Impact of farming non-indigenous scallop *Argopecten irradians* on benthic ecosystem functioning: a case-study in Laizhou Bay, China

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ABSTRACT: The farming of the non-indigenous bay scallop *Argopecten irradians* in coastal waters generates large amounts of biodeposits that potentially change the trophic pathways and quality of the benthic food web at lower trophic levels such as meiobenthos. To understand the trophic link between faecal pellets of bay scallop and meiobenthos in the aquaculture area, we investigated the resource use of harpacticoid copepods and nematodes inside and outside of 3 bay scallop farms in Laizhou Bay (Bohai Sea, China) using natural abundance of stable carbon and nitrogen isotopes together with fatty acid profiling. Faeces were found to be enriched in $\delta^{15}$N compared to all other food sources, which made faecal matter traceable. The enriched $\delta^{15}$N in several meiobenthos at the farms, together with the mixing model results, indicated that faeces could be a new food source for most of harpacticoid copepods and some nematodes. The quantities and the pathways of assimilation differed between the copepod families, depending on their feeding behaviors and the receiving environment. Furthermore, due to the presence of higher levels of polyunsaturated fatty acids, in particular docosahexaenoic acid, the dominant copepod family Canuellidae that abundantly consumed scallop faeces showed enhanced nutritional quality compared with those in the control sites. Thus, aquaculture of non-indigenous bay scallops provided a food source that was directly and indirectly consumed by meiobenthos underneath the scallop farms and improved the quality of lower level consumers as a food item in the benthic food web.

KEY WORDS: Mariculture · Non-indigenous species · *Argopecten irradians* · Harpacticoid copepods · Nematodes · Food source · Stable isotopes · Fatty acids
change the benthic energy flow (Peterson & Heck 1999, Callier et al. 2008). Although many studies have recorded the influence of biodeposition on benthic assemblages (Hartstein & Rowden 2004, Callier et al. 2008), little is known about the fate of biodeposits and their functional effect on the benthic ecosystem, e.g. through the assimilation of these deposits by benthic organisms. While biodeposits have a good nutritional value (McKindsey et al. 2011), it is unclear whether the benthos can benefit from incorporating these deposits into their diet.

China is the biggest shellfish aquaculture producer worldwide and scallop farming using longlines in coastal marine waters is a major part of its aquaculture industry (FAO 2016). Among the cultured scallop species, the non-indigenous species (NIS) Argopecten iradians (a.k.a. bay scallop), introduced from North America, has dominated the Chinese scallop production over the past 30 yr (Guo & Luo 2016). In terms of economic output, A. iradians has an advantage over the native scallop Chlamys farreri due to the faster growth rate (Guo & Luo 2016). However, the high biodeposition rate of A. iradians (almost 10 times higher than C. farreri) also poses a threat to the benthic ecosystem (Zhou et al. 2006, Li et al. 2009, Wang 2015). However, little is known about whether and how the biodeposition of non-indigenous A. irradians can potentially affect organisms in the sediment and the overall functioning of the benthic ecosystem.

Ecosystem functioning integrates the energy flux within a system (Power 1992). The process of resource utilization by benthic consumers, especially meioiobenthos (the smaller fraction of metazoans passing through a 1 mm sieve but being retained on a 38 μm sieve), is crucial to understand the energy flux of an ecosystem, because: (1) meioiobenthos are highly abundant and form a link between primary producers and higher trophic levels (TLs) (Leduc et al. 2009); and (2) due to their small size, short generation times and close associations with sediments, they are sensitive to stressors and respond functionally to them (Kennedy & Jacoby 1999). However, there is little information on functional responses (in particular, resource utilization) of meioiobenthos to aquaculture biodeposition. Since the effects of aquaculture waste are known to differ among trophic guilds and feeding behaviors of animals (Wai et al. 2011), it is important to incorporate the knowledge of functional responses of different taxonomic groups in order to understand the impact on ecosystem functioning. Furthermore, harpacticoid copepods are important food sources and providers of fatty acids (FAs), especially highly polyunsaturated fatty acids (PUFAs), to higher marine consumers (de Lima et al. 2013). Their FA profiles depend on their food sources and environmental conditions (Nanton & Castell 1999, De Troch et al. 2012, de Lima et al. 2013). Determining the presence of quality-indicator FAs, such as PUFA, 20:5ω3 (eicosapentaenoic acid; EPA) and 22:6ω3 (docosahexaenoic acid; DHA), in harpacticoid copepods will contribute to the assessment of the functional impacts of biodeposition by bay scallop in aquaculture areas.

The analysis of the natural abundance of stable isotopes together with FA profiles is an efficient tool to investigate the diet of meioiobenthos (Leduc & Probert 2009, Cnudde et al. 2015). Stable carbon ratios reflect the food sources of consumers, and nitrogen ratios are used to determine their trophic positions (DeNiro & Epstein 1978, Minagawa & Wada 1984). Moreover, stable isotopes help to trace the fate of aquaculture waste in different communities (Gondwe et al. 2012, Vizzini & Mazzola 2012, Sanz-Lázaro & Sanchez-Jerez 2017). FA profiles provide information on food sources such as diatoms and bacteria (Kelly & Scheibling 2012).

The aim of this study was to evaluate the impact of aquaculture of the NIS bay scallop A. iradians on the benthic ecosystem functioning. Specifically, we used stable isotopes and FAs to test the following hypotheses: (1) the presence of a scallop farm affects the isotopic values of primary organic sources in the sediment; (2) scallop faeces are consumed by meioiobenthos in the receiving sediment; (3) the quantity of faecal consumption differs among taxa/families of meioiobenthos; and (4) the presence of a scallop farm changes the FA profile of the harpacticoid copepods occurring in the sediment.

**MATERIALS AND METHODS**

**Study area**

Samples were collected in scallop farms located in the eastern part of the Laizhou Bay (37°00’ to 38°30’ N, 118°45’ to 120°30’ E). Laizhou Bay is located in the south Bohai Sea, Shandong Province, on the north coast of China. It has a mean depth of 9 m (maximum ~18 m), a coastline of 320 km and a total area of ~700000 ha (Zhuang et al. 2014). The northeast–southwest currents dominate with a mean velocity of 20 cm s⁻¹, indicating a poor exchange of water (Zhao & Chen 2001). The area is an important spawning and breeding ground for many fishes but is currently under pressure due to human activities such as intensive mariculture (Jin et al. 2013).
The bay scallop *Argopecten irradians* is a major mariculture species, which has been cultivated using the suspended-longline method in this area for 30 yr. Bay scallop mariculture in Laizhou Bay is one of the most important sources of scallop production in China, with a total area of 500 ha and a density of 200 ind. m$^{-2}$ yielding a production of around 20 000 t yr$^{-1}$ (Liu et al. 2004, Wang 2015). Bay scallops are cultured from May to November each year without adding any feed. The rest of the year, the area is devoid of any aquaculture activities.

**Sampling design**

We selected 3 of the largest farms and 3 corresponding control sites. To exclude any direct effect of aquaculture on the control sediment, each control site was located 2 km to one side of the farm, at the same depth as the farm, so that it was exposed laterally to the predominant current that also flowed through the farm. Each pair of sites, ‘farm’ and ‘control’, was considered as a single station, and the 3 stations were characterized by sediment type as sandy-shallow (Stn SS) (2% clay, 24–28% silt, 69–73% sand; water depth 5 m), muddy-shallow (Stn MS) and muddy-deep (Stn MD) (4–6% clay, 53–55% silt, 39–42% sand; water depth 7.5 and 12 m respectively). Previous observations in Laizhou Bay revealed that biodeposition rates of bay scallops peaked in November, and rates of larger individuals were higher than those of smaller ones (Wang 2015). We thus conducted sampling for stable isotopes and FA analysis in October–November 2015 and 2016, before the harvest of scallops, in order to cover the period of high biodeposition rates in this area. For FA samples, an additional sampling campaign was conducted in May 2016 before the scallop lantern nets were placed in this area.

