Spatial Differences in East Scotia Ridge Hydrothermal Vent Food Webs: Influences of Chemistry, Microbiology and Predation on Trophodynamics

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Abstract

The hydrothermal vents on the East Scotia Ridge are the first to be explored in the Antarctic and are dominated by large pelagic gastropods, stalked barnacles (Vulcanolepas sp.) and anomuran crabs (Kiwa sp.) but their food webs are unknown. Vent fluid and macroconsumer samples were collected at three vent sites (E2, E9N and E9S) at distances of tens of metres to hundreds of kilometres apart with contrasting vent fluid chemistries to describe trophic interactions and identify potential carbon fixation pathways using stable isotopes. δ13C of dissolved inorganic carbon from vent fluids ranged from −4.6‰ to 0.8‰ at E2 and from −4.4‰ to 1.5‰ at E9. The lowest macroconsumer δ13C was observed in pelagic gastropods (−30.0‰ to −31.1‰) and indicated carbon fixation via the Calvin-Benson-Bassham (CBB) cycle by endosymbiotic gamma-Proteobacteria. Highest δ13C occurred in Kiwa sp. (−19.0‰ to −10.5‰), similar to that of the epibionts sampled from their ventral seata. Kiwa sp. δ13C differed among sites, which were attributed to spatial differences in the epibiont community and the relative contribution of carbon fixed via the reductive tricarboxylic acid (rTCA) and CBB cycles assimilated by Kiwa sp. Site differences in carbon fixation pathways were traced into higher trophic levels e.g. a stichasterid asteroid that predates on Kiwa sp. Sponges and anemones at the periphery of E2 assimilates a proportion of epipelagic photosynthetic primary production but this was not observed at E9. Differences in the δ13C and δ34S values of vent macroconsumers between E2 and E9 sites suggest the relative contributions of photosynthetic and chemosymbiotic carbon fixation (rTCA v CBB) entering the hydrothermal vent food webs vary between the sites.

Introduction

Deep-sea hydrothermal vents are chemically reducing habitats occurring on mid-ocean and back-arc spreading centres, seamounts, volcanic hotspots and off-axis ridge settings [1,2,3]. They are distinct from the surrounding deep sea with respect to environmental conditions, the energy sources sustaining life and their biological communities [4,5]. High densities of organisms are found to thrive at the interface where hot, mineral-rich fluids discharge from the seafloor and mix with colder, oxygenated seawater. The hot fluids emitted from the seafloor may differ in pH and are enriched in reduced gases (e.g. H2S, CH4, H2) and metals (e.g. Fe2+, Cu, Mn) relative to seawater [6]. Microorganisms oxidise the reduced species in vent fluids and utilise the energy released to fix CO2 or other single carbon compounds (e.g. CO2, CH4) into cellular material [7]. This results in microbial chemoautotrophy replacing photosynthetic primary production at the base of the food chain [7].

Sulfide oxidation appears to be the principal energy acquisition pathway, which microorganisms use to drive carbon fixation [3,7,8]. The most important carbon fixation pathways at the base of the metazoan hydrothermal vent food webs are the Calvin-Benson-Bassham (CBB) and reductive tricarboxylic acid (rTCA) cycles [9,10,11]. Methane oxidation (methanotrophy) is a further carbon fixation process at hydrothermal vents with CH4 of thermogenic, biogenic or magmatic origin available depending on the host substrate [3,12]. Epipelagic photosynthetic primary production may also provide some nutrition to vent macroconsumers, although the relative contribution to vent fauna is thought to be negligible [12,13]. Macroconsumers utilise the vent organic carbon through endo- and epipsymbiotic relationships, consumption of free-living microorganisms either from various surfaces or the water column and indirectly through predation and scavenging [14,15,16].
The relative contributions of different carbon sources and complexity of hydrothermal vent food webs vary greatly depending on the species present, the geological host substrate and the vent fluid chemistry [17,18,19]. The first Antarctic hydrothermal vent communities were discovered recently on the East Scotia Ridge (ESR), a back-arc spreading centre in the Atlantic sector of the Southern Ocean [20,21]. The two basalt-hosted vent fields occur on the ridge segments E2 and E9, which lack the characteristic alvinocarid shrimps, bathymodiolid mussels and siboglinid worms found at Atlantic, Indian and Pacific hydrothermal vents, respectively [21]. Instead, biomass at the ESR vents is dominated by anomuran crabs (Kiwa sp.), stalked barnacles (Fulacopecten sp.) and large peltopsiodont gastropods [22], indicating a new biogeographic province [21]. Furthermore, there are differences in the end-member vent fluid chemistry between the E2 and E9 vent fields as well as within field between northern (E9N) and southern (E9S) areas of E9 [21].

