As in all staining techniques, staining is seldom an all-or-nothing phenomenon. The copepods and nematodes stained with RB showed different intensity in colouration, but only the empty exuvia of copepods or unstained nematodes could be definitely determined as dead. When labelled with CTG, some organisms shined brightly green, and were classified as living at the time of sampling (Fig. 2 left, Fig. 4 left). Another set of individuals coloured only minimally – pale green – and were classified as dead at the time of sampling (Fig. 2 right, Fig. 4 right). The difference between the live and dead animals is similar to that outlined for foraminiferans by Pucci et al. (2009) and for loriciferans by Danovaro et al. (2010). Loriciferans that were frozen prior to the CTG labelling (and therefore dead; the enzymatic activity was inhibited and CTG could not label them) coloured pale green (Danovaro et al., 2010). It was helpful to pull all individuals together to simplify the distinction between these two levels of colouration. The intensity of the staining (RB) or fluorescence (CTG) varied also with different species of copepod or nematode. Elliott and Tang (2009) faced similar problems while staining with neutral red, also a vital stain. Moreover, bigger species absorbed more stain, as already reported by Elliott and Tang (2009).

RB staining is a classical procedure in benthic ecology (Elefteriou and McIntyre, 2005; Giere, 2009) and stains proteins that can be preserved well in dead animals for lengthier periods (Murray and Bowser, 2000). The CTG method, in contrast, enables the detection of actively metabolizing cells as it identifies the presence of respiratory activity (Bernhard et al., 2006; Pucci et al., 2009). The CTG technique, however, can have certain peculiarities. The first involves the duration of post-mortem enzymatic activity (Borrelli et al., 2011). The residual esterase activity in microbes produced around 25 % of fluorescence after heat killing, still 48 h after treatment (Kaneshiro et al., 1993). At the same time, foraminifera killed 1 h prior to staining did not exhibit any fluorescence (Bernhard et al., 1995), indicating only short-lived post-mortem esterase activity. The latter is currently

![Fig. 6. First-order interactions of three-way ANOVA for factors (a) oxygen X method, (b) oxygen X depth and (c) depth X method. The geomean of copepod abundance is given. For each graph, the third factor (that is not shown) was averaged.](http://www.biogeosciences.net/10/4565/2013/)
unknown in copepods, but Bergvik et al. (2012) recently showed that planktonic calanoid copepods exhibited activity of autolytic enzymes soon after death, speeding up their decomposition. We therefore assume that vital functions, necessary to activate the fluorescent probe CTG, are diminished soon after death. The second feature of the CTG technique is that it also labels bacterial stocks (Pucci et al., 2009). In our samples, for example, the empty exuvia (carapace) of the copepods were not labelled, but the area of joints was shiny due to bacterial aggregation here (i.e. the largest copepod individual on the top right side in Fig. 2); this did not affect the distinction between dead and living individuals. The third CTG feature to consider is that the working time must be rationalized because CTG fluorescence slowly fades with working time under UV light. Finally, although Danovaro et al. (2010) incubated the Loriciferans with CTG probe at sea level pressure, Elliott and Tang (2009) suggest that such live stains/labels should not be applied for deep-sea animals or other animals inhabiting specific environments whose variables cannot be reproduced in the laboratory, causing the animals to die during incubation.

The examination of living fauna (Steyaert et al., 2007) prior to fixation is clearly one alternative to CTG. However, the determination of live copepods or nematodes can be misleading here as well: individuals can be immobile but actually still alive. Moreover, the time needed to examine the living fauna in multiple replicated and big samples is typically too long to treat all samples equally.

4 Conclusions

In conclusion, our results show that the CTG technique well distinguished living from dead organisms and is a suitable staining method to document the response of selected meiofauna groups to short-term disturbances. The common trend is that RB markedly overestimates the number of survivors in both taxa. Specifically, the CTG technique was able to pinpoint a turning point in copepod density from normoxia to anoxia, whereas traditional RB staining failed to resolve this event properly. When the copepods and nematodes are dead, but not yet decomposed, the RB will stain them equally as live ones, especially if they are freshly dead. Therefore, copepods and nematodes that died in the course of the experiment can only be enumerated, and excluded, with CTG labelling.

Importantly, CTG helps to better determine more tolerant species that survived the respective level of disturbance. We therefore conclude that for normal monitoring of meiofauna diversity under undisturbed conditions, RB is an efficient method. If the task is to examine survival of animals in hypoxia, anoxia or other types of disturbance, and a binocular microscope equipped with proper UV production light and filter is available, then we highly recommend the CTG technique. This echoes the recommendation of foraminiferan researchers.
References


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