**Sampling procedure**

We considered that meioobenthos from each farm and control site were exposed to the following common primary organic sources: (1) phytoplankton from the water column, (2) microphytobenthos (MPB) in the sediment surface, and (3) fragments of seagrass *Zostera marina* leaves in the sediment (senescent fragments of *Z. marina* verified by microscopical observation). Bay scallop faeces were only considered as a potential food source at the farm sites. Phytoplankton was considered to be the major component in the pre-filtered particulate organic matter (POM) because the weight-to-weight ratio of particulate organic carbon to chlorophyll $a$ (POC/chl $a$) ranged from 19 to 37 in October-November in our sampling area (Wang 2015). This is within the known range of 2 to 200 for POC/chl $a$ in algae (Cifuentes et al. 1988). POM samples were obtained at each control site in order to avoid the potential addition of aquaculture-derived organic matter. About 500 ml seawater was filtered through a 58 μm net to remove zooplankton and large detritus, and subsequently filtered on pre-combusted (450°C, 4 h) 0.7 μm Whatman GF/F glass fiber filters. MPB samples were collected at each farm and control sites in 2016. Due to the insufficient MPB biomass for reliable stable isotope analysis, we pooled farm and control together for each station, which is justified by the similar $\delta^{13}C$ and $\delta^{15}N$ values of the surface sediment at each farm and control site (see ‘Data analysis’ below). MPB were separated from the sediment surface using a modified method of Doi et al. (2003): the top 1 cm sediment collected by a Van Veen grab was covered by a 2 mm layer of quartz sands (25 to 65 μm diameter, pre-combusted at 500°C for 2 h), a nylon net (75 μm), and another layer of 2 mm pre-combusted quartz sands. The dishes were illuminated for 24 h, while moisture was maintained with continuous spraying of filtered seawater on the sand. After illumination, the upper sand layer was scraped off and resuspended in filtered seawater. The supernatants were filtered on pre-combusted Whatman GF/F filters. We also collected sediment surface organic matter (SSOM) to evaluate the impact on the organic pool of surface sediments. SSOM was collected at each farm and control site by scraping the top layer (0 to 1 cm) of sediment from the Van Veen grab. To collect the bay scallop faeces, 20 individuals of scallop (average body length of 5.5 cm) were placed in the containers with filtered seawater overnight and faeces were obtained by filtering the water from the containers on pre-combusted GF/F filters. Samples ($n = 2$ to 4) were taken for POM, MPB, SSOM and scallop faeces. Meiothentos for stable isotopes and FAs were collected qualitatively by scraping the top 1 cm layer of the sediment from 3 randomly selected patches (approx. $1 \times 1$ m) that were at least 1 m apart at the farm and control sites.

**Stable isotope analysis**

The POM, MPB, and faeces samples were oven-dried at 60°C to a constant weight and divided into 2
subsamples. One set was treated with hydrochloric acid (HCl) fume to remove carbonates for \( ^{13}\text{C} \), and the other, without acid treatment, was used for \( ^{15}\text{N} \) analysis. All samples were put into tin capsules (Elemental Microanalysis, 8 x 5 mm) and pinched closed. 30 to 60 mg of sediment was acidified in silver capsules (Elemental Microanalysis, 8 x 5 mm) with diluted HCl to remove carbonates and washed with distilled water. The capsules were dried and pinched closed afterwards.

Sediments were sieved with filtered seawater through a 500 μm and a 150 μm sieve. The fraction retained on the 150 μm sieve was frozen and transported to the lab. The frozen samples were thawed and meiobenthos were handpicked with a needle under a stereomicroscope. Meiobenthos were sorted into nematodes (150 to 300 ind. per sample) and copepods. For the samples collected in 2015, copepods were pooled together (80 to 100 ind. per sample) while copepod samples in 2016 were sorted to family level (80 to 120 ind. per sample). Meiobenthos were rinsed with MilliQ water twice before being transferred to tin capsules, oven-dried overnight at 60°C and pinched closed. All samples were stored in a desiccator prior to further analysis.

C and N stable isotope ratios were measured with an isotope ratio mass spectrometer (type Europa Integra) at UC Davis Stable Isotope Facility (University of California). Isotope values are expressed as \( \delta^{13}\text{C} \) and \( \delta^{15}\text{N} \) (%) determined by the following equation:

\[
\delta^{13}\text{C or } \delta^{15}\text{N} = \left( \frac{R_{\text{sample}}}{R_{\text{standard}}} - 1 \right) \times 10^3 \tag{1}
\]

where \( R = {^{13}\text{C}}/{^{12}\text{C}} \) or \( {^{15}\text{N}}/{^{14}\text{N}} \). Reference standards were PDB for \( \delta^{13}\text{C} \) and atmospheric nitrogen for \( \delta^{15}\text{N} \).

### Fatty acid profiling

FA samples included only the omnivorous family Canuellidae collected at Stn SS. The abundances of the rest of the meiobenthos at the other sites were too low to obtain enough biomass for a reliable analysis. Canuellidae were extracted alive from sediments following the method of Svensson et al. (2010) and stored at room temperature overnight to clear their gut content. The next day, 100 to 150 individuals were picked, washed with filtered seawater, and transferred to glass tubes for storage at \(-80^\circ\text{C}\) prior to FA extraction.

Lipid extraction, methylation to fatty acid methyl esters (FAMEs), and FAME analysis followed the procedure of De Troch et al. (2012). FAME of 19:0 (Fluka 74208) was added as internal standard. The FAMEs were analyzed with a gas chromatograph (HP 6890N) coupled to a mass spectrometer (HP 5973). FAMEs were identified by comparing the retention time and mass spectra with authentic standards and mass spectral libraries (WILEY, NITS05) and then analyzed with the software MSD ChemStation (Agilent Technologies). Individual FAMEs were quantified by using a component FAME and BAME mix (Supelco #47885 and #47080 respectively, Sigma-Aldrich) and additional standards (Larodan). Shorthand FA notations were expressed as A:BωX, where A gives the number of carbon atoms, B represents the number of double bonds and X is the position of the first double bond closest to the terminal methyl group.

### Data analysis

Variations in \( \delta^{13}\text{C} \) and \( \delta^{15}\text{N} \) values of potential food sources, SSOM, and meiobenthos were tested using analysis of variance (ANOVA) followed by Student-Newman-Keuls pairwise comparisons. Prior to ANOVA, the assumption of homogeneity of variances were diagnosed with Kolmogorov-Smirnov tests and Levene’s tests, respectively. Log transformations were used to meet this assumption if necessary. Non-parametric Kruskal-Wallis tests were conducted when homogeneity of variation could not be reached. Differences between farm and control at each station for \( \delta^{13}\text{C} \) and \( \delta^{15}\text{N} \) values of SSOM, TL, and FA indicators were assessed using t-tests. ANOVA and t-tests were conducted with the software SPSS 20.0.

To identify the differences in isotopic values (\( \delta^{13}\text{C} \) and \( \delta^{15}\text{N} \)) of meiobenthos between farm and control, the \( \delta^{13}\text{C} \) and \( \delta^{15}\text{N} \) ratios of taxon or family co-occurring in farm and control sites were plotted in the same biplot (i.e. \( \delta^{13}\text{C} \) control/\( \delta^{13}\text{C} \) farm and \( \delta^{15}\text{N} \) control/\( \delta^{15}\text{N} \) farm). Taxa or families were considered to have similar \( \delta^{13}\text{C} \) or \( \delta^{15}\text{N} \) values if the ratio fell within the 95% confidence interval (CI) encompassing the 1:1 correlation between farm and control isotopic values.