Stable isotopes of carbon ($^{13}$C/$^{12}$C expressed as $\delta^{13}$C), nitrogen ($^{15}$N/$^{14}$N expressed as $\delta^{15}$N) and sulfur ($^{34}$S/$^{32}$S expressed as $\delta^{34}$S) have been used to examine hydrothermal vent community biomass [23,24]. $\delta^{13}$C can be used to characterise the various carbon sources utilised by vent macroconsumers [25]. This is done by comparing the expected carbon fractionation between dissolved inorganic carbon (DIC) and the macroconsumer’s tissue. Enzymatic reactions catalysed by the ribulose-1,5-biphosphate carboxylase/oxygenase form 1 (RuBiCO form 1) of the CBB cycle (22%) to 30% [26,27,28]) exhibit greater fractionation than those of the RTCA cycle (2% to 14%) [29,30,31]). Once organic material is incorporated into the macroconsumer food web, carbon trophic discrimination ($\Delta^{13}$C) is small, ranging from 0 to 1.5% between the food source and consumer [32]. $\delta^{34}$S also identifies energy sources (sulfur trophic discrimination, $\Delta^{34}$S, $\Delta^{15}$N, $\Delta^{34}$S) between seawater sulphate and sulfides at hydrothermal vents [33] results in organic matter of photosynthetic ($\pm$16% to 19%) and chemosynthetic ($\pm$9% to 10%) origin having distinctive $\Delta^{34}$S values [34,35]. The greater trophic discrimination (2% to 5%) in $\delta^{15}$N between consumer and food source provides information on the trophic position of an organism relative to a primary consumer [32]. Therefore, the isotopic value of a vent macroconsumer is the product of the following factors: (1) the inorganic substrate and its isotopic value used by the chemoautotroph; (2) the isotopic discrimination processes occurring during metabolic reactions involving inorganic substrates to create organic compounds (e.g. CBB or RTCA cycles) by the chemoautotroph; (3) food-source macroconsumer trophic interactions (e.g. endosymbiont-host, predator-prey) that occur as a function of (1) and (2); and (4) the physiology associated with the macroconsumer’s isotopic trophic discrimination.

The goal of the present research was to investigate intra- and inter-site patterns in the trophic assemblages of macroconsumers occurring at hydrothermal vents on the ESR using $\delta^{13}$C, $\delta^{15}$N and $\delta^{34}$S. Specifically, the aims were to: (1) compare $\delta^{13}$C$_{DIC}$ among vent sites and thus establish difference in the isotopic inorganic substrates used by chemoautotrophs; (2) compare $\delta^{15}$C, $\Delta^{15}$N and $\delta^{34}$S between vent and benthic non-vent fauna to assess any photosynthetic inputs into the hydrothermal vent food web; (3) investigate differences in trophic structures among the three sites; and (4) assess which species are driving any differences in trophic structure. The investigation provides a unique opportunity to examine differences in trophic structure at the scale of tens of metres to 100s of kilometres in a newly discovered hydrothermal vent biogeographical province.

Materials and Methods

Ethics Statement

Permits for the fieldwork were granted by the United Kingdom Foreign and Commonwealth Office. This study met the ethical requirements of the affiliated research institutions for research utilising animal tissues. No animal husbandry or laboratory controlled experiments were part of the research that required permits from the UK Home Office. The fish were collected at a water depth of 2500 m, which meant that they were dead when they arrived on deck as a result of changes in pressure. This was the case with the majority of the animals dissected within this study. The research also adhered to the Inter Ridge code of conduct for sampling hydrothermal vents (http://www.interridge.org/IRStatement).

Study Sites

The E2 and E9 vent fields are situated approximately 440 km apart at 56° 05.35’S, 30° 19.20’W and 60° 02.50’ S, 29° 58.93’ W, respectively (Fig. 1). E2 is at a depth of ~2600 m and seafloor topography is complex with a series of terraced features and lobed pillow basalt filling a major north-south steep-sided fissure [21]. The main high-temperature and diffusive venting occurred at an intersection between this fissure and an east-west running fault or scarp [21]. E9 was located at ~2400 m depth and its topography was relatively flat with sheet lava, a series of lava drain back features and collapsed pillow basalt. A series of north-south fissures were found with venting mainly occurring on the most western [21,22]. The end-member fluid chemistry exiting chimneys differed between the northern and southern sections of E9 [21], therefore E9N and E9S are here considered to be separate sites. Ambient seabed water temperatures were 0.0°C at E2 and between ~0.1°C and ~1.3°C at E9 [21].