The TL of meiobenthos was estimated based on \( \delta^{15}\text{N} \) values:

\[
\text{TL}_{\text{consumer}} = 2 + (\delta^{15}\text{N}_{\text{consumer}} - \delta^{15}\text{N}_{\text{base}})/\Delta^{15}\text{N} \tag{2}
\]

where \( \delta^{15}\text{N}_{\text{base}} \) is the \( \delta^{15}\text{N} \) of primary consumer i.e. the family of harpacticoid copepods that has the lowest \( \delta^{15}\text{N} \) in each site. The \( \Delta^{15}\text{N} \) of 2.3% for marine organisms was adopted (Zanden & Rasmussen 2001).
A mixing model MixSIR applied Bayesian method was used to calculate the potential food sources of meiobenthos (Moore & Semmens 2008). Trophic enrichment factors (TEF) of $0.3 \pm 1.3\% \delta^{13}C$ and $2.3 \pm 1.8\% \delta^{15}N$ were adopted for each trophic step (Zanden & Rasmussen 2001). The isotopic values of the seagrass Z. marina were adopted from Hoshika et al. (2006). We calculated isotopic data only for the year 2016 because MPB was not collected in 2015. MPB and POM were pooled as a logical group representing the microalgae-derived organic matter (Phillips et al. 2005).

A 2-way permutational multivariate analysis of variance (PERMANOVA, main test and pairwise test) and analysis of similarity (ANOSIM) were run on the relative FA profiles of Canuellidae in Stn SS in 2016. Time (‘Before vs. During’, where ‘Before’ refers to the period prior to the start of seasonal aquaculture production) and site (‘Farm vs. Control’) were orthogonal and fixed factors. A distance-based test for homogeneity of multivariate dispersions (PERMDISP) was used to test the homogeneity of multivariate dispersion (Anderson 2006). To visualize the degree of dissimilarity of FA composition between groups, non-metric multi-dimensional scaling (nMDS) was conducted based on a Bray-Curtis resemblance matrix of untransformed relative FA profiles. The contribution of individual FA to these clusters was tested by similarity percentages (SIMPER) analysis. All multivariate analyses were performed with Primer V6 (Clarke & Gorley 2006), using the PERMANOVA+ add-on package (Anderson et al. 2008).

**RESULTS**

**Stable isotopes of food sources**

Scallop faeces showed enriched $\delta^{15}N$ values ($9.40 \pm 1.03\%$ in 2015 and $9.44 \pm 0.93\%$ in 2016, Fig. 1), which were higher than those of any other potential food source (ANOVA for 2016: $F_{6,13} = 20.999$, $p < 0.001$; 2015: $F_{3,11} = 8.666$, $p = 0.003$). For the $\delta^{13}C$ values, significant differences were found, with decreasing signatures from seagrass (Zostera marina) to scallop faeces to POM and MPB (Fig. 1A). Isotopic composition of POM and MPB varied among stations in terms of $\delta^{15}N$ values in 2016 (ANOVA, POM in 2016: $F_{2,6} = 6.806$, $p = 0.029$; MPB in 2016: $F_{2,5} = 13.900$, $p = 0.009$) (Fig. 1B). Within each station, only Stn MD and Stn SS showed significant differences in isotopic signatures. At Stn MD the MPB had more depleted $\delta^{15}N$ values compared with POM, while there were no differences at SS and MS ($t$-test, SS: $t_{1.074} = 6.157$, $p = 0.091$; MS: $t_4 = 1.622$, $p = 0.180$; MD: $t_4 = 3.715$, $p = 0.021$). At Stn SS $\delta^{13}C$ values of POM
were more depleted than those of MPB, while there were no differences at MS and MD (t-test, SS: \( t_3 = 9.203, p = 0.003 \); MS: \( t_4 = 6.908, p = 0.415 \); MD: \( t_4 = 1.050, p = 0.353 \)). The \( \delta^{15}N \) values of POM at Stns SS and MS were lower in 2015 compared with those in 2016 (Fig. 1B), and Stns SS and MD in 2015 displayed more depleted \( \delta^{15}N \) than in 2016 (\( F_{5,14} = 3.477, p = 0.030 \); post hoc test \( \alpha = 0.014 \) for Stn SS and \( \alpha = 0.023 \) for Stn MS).

**Stable isotopes of SSOM**

Results of t-tests showed there was no significant difference for either \( \delta^{13}C \) or \( \delta^{15}N \) of SSOM between farm and control sites at each station (Fig. 2; see also Table S1 in the Supplement at www.int-res.com/articles/suppl/q010p227_supp.pdf; t-test: \( p > 0.05 \)). The ANOVA test for the \( \delta^{15}N \) values of SSOM at 3 stations in 2015 showed a spatial variation (ANOVA: \( F_{5,12} = 33.338, p < 0.001 \)) with Stn SS having slightly lower \( \delta^{15}N \) values (Fig. 2).

**Stable isotopes and trophic level of meiobenthos**

Copepods and nematodes were the major meiobenthos taxa in all stations, accounting from 8 to 15% and 45 to 92% for meiobenthos biomass respectively (authors’ unpubl. data). For copepods, in total 4 families were identified: Canuellidae, Laophontidae, Ectinosomatidae, and Miracididae. At Stns SS and MS, Canuellidae was the dominant copepod family at both control and farm sites. At Stn MD, copepods were more diverse at the farm site, where Laophontidae, Ectinosomatidae, Miracidae, and Canuellidae were abundant, while at the control site only Laophontidae and Ectinosomatidae were abundant. C and N isotopic values varied spatially and were not always the same in the 2 sampling events, ranging from \(-23.02 \pm 0.76\%\) to \(-19.16 \pm 0.20\%\) for \( \delta^{13}C \) and \( 8.13 \pm 0.13\%\) to \( 14.37 \pm 0.17\%\) for \( \delta^{15}N \) (Table 1). Carbon and nitrogen isotopic values varied among meiobenthos taxa (in 2015, \( \delta^{13}C: F_{12,31} = 31.126, p < 0.001 \); \( \delta^{15}N: F_{12,31} = 9.832, p < 0.001 \); in 2016, \( \delta^{13}C: F_{13,31} = 30.035, p < 0.001 \); \( \delta^{15}N: F_{13,31} = 8.089, p < 0.001 \)). In general, nematodes had higher \( \delta^{15}N \) values compared with copepods from the same site, except for the nematodes from 2015 Stn SS that showed a slightly lower value at the farm and a similar value at the control site (Table 1). For the common copepod family Canuellidae, isotopic values displayed high variation (mean values ranged from \(-21.27 \) to \(-19.16\%\) for \( \delta^{13}C \) and \( 8.13 \) to \( 12.55\%\) for \( \delta^{15}N \) respectively). Family Ectinosomatidae showed the highest \( \delta^{15}N \) among copepods at the farm at Stn MD in 2016.

With regard to TL, nematodes at Stns MD and MS showed higher values than any other meiobenthos, ranging from 2.9 to 3.7 (ANOVA: \( F_{19,47} = 6.024, p < 0.001 \)). Compared with nematodes from the same
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Table 1. $\delta^{13}$C and $\delta^{15}$N values (mean ± SD, with number of replicates shown in parentheses) and trophic levels (TL, mean) of harpacticoid copepods and nematodes from scallop farms in Laizhou Bay, China in 2015 and 2016. Samples were taken from ‘control’ and ‘farm’ sites at 3 stations, characterized by sediment type as sandy-shallow (SS), muddy-shallow (MS) and muddy-deep (MD). Results of $t$-tests of TL in taxa co-occurring in farm and control sites are shown. ***$p < 0.001$; **$p < 0.01$; *$p < 0.05$; NS = non-significant; n/a: not applicable.

| Station | Taxon      | Control $\delta^{13}$C | Control $\delta^{15}$N | Control TL | Farm $\delta^{13}$C | Farm $\delta^{15}$N | Farm TL | $t$-test for TL ($t$ (df) p-value) |
|---------|------------|-------------------------|-------------------------|------------|----------------------|----------------------|---------|-------------------------------|
| 2016    | Canuellida | −20.32 ± 0.08 (3)       | 11.63 ± 0.08 (3)       | 1.9        | −20.40 ± 0.01 (4)   | 12.55 ± 0.03 (4)   | 2       | 0.818 (5) 0.451NS            |
| SS      | Canuellida | −19.28 ± 0.06 (3)       | 8.52 ± 0.05 (3)        | 2          | −19.50 ± 0.07 (3)   | 8.13 ± 0.13 (3)    | 2       | 0.088 (4) 0.943NS            |
| MS      | Nematoda   | −19.68 ± 0.12 (3)       | 12.04 ± 0.29 (3)       | 3.5        | −20.21 ± 0.16 (3)   | 12.07 ± 0.20 (3)   | 3.7     | 1.922 (4) 0.117NS            |
| MD      | Laophontidae | −23.02 ± 0.76 (3)     | 8.71 ± 0.97 (3)        | 2          | −20.98 ± 0.18 (3)   | 9.61 ± 0.19 (3)    | 2.4     | 1.705 (4) 0.163NS            |
|           | Ectinosomatidae | −20.39 ± 0.11 (3) | 9.68 ± 0.19 (3)        | 2.4        | −19.51 ± 0.26 (3)   | 11.41 ± 0.48 (3)   | 3.2     | 6.014 (4) 0.004**           |
|           | Miraciidae | n/a                    | n/a                    | n/a        | −19.72 ± 0.31 (3)   | 9.67 ± 0.25 (3)    | 2.4     | n/a (n/a) NS                  |
|           | Canuellida | n/a                    | n/a                    | n/a        | −19.16 ± 0.20 (3)   | 8.63 ± 0.28 (3)    | 2       | n/a (n/a) NS                  |
|           | Nematoda   | −21.36 ± 0.21 (3)      | 11.83 ± 0.26 (3)       | 3.4        | −19.88 ± 0.32 (4)   | 12.04 ± 0.42 (4)   | 3.5     | 0.987 (5) 0.369NS            |
| 2015    | Canuellida | −20.35 ± 0.42 (4)      | 8.98 ± 0.22 (4)        | 2          | −20.58 ± 0.33 (4)   | 10.66 ± 0.11 (4)   | 2       | 1.731 (6) 0.134NS            |
| SS      | Nematoda   | −19.54 ± 0.65 (3)      | 8.25 ± 0.51 (3)        | 1.7        | −19.67 ± 0.72 (4)   | 10.89 ± 0.31 (4)   | 2.1     | 3.820 (5) 0.012*             |
| MS      | Canuellida | −21.27 ± 0.83 (4)      | 10.36 ± 0.38 (4)       | 2          | −20.95 ± 0.07 (4)   | 11.55 ± 0.11 (4)   | 2       | 0.986 (6) 0.362NS            |
|           | Nematoda   | −20.83 ± 0.19 (3)      | 13.28 ± 0.21 (3)       | 3.3        | −20.38 ± 0.12 (3)   | 13.97 ± 0.09 (3)   | 3       | 3.960 (4) 0.017*             |
| MD      | Bulk copepoda | −21.08 (1)            | 11.66 (1)              | 2          | −21.45 ± 0.04 (2)   | 12.0 ± 0.4 (2)     | 2       | n/a (n/a) NS                  |
|           | Nematoda   | −20.88 ± 0.29 (2)      | 13.65 ± 0.01 (2)       | 2.9        | −20.22 ± 0.07 (2)   | 14.37 ± 0.17 (2)   | 3       | n/a (n/a) NS                  |

*aFamily Canuellidae represented >70% of sampled organisms per replicate.

Fig. 3. Comparison of (A) $\delta^{13}$C and (B) $\delta^{15}$N values for harpacticoid copepod and nematode taxa between ‘farm’ and ‘control’ sites at 3 stations (SS, MS and MD) at scallop farms in Laizhou Bay, China in 2015 and 2016. See Fig. 1 legend for abbreviations of stations. In each panel, the solid line represents a 1:1 correlation between $\delta^{13}$C or $\delta^{15}$N values at farm and control sites, and the 2 dashed lines show the 95% CI. Values outside the 95% CI are significantly different. Nema: nematodes; Cop Can.: copepod family Canuellidae; Cop Ect.: copepod family Ectinosomatidae; Cop Lao.: copepod family Laophontidae; Cop other: rest of the pooled copepods. In 2015, all copepods are pooled but still represented by the dominant family: Canuellidae, representing >70% per replicate.
farm and control sites since they fell within the 1:1 correlation 95% CI. For $\delta^{15}$N, nematodes at Stns MS and MD had similar values between farm and control sites, but nematodes at the farm site at Stn SS showed a more enriched $\delta^{15}$N compared to the control site. All copepods in 2015 and all the abundant families in 2016 (except family Canuellidae at Stn MS) showed more enriched $\delta^{15}$N at farm sites compared to control sites.

**Mixing model estimation of utilization by meio-benthos**

In general, faeces-derived materials were utilized by many copepods under the farms, contributing 12 to 61%, 13 to 60%, 44 to 83%, and 49 to 83% to the diets of Miraciidae, Laophontidae, and Ectinosomatidae at Stn MD, and Canuellidae at Stn SS, respectively (Table 2). However, the quantities of faeces consumed by copepods differed among the stations. For Canuellidae, scallop faeces were predominantly consumed at the farm site at Stn SS (contributing at least almost half of their diet).

**Table 2. Contribution (mean, with 95% CIs in parentheses) of particulate organic matter plus microphytobenthos (POM+MPB), seagrass and faeces in the diet of meio-benthos taxa collected from ‘farm’ and ‘control’ sites at 3 stations at scallop farms in Laizhou Bay, China in 2015 and 2016. See Table 1 legend for abbreviations of stations. The values were calculated using the Bayesian stable isotope mixing model (MixSIR). n/a: not applicable**
ets), while faeces were less important at the farm sites at Stns MS and MD (Table 2, Fig. 4). By contrast, nematodes at Stns MS and MD consumed very little faeces-derived material (<1%).

**Fatty acid profiles of the copepod family Canuellidae**

There was a significant difference in FA profiles over time (PERMANOVA, p < 0.01; ANOSIM, R > 0.75; Table 3), indicating that the FA profile of Canuellidae changed in time. In line with this, the nMDS showed a grouping for samples before aquaculture activity (T0) and the ones collected during aquaculture activity (T1) (Fig. 5). Based on the results of SIMPER, DHA contributed the most to this dissimilarity (29.12%). Before aquaculture activity, 16:0 was the major component of the total FA, contributing 46.1 and 47.0% of the total FA of family Canuellidae at farm and control sites respectively, while DHA was the most important FA during aquaculture activities, contributing 39.01 and 29.38% at farm and control sites respectively (Table 4).

In addition, a seasonal effect contributed to this pattern: significant differences were found between before and during aquaculture (‘Before vs. During’) within control and farm sites (see pairwise PERMANOVA tests in Table 3). More importantly, the difference ‘Before vs During’ was more significant within ‘Control’ than within ‘Farm’ (p < 0.05 and p < 0.001 respectively; Table 3), demonstrating that aquaculture made a crucial contribution to the FA profile of Canuellidae at Stn SS. Pairwise PERMANOVA also showed that farm and control sites strongly differed during the aquaculture stage (‘Farm vs. Control’ with ‘During’: p < 0.001; Table 3), though more replicates were needed to confirm this pattern in a pairwise test between farm and control within ‘Before’.

When excluding the seasonal effect, SIMPER showed that DHA contributed 33.71% to the difference between farm and control sites during aquaculture. During aquaculture at Stn SS, Canuellidae at the farm site had higher amounts of DHA and PUFAs than at the control site (Fig. 6, Table 4, Table S2 in the Supplement). 20:1ω9 and PUFA/SFA, considered as the indicators of carnivorous diet (Stevens et al. 2004), showed higher values in Canuellidae at farm sites during aquaculture at Stn SS. The trophic marker of diatoms/dinoflagellates, i.e. EPA/DHA (Cripps & Atkinson 2000), was lower at farm sites, indicating the different diets of Canuellidae at farm sites compared to those at control sites. Other trophic FA biomarkers showed no difference between farm and control sites (Fig. 6; see also Table S2; p > 0.05).

**DISCUSSION**

**Fate of bay scallop biodeposits in the aquaculture area**

Stable isotopes analysis has been applied to trace the fate of aquaculture waste in several studies...
Table 4. Relative fatty acid composition of the harpacticoid copepod family Canuelliidae from ‘farm’ and ‘control’ sites at Stn SS (see Table 1 legend for abbreviation) at a scallop farm in Laizhou Bay, China before and during seasonal aquaculture in 2016. Values for ‘during aquaculture’ are mean ± SD, with the number of replicates given in parentheses. ALA: alpha-linolenic acid; EPA: eicosapentaenoic acid; DHA: docosahexaenoic acid; n/a: not applicable

| Fatty acid | Before aquaculture | During aquaculture |
|------------|--------------------|--------------------|
|            | Farm | Control | Farm | Control |
| 14:0       | 6.61 (1) | 7.52 (1) | 1.03 ± 0.15 (4) | 1.63 ± 0.59 (3) |
| 15:0       | 5.95 (1) | 6.67 (1) | 0.88 ± 0.12 (4) | 1.18 ± 0.19 (3) |
| 16:0       | 46.1 (1) | 47.0 (1) | 14.67 ± 1.19 (4) | 20.54 ± 3.80 (3) |
| 16:1ω7     | n/a | n/a | 3.34 ± 0.55 (4) | 3.91 ± 0.73 (3) |
| 17:0       | 7.09 (1) | 7.4 (1) | 3.41 ± 0.14 (4) | 3.00 ± 0.13 (3) |
| 17:1ω7     | n/a | n/a | n/a | 0.06 ± 0.11 (3) |
| 18:0       | 27 (1) | 25.6 (1) | 12.47 ± 0.43 (4) | 17.60 ± 1.09 (3) |
| 18:1ω9t    | n/a | n/a | 2.62 ± 0.83 (4) | 2.79 ± 0.92 (3) |
| 18:1ω9c    | n/a | n/a | 4.90 ± 0.14 (4) | 4.25 ± 0.75 (3) |
| 18:2ω6c    | n/a | n/a | 1.03 ± 0.69 (4) | 1.30 ± 0.13 (3) |
| 20:0       | 1.53 (1) | 1.01 (1) | 0.50 ± 0.08 (4) | 0.50 ± 0.04 (3) |
| 18:3ω3 ALA | n/a | n/a | 0.77 ± 0.08 (4) | 0.74 ± 0.07 (3) |
| 20:1ω9     | n/a | n/a | 2.05 ± 0.19 (4) | 1.30 ± 0.02 (3) |
| 20:2ω6     | n/a | n/a | 0.38 ± 0.02 (4) | n/a |
| 22:0       | 2.01 (1) | 1.49 (1) | 0.83 ± 0.01 (4) | 0.56 ± 0.04 (3) |
| 20:3ω6     | n/a | n/a | n/a | 0.16 ± 0.28 (3) |
| 22:1ω9     | n/a | n/a | 0.13 ± 0.26 (4) | n/a |
| 23:0       | 0.96 (1) | 0.74 (1) | 0.61 ± 0.02 (4) | 0.25 ± 0.22 (3) |
| 20:5ω3 EPA  | n/a | 0.45 (1) | 9.82 ± 0.64 (4) | 9.77 ± 1.45 (3) |
| 24:0       | 2.23 (1) | 1.62 (1) | 0.93 ± 0.03 (4) | 0.60 ± 0.02 (3) |
| 24:1ω6     | n/a | n/a | 0.51 ± 0.03 (4) | 0.38 ± 0.36 (3) |
| 22:6ω3 DHA | n/a | n/a | 39.01 ± 1.84 (4) | 29.38 ± 3.59 (3) |

(Kon et al. 2009, Gondwe et al. 2012, Callier et al. 2013), since the aquaculture-derived waste has distinctive isotopic values. In our study, the enriched δ15N of bay scallop faeces compared to other organic sources was a prerequisite in order to be able to trace its fate. Contrary to some studies that demonstrate the accumulation of aquaculture waste in the sediment organic matter (OM) pool, SSOM in our study did not show the enrichment of 15N by scallop farming activity. This could be explained by 2 hypotheses. First, the aquaculture-derived OM was diluted in the water column and thus did not sink to the sea floor (Vizzini & Mazzola 2012). Second, the local consumers rapidly incorporated the aquaculture-derived matter (Kon et al. 2009). As we found more enriched δ15N values of most copepods and some nematodes in the sediment under the farms (see ‘Effects of bay scallop aquaculture on the quality of harpacticoid copepods for higher trophic levels’), the second explanation is more plausible. Callier et al. (2013) reported a similar outcome; i.e. isotopic values shifted in invertebrates rather than in SSOM. Taken together with the results of our study, this suggests that analyzing the response of the benthos (i.e. a biotic response) to evaluate the effect of aquaculture will provide a more comprehensive view than investigating only the response of the sediment (i.e. an abiotic response).

Biodeposits of bay scallop as food sources to meiobenthos

The resource utilization of meiobenthos appears to be a promising tool to evaluate the effect of aquaculture effluents on ecosystem functioning in terms of energy flow (Kennedy & Jacoby 1999, De Troch et al. 2013). The incorporation of scallop faeces into the diets of most harpacticoid copepods at all stations and nematodes in the sandy station under the farms demonstrates that bay scallop farming changed the energy flow in the basal part of the food web. This evidence agrees with findings of other studies that aquaculture effluents serve as alternative food sources for the benthos (Dubois et al. 2007, Callier et al. 2013).

The flux of OM is usually enhanced in the shellfish aquaculture area (Newell 2004, McKindsey et al. 2011); consequently the food availability for benthic organisms increases. Moreover, in terms of food source profitability, biodeposits are considered to be of good nutritional value because of their high carbon and nitrogen contents, large proportions of labile OM, and the low C/N ratios (Kautsky & Evans 1987, Miller et al. 2002). Biodeposits are mucus-enriched and function as good substrates for bacteria (Hargrave 1976). Thus, abundant microorganisms colonize on the pellet particles during gut passage (Werry & Lee 2005, Cnudde et al. 2011) and after defecation (Fabiano et al. 1994), and efficiently rework the labile components within hours to days (Carlsson et al. 2010). They break down the refractory OM from the faecal pellets and also produce microbial nutrients (e.g. extracellular protein and exudates) that make them available to other benthic organisms (Kautsky & Evans 1987, Wotton & Malmqvist 2001). Furthermore, increasing densities of microorganisms (e.g. bacteria, ciliates and dinoflagellates) themselves provide more foods for meiobenthos (Epstein 1997, Moens & Vincx 1997). Therefore, biodeposit-derived materials can easily be consumed by meiobenthos and enter the basal food web.
Effect of bay scallop aquaculture on the diets of meiobenthos

While aquaculture-derived OM is a nutritional food source (McKindsey et al. 2011, Callier et al. 2013), the quantities and pathways of consumption by benthos vary among trophic groups and feeding modes (Dubois et al. 2007, Wai et al. 2011). In accordance with the large trophic diversity of meiobenthos in coastal areas (Hicks & Coull 1983, Jensen 1987), we found that the quantities and pathways of faeces consumption differed according to the feeding behaviors of the meiobenthos. This was clearly illustrated by the harpacticoid copepod families. Substrate browsers, like Laophontidae and Miraciidae, possibly took faecal OM through scraping or sweeping off the attached bacteria (Hicks & Coull 1983, Cnudde et al. 2013, Mascart et al. 2013). For Canuellidae, faecal OM were presumably assimilated by filtering-feeding the small particles suspended in the water column (Cnudde et al. 2015). With the provision of bacteria and protists on the biodeposits (Wotton & Malmqvist 2001, Bongiorni et al. 2005), Ectinosomatidae showed an increased TL, suggesting that the feeding type of this taxon changed from omnivorous to carnivorous. It is possible that the dominant species of Ectinosomatidae has been modified by farming, since this is a species-rich family with various resource utilization strategies including diatom-feeding, microvory, point-feeding, and predation (Coull & Dudley 1976, Seifried & Dürbaum 2000). Further detailed screening of the trophic ecology of meiobenthos is required to prove this.

In addition, the copepod family Canuellidae demonstrated that the effect of bay scallop biodeposition depends on the receiving environment. With a broad range of niche breadth (De Troch et al. 2003), Canuellidae could switch their diets to more nutritious organic sources such as scallop biodeposits if other sources were insufficient, for instance in a sandy bottom with low chlorophyll levels and poor organic resources (Cartaxana et al. 2006). At muddy stations, the small contributions of biodeposits to Canuellidae may be explained by the greater availability of autotrophic production.

As for nematodes, their TLs indicate that the communities were different according to the sediment type. Nematodes at muddy stations were carnivores...
and did not incorporate faeces-derived materials into their diets. The data suggest that even when biodeposits were present, they did not modify their feeding mode i.e. they were either consuming bacteria derived from MPB extracellular polymeric substrate (EPS) or predating on small nematodes that fed on EPS of MPB (Moens et al. 2005, Rzeznik-Orignac et al. 2008, Majdi et al. 2012). It is also possible that this group of nematodes resided in deeper sediment (Steyaert et al. 2003) and could not access the biodeposits, which had either already been consumed by surface locating copepods or were not buried deep enough into the sediments. In this case, bay scallop farming has no impact on the resource utilization by nematodes that are trophic specialists. In contrast, nematodes at the sandy station incorporated biodeposits under the scallop farm. Microvory of nematodes has been reported to be dominant below bay scallop farms (Netto & Valgas 2010), where they presumably benefit from higher microbial densities by organic loading (Mrito et al. 2000). In our study, the feeding group of nematodes was likely changed by bay scallop farming.

Effect of bay scallop aquaculture on the quality of harpacticoid copepods for higher trophic levels

PUFA concentrations, and especially those of DHA increased in Canuellidae (the most abundant harpacticoid copepods) as they consumed certain amounts of scallop faeces at farm sites during the high biodeposition stage. PUFAs are important compositions of cell membranes and are needed in animals at all taxonomic levels, including copepods (Ederington et al. 1995, Hartwich et al. 2013). Especially at low temperature, PUFAs promote membrane fluidity (Farkas 1979, Stillwell & Wassall 2003). Also, with higher levels of PUFA, copepods have higher reproduction ability in terms of eggs production (Ederington et al. 1995). In our study, as the winter had just started during our sampling season, we hypothesize that the higher contents of PUFA provided Canuellidae with better opportunities to reproduce and deal with the low temperature and food-stress periods.

Furthermore, PUFAs, EPA and DHA have been recognized as good biomarkers to describe the quality of food (Boon & Duineveld 1996, Goedkoop et al. 2000). As higher level consumers in marine ecosystems cannot synthesize highly unsaturated fatty acids (HUFAs) such as DHA, their HUFA levels are derived entirely from their food sources (Iverson 2009, Hartwich et al. 2013). As harpacticoid copepods are important food items for epibenthic organisms such as shrimps and juvenile fishes (Coull et al. 1995), their quality in terms of PUFA or HUFA levels becomes an important factor influencing the quality of higher consumers as a food source and thus enriching the entire food web. Thus, the fact that Canuellidae consuming biodeposits of bay scallops contained more PUFAs (especially DHA) implies that they constituted a more nutritious food item for higher level consumers.

However, the mechanisms behind the increasing levels of PUFA (especially DHA) are not clear yet because the pathways to accumulate DHA are complicated and depend on several factors (Bell & Tocher 2009, Werbrouck et al. 2016). Copepods may gain DHA directly from the food or synthesize DHA from shorter chain of PUFAs (Schlechtriem et al. 2006, De Troch et al. 2012). It has been suggested that organisms using ‘foreign foods’ (i.e. foods not originating from their habitat) were not accustomed to these, and this might have stimulated the compensatory biochemical pathways (Iverson 2009). It is possible that loading of biodeposits induces the pathway to accumulate PUFAs and DHA in Canuellidae. To further elucidate the role of faeces in the ecosystem functioning, there are many options for future investigation; for example to determine whether, how, and to what extent aquaculture-derived PUFAs is transferable through the food chain.

Bay scallop as a non-indigenous species for the local environment: a positive perspective

It is always a concern that culturing NIS, especially non-indigenous shellfish, in coastal marine ecosystem, includes ecological risks (Newell 2004, Shelton & Rothbard 2006, Minchin et al. 2009). Our study showed that, to some extent, farming the non-indigenous bay scallop provides an extra food source for the benthos and consequently improves the quality of the local benthic environment. This suggests that the farming bay scallop has a positive effect on ecosystem functioning. To make a more complete assessment of NIS aquaculture, further research is needed to compare the effects with those of culturing native species.

Our observations may also apply to other shellfish farms, but additional factors should be considered, such as aquaculture characteristics (e.g. cultured species, stocking densities, etc.), and the hydrodynamics and sediment type of the receiving environment, because these can also affect the activities of the benthos (Chamberlain et al. 2001, Giles et al. 2006).
CONCLUSIONS

Our study showed that stable isotopes, especially δ15N, are a powerful tool to trace possible functional changes due to aquaculture activities. We also showed that the biodeposits were directly/indirectly consumed by meiobenthos, and accordingly, harpacticoid copepods residing under the scallop farms improved their quality as food items for the next trophic level. Therefore, we demonstrate a positive effect of intensive farming of bay scallop Argopecten irradians on ecosystem functioning.

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LITERATURE CITED

Anderson MJ (2006) Distance-based tests for homogeneity of multivariate dispersions. Biometrics 62:243–253
Anderson MJ, Gorley R, Clarke K (2008) PERMANOVA+ for PRIMER: guide to software and statistical methods. PRIMER-E, Plymouth
Bell MV, Tocher DR (2009) Biosynthesis of polyunsaturated fatty acids in aquatic ecosystems: general pathways and new directions. In: Arts MT, Brett MT, Kainz MJ (eds) Lipids in aquatic ecosystems. Springer, New York, NY, p 211–236
Bongiorni L, Mirto S, Pusceddu A, Danovaro R (2005) Response of benthic protozoa and thraustochytrid protists to fish farm impact in seagrass (Posidonia oceanica) and soft-bottom sediments. Microb Ecol 50:268–276
Boon A, Duineveld G (1996) Phytopigments and fatty acids as molecular markers for the quality of near-bottom particulate organic matter in the North Sea. J Sea Res 35: 279–291
Callewaert M, McKindsey CW, Desrosiers G (2008) Evaluation of indicators used to detect mussel farm influence on the benthos: two case studies in the Magdalen Islands, Eastern Canada. Aquaculture 278:77–88
Callewaert M, Lefebvre S, Dunagan MK, Bataille MP, Coughlan J, Crowe TP (2013) Shift in benthic assemblages and organisms’ diet at salmon farms: community structure and stable isotope analyses. Mar Ecol Prog Ser 483: 153–167
Carlsson MS, Glud RN, Petersen JK (2010) Degradation of mussel (Mytilus edulis) fecal pellets released from hanger long-lines upon sinking and after settling at the sediment. Can J Fish Aquat Sci 67:1376–1387
Cartaxana P, Mendes C, Van Leeuwe M, Brotas V (2006) Comparative study on microphytobenthic pigments of muddy and sandy intertidal sediments of the Tagus estuary. Estuar Coast Mar Sci 66:225–230
Chamberlain J, Fernandes T, Read P, Nickell T, Davies I (2001) Impacts of biodeposits from suspended mussel (Mytilus edulis L.) culture on the surrounding surficial sediments. ICES J Mar Sci 58:411–416
Cifuentes L, Sharp J, Fookes ML (1988) Stable carbon and nitrogen isotope biogeochemistry in the Delaware estuary. Limnol Oceanogr 33:1102–1115
Clarke KR, Gorley RN (2006) PRIMER V6: user manual - tutorial. Plymouth Marine Laboratory, Plymouth
Cnudde C, Willems A, Van Hoorde K, Vyverman W, Moens T, De Troch M (2011) Effect of food preservation on the grazing behavior and on the gut flora of the harpacticoid copepod Paraphaiaacea tulfofasiata. J Exp Mar Biol Ecol 407:63–69
Cnudde C, Moens T, Willems A, De Troch M (2013) Substrate-dependent bacterivory by intertidal benthic copepods. Mar Biol 160:327–341
Cnudde C, Moens T, Werbrouck E, Leport G, Van Gansbeke D, De Troch M (2015) Trophodynamics of estuarine intertidal harpacticoid copepods based on stable isotope composition and fatty acid profiles. Mar Ecol Prog Ser 524:225–239
Coull BC, Dudley BW (1976) Delayed naupliar development of meiobenthic copepods. Biol Bull 150:38–46
Coull BC, Greenwood JG, Fiedler DR, Coull BA (1995) Sub-tropical Australian juvenile fish eat meiofauna: experiments with winter whiting Sillago maculata and observations on other species. Mar Ecol Prog Ser 125:13–19
Crawford CM, Macleod CKA, Mitchell IM (2003) Effects of shellfish farming on the benthic environment. Aquaculture 224:117–140
Cripps G, Atkinson A (2000) Fatty acid composition as an indicator of carnivory in Antarctic krill, Euphausia superba. Can J Fish Aquat Sci 57:31–37
de Lima LCM, Navarro DMAF, Souza-Santos LP (2013) Effect of diet on the fatty acid composition of the copepod Tisbe Battioni. J Crustac Biol 33:372–381
De Troch M, Fiers F, Vinckx M (2003) Niche segregation and habitat specialisation of harpacticoid copepods in a tropical seagrass bed. Mar Biol 142:345–355
De Troch M, Boeckx P, Cnudde C, Van Gansbeke D, Vanreusel A, Vinckx M, Caramujo MJ (2012) Bioconversion of fatty acids at the basis of marine food webs: insights from a compound-specific stable isotope analysis. Mar Ecol Prog Ser 465:53–67
De Troch M, Riede C, De Troch M, Doi H, Kikuchi E, Hino S, Takagi S, Itoh T, Shikano S (2003) Impacts of oyster farming on macrofaunal assemblages and functional responses of harpacticoid copepods to anoxia in the Northern Adriatic: an experimental approach. Biogeosciences 10:4259–4272
DeNiro MJ, Epstein S (1978) Influence of diet on the distribution of carbon isotopes in animals. Geochim Cosmochim Acta 42:495–506
Doi H, Kikuchi E, Hino S, Itoh T, Takagi S, Shikano S (2003) Seasonal dynamics of carbon stable isotope ratios of particulate organic matter and benthic diatoms in strongly acidic Lake Katunuma. Aquat Microb Ecol 33:87–94
Dubois S, Marin-Leal JC, Rupert M, Lefebvre S (2007) Effects of oyster farming on macrofaunal assemblages associated with Lanice conchilega tubeworm popula-
tions: a trophic analysis using natural stable isotopes. Aquaculture 271:336–349

Ederington MC, McManus GB, Harvey HR (1995) Trophic transfer of fatty acids, sterols, and a triterpenoid alcohol between bacteria, a ciliate, and the copepod *Acartia tonsa*. Limnol Oceanogr 40:860–867

Epstein SS (1997) Microbial food webs in marine sediments. II. Seasonal changes in trophic interactions in a sandy tidal flat community. Microb Ecol 34:199–209

Fabiano M, DANOVARO R, Oliveri E, Misco C (1994) Decomposition of faecal matter and somatic tissue of *Mytilus galloprovincialis*: changes in organic matter composition and microbial succession. Mar Biol 119:375–384

FAO (Food and Agriculture Organization of the United Nations) (2016) The state of world fisheries and aquaculture 2016. FAO, Rome

Farkas T (1979) Adaptation of fatty acid compositions to temperature — a study on planktonic crustaceans. Comp Biochem Physiol B 64:71–76

Giles H, Pilditch CA, Bell DG (2006) Sedimentation from mussel (*Perna canaliculus*) culture in the Firth of Thames, New Zealand: impacts on sediment oxygen and nutrient fluxes. Aquaculture 261:125–140

Goedkoop W, Sonesten L, Ahlgren G, Boberg M (2000) Fatty acids in profound benthic invertebrates and their major food resources in Lake Erken, Sweden: seasonal variation and trophic indications. Can J Fish Aquat Sci 57:2267–2279

Gondwe MJ, Guildford SJ, Hecky RE (2012) Tracing the flux of aquaculture-derived organic wastes in the southeast arm of Lake Malawi using carbon and nitrogen stable isotopes. Aquaculture 350–353:8–18

Guo X, Luo Y (2016) Scallops and scallop aquaculture in China. In: Shumway SE, Parsons GJ (eds) Scallops, 3E. Elsevier Science, Oxford, p 937–952

Hargrave BT (1976) The central role of invertebrate faeces in sediment decomposition. In: Anderson JM, Macfadyen A (eds) The role of terrestrial and aquatic organisms in decomposition processes. Blackwell Scientific, Oxford, p 301–321

Hartstein ND, Rowden AA (2004) Effect of biodeposits from mussel culture on macroinvertebrate assemblages at sites of different hydrodynamic regime. Mar Environ Res 57:339–357

Hartwich M, Martin-Creuzburg D, Wacker A (2013) Seasonal changes in the accumulation of polyunsaturated fatty acids in zooplankton. J Plankton Res 35:121–134

Hicks GRF, Coull BC (1983) The ecology of marine harpacticoid copepods. Oceanogr Mar Biol Annu Rev 21:67–175

Hoshika A, Sarker MJ, Ishida S, Mishima Y, Takai N, Hayashizaki KI, Kurokura H (2009) Effect of shrimp farming organic waste on food availability for deposit feeder crabs in a mangrove estuary, based on stable isotope analysis. Fish Sci 75:715–722

Hoshika A, Sarker MJ, Ishida S, Mishima Y, Takai N (2006) Feeding of biofilm-dwelling nematodes examined using HPLC-analysis of gut pigment contents. Hydrobiologia 680:219–232

Iverson SJ (1976) The central role of invertebrate faeces in sediment decomposition. In: Anderson JM, Macfadyen A (eds) The role of terrestrial and aquatic organisms in decomposition processes. Blackwell Scientific, Oxford, p 301–321

Jin X, Shan X, Li X, Wang J, Cui Y, Zuo T (2013) Long-term changes in the fishery ecosystem structure of Laizhou Bay, China. Sci China Earth Sci 56:366–374

Kalantzis I, Karakassi I (2006) Benthic impacts of fish farming: meta-analysis of community and geochemical data. Mar Pollut Bull 52:484–493

Kautsky N, Evans S (1987) Role of biodeposition by *Mytilus edulis* in the circulation of matter and nutrients in a Baltic coastal ecosystem. Mar Ecol Prog Ser 38:201–212

Kelly JR, Scheibing RE (2012) Fatty acids as dietary tracers in benthic food webs. Mar Ecol Prog Ser 446:1–22

Kennedy AD, Jacoby CA (1999) Biological indicators of marine environmental health: meiofauna — a neglected benthic component? Environ Monit Assess 54:47–68

Kon K, Kawakubo N, Aoki JL, Tongnunui P, Hayashizaki KI, Kurokura H (2009) Effect of shrimp farming organic waste on food availability for deposit feeder crabs in a mangrove estuary, based on stable isotope analysis. Fish Sci 75:715–722

Leduc D, Probert PK (2009) The effect of bacterivorous nematodes on detritus incorporation by macrofaunal detritivores: a study using stable isotope and fatty acid analyses. J Exp Mar Biol Ecol 371:130–139

Leduc D, Probert PK, Duncan A (2009) A multi-method approach for identifying meiofaunal trophic connections. Mar Ecol Prog Ser 383:95–111

Li Y, Veilleux DJ, Wikfors GH (2009) Particle removal by Northern bay scallops *Argopecten irradians irradians* in a semi-natural setting: application of a flow-cytometric technique. Aquaculture 296:237–245

Liu H, Fang J, Zhu J, Dong S and others (2004) Study on limiting nutrients and phytoplankton at long-line-culture areas in Laizhou Bay and Sanggou Bay, northeastern China. Aquat Conserv 14:551–574

Majdi N, Tackx M, Traunspurger W, Buffan-Dubau E (2012) Influence of suspended and off-bottom mussel culture on the sea bottom and benthic habitats: a review. Can J Zool 89:622–646

Mascart T, Lepoint G, De Troch M (2013) *Meiofauna and harpacticoid copepods* in different habitats of a Mediterranean seagrass meadow. J Mar Biol Assoc UK 93:1557–1566

McCandless CW, Archambault P, Callier MD, Olivier F (2011) Influence of suspended and off-bottom mussel culture on the sea bottom and benthic habitats: a review. Can J Zool 89:622–646

Miller DC, Norkko A, Pilditch CA (2002) Influence of diet on dispersal of horse mussel *Atrina zelandica* biodeposits. Mar Ecol Prog Ser 242:153–167

Minagawa M, Wada E (1984) Stepwise enrichment of $^{15}$N and animal age. Geochim Cosmochim Acta 48:1135–1140

Minchin D, Gollasch S, Cohen AN, Hewitt CL, Olenin S (2009) Characterizing vectors of marine invasion. In: Rilov G, Crooks JA (eds) Biological invasions in marine ecosystems. Ecological Studies 204. Springer, Berlin, p 153–172

Mirti S, La rosa T, Danovaro R, Mazzola A (2000) Microbial and meiofaunal response to intensive mussel-farm biodeposition in coastal sediments of the western Mediterranean. Mar Pollut Bull 40:244–252

Moens T, Vincx M (1997) Observations on the feeding ecology of estuarine nematodes. J Mar Biol Assoc UK 77:211–227
Moens T, Bouillon S, Gallucci F (2005) Dual stable isotope abundances unravel trophic position of estuarine nematodes. J Mar Biol Assoc UK 85:1401−1407
Moore JW, Semmens BX (2008) Incorporating uncertainty and prior information into stable isotope mixing models. Ecol Lett 11:470−480
Nanton DA, Castell JD (1999) The effects of temperature and dietary fatty acids on the fatty acid composition of harpacticoid copepods, for use as a live food for marine fish larvae. Aquaculture 175:167−181
Netto SÁ, Valgas I (2010) The response of nematode assemblages to intensive mussel farming in coastal sediments (Southern Brazil). Environ Monit Assess 162:81−93
Newell RI (2004) Ecosystem influences of natural and cultivated populations of suspension-feeding bivalve molluscs: a review. J Shellfish Res 23:51−62
Peterson BJ, Heck KL (1999) The potential for suspension-feeding bivalves to increase seagrass productivity. J Exp Mar Biol Ecol 240:37−52
Phillips DL, Newsome SD, Gregg JW (2005) Combining sources in stable isotope mixing models: alternative methods. Oecologia 144:520−527
Pillay TVR (2008) Aquaculture and the environment, 2nd edn. Blackwell, London
Post DM (2002) Using stable isotopes to estimate trophic position: models, methods, and assumptions. Ecology 83:703−718
Power ME (1992) Habitat heterogeneity and the functional significance of fish in river food webs. Ecology 73:1675−1688
Rzeznik-Orignac J, Boucher G, Fichet D, Richard P (2008) Stable isotope analysis of food source and trophic position of intertidal nematodes and copepods. Mar Ecol Prog Ser 359:145−150
Sanz-Lázaro C, Sanchez-Jerez P (2017) Mussels do not directly assimilate fish farm wastes: shifting the rationale of integrated multi-trophic aquaculture to a broader scale. J Environ Manage 201:82−88
Schlechtriem C, Arts M, Zellmer I (2006) Effect of temperature on the fatty acid composition and temporal trajectories of fatty acids in fasting Daphnia pulex (Crustacea, Cladocera). Lipids 41:397−400
Seifried S, Dürrbaum J (2000) First clear case of carnivory in marine Copepoda Harpacticoida. J Nat Hist 34:1395−1618
Shelton WL, Rothbard S (2006) Exotic species in global aquaculture—a review. Isr J Aquicult Bamidgeh 58:3−28
Stevens CJ, Deibel D, Parrish CC (2004) Incorporation of bacterial fatty acids and changes in a wax ester-based omnivory index during a long-term incubation experiment with Calanus glacialis Jaschnov. J Exp Mar Biol Ecol 303:135−156
Steyaert M, Vanaverbeke J, Vanreusel A, Barranguet C, Lucas C, Vincx M (2003) The importance of fine-scale, vertical profiles in characterising nematode community structure. Estuar Coast Mar Sci 58:353−366
Stillwell W, Wassall SR (2003) Docosahexaenoic acid: membrane properties of a unique fatty acid. Chem Phys Lipids 126:1−27
Svensson CJ, Hynes GA, Laverty PS (2010) An efficient method for collecting large samples of live copepods free from detritus. Mar Freshw Res 61:621−624
Vizzini S, Mazzola A (2012) Tracking multiple pathways of waste from a northern bluefin tuna farm in a marine-coastal area. Mar Environ Res 77:103−111
Wai TC, Leung KM, Wu RS, Shin PK, Cheung SG, Li X, Lee JH (2011) Stable isotopes as a useful tool for revealing the environmental fate and trophic effect of open-sea cage fish farm wastes on marine benthic organisms with different feeding guilds. Mar Pollut Bull 63:77−85
Wang F (2015) Feeding ecology of two megagrazers in two typical bays. MSc thesis, University of Chinese Academy of Sciences, Beijing
Werbrouck E, Van Gansbeke D, Vanreusel A, De Troch M (2016) Temperature affects the use of storage fatty acids as energy source in a benthic copepod (Platycheilus littoralis, Harpacticoida). PLOS ONE 11:e0151779
Werry J, Lee SY (2005) Grapsid crabs mediate link between mangrove litter production and estuarine planktonic food chains. Mar Ecol Prog Ser 293:165−176
Wotton RS, Malmqvist B (2001) Feces in aquatic ecosystems. Bioscience 51:537−544
Zanden M, Rasmussen JB (2001) Variation in δ15N and δ13C trophic fractionation: implications for aquatic food web studies. Limnol Oceanogr 46:2061−2066
Zhao J, Chen JF (2001) Research on aquacultural hydro-environment of Eastern Laizhou Bay. Mar Fish Res 22:62−67 (in Chinese with English Abstract)
Zhou Y, Yang H, Zhang T, Liu S and others (2006) Influence of filtering and biodeposition by the cultured scallop Chlamys farreri on benthic-pelagic coupling in a eutrophic bay in China. Mar Ecol Prog Ser 317:127−141
Zhuang W, Gao X, Zhang Y, Xing Q, Tosi L, Qin S (2014) Geochemical characteristics of phosphorus in surface sediments of two major Chinese mariculture areas: the Laizhou Bay and the coastal waters of the Zhangzi Island. Mar Pollut Bull 83:343−351

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