Sample Collection and Ship-board Processing

Samples were collected onboard the R. R. S. James Cook during the 2010 austral summer (7 January to 21 February) using the remotely operated vehicle (ROV) Isis. High temperature and diffuse flow fluids were collected for DIC using titanium samplers, equipped with an inductively coupled link high temperature sensor. The nozzle of the titanium sampler was inserted into the chimney orifice for high temperature fluid samples and once the temperature reading became stable the fluid was collected. For diffuse flow samples, a circular titanium housing was placed over the area of diffuse venting to minimise the entrainment of seawater. Once the diffuse flow was visible exiting the top of the housing, the titanium sampler was inserted into the opening and the diffuse flow sample was collected once the temperature reading was stable. On board, an aliquot for stable isolate analysis of DIC was sampled to exclude air and poisoned with mercury chloride.

Vent macroconsumers were collected by suction sampler or scoop with species separated into a series of acrylic chambers or perspex boxes to avoid predation or contamination. Six species were collected at all three sites. No female Kiwa sp. were collected from E9N or E9S. Fish and pycnogonids were collected using large collapsible and small metal baited traps deployed from the ROV. Non-vent macroconsumers were collected from metres to tens of metres away from active venting where there were no obvious signs of hydrothermal influence, i.e. no bacterial mat, and where temperature was consistent with local Antarctic bottom water. Non-vent samples were collected on separate dives from those for vent fauna to avoid contamination. Only one non-vent species was collected from E2 and sampling was limited to the areas adjacent to E9N because of ROV operational time.
constraints. Potential food sources were collected by scraping material from rocks collected by ROV manipulators and epibionts from the ventral setae of the decapod *Kiwa* sp. Particulate suspended material was collected from the acrylic chambers, which was sampled incidentally during faunal collection. Samples were sorted on board to the lowest possible taxonomic resolution. The majority of the vent species are undescribed to date.

Faunal samples were frozen at $-80^\circ$C whole or after dissection, depending on their size, for stable isotope analysis. Muscle was removed from the chelipeds of *Kiwa* sp., foot dissected from Peltospioidea sp., tube feet removed from the asteroids Stichasteridae sp. and *Freyella cf. fragilissima* and tentacles removed from the anemones. Legs were removed from the pycnogonids Colossendeis cf. concedis and C. cf. elephantis, while *Sericosura* spp. was sampled whole. The gastropods Provannidae sp. 1 and 2, *Lepetodrilus* sp., and juvenile Peltospioidea sp. (<7 mm shell length), and the stalked barnacle *Vulcanolepas* sp. were removed from their shells and sampled whole. White muscle tissue was dissected from the anterior dorso-lateral region of the zoarcid fish.

Sample Processing Onshore

Each end-member and diffuse flow DIC sample was prepared for isotopic analysis by removing a 1 mL water sample and transferring it into a separate vial. The headspace was flushed with helium, phosphoric acid was injected into the vial and then the contents were vortex mixed. The samples were then left to react for 24 hours to ensure complete conversion of all DIC to CO$_2$ for isotopic analysis. The CO$_2$ was then analysed by continuous-flow isotope ratio mass spectrometry (IRMS) using a Europa Scientific 20–20 IRMS by Iso-Analytical (Crewe, United Kingdom). Samples were run in duplicate and the mean is reported. An internal reference gas (IA-R060, $\delta^{13}$C = $-36.08\pm 0.13$) was used to determine the $\delta^{13}$CDIC values and is traceable to the International Atomic Energy Agency standard, NBS-19. Concentrations of CH$_4$ in the water samples were insufficient for isotope analysis.

Faunal tissue samples were freeze dried and ground to a homogenous powder using a pestle and mortar. Aliquots of fauna, particulate suspended material and material scraped from rocks were tested for carbonates prior to analysis with 0.1 N HCl. If the sample effervesced, this indicated carbonates were present and it was subsequently acidified by further addition of HCl until the effervescence ceased. Samples were re-dried at 50°C for 48 hours. If the sample did not effervesce, no acidification was carried out. Aliquots for $\delta^{13}$C analysis were not lipid extracted. Any confounding lipid effects due to metabolic processes would not affect the interpretation of the ultimate carbon sources of the vent fauna described by $\delta^{13}$C because of the large differences in the $\delta^{13}$C values of trophic end-members.
Approximately 0.7 mg of powder was weighed into a tin capsule for carbon and nitrogen IRMS. For sulfur, 2 mg of sample and 4 mg of the catalyst vanadium pentoxide were weighed into each tin capsule. Dual stable carbon and nitrogen isotope ratios were measured by continuous-flow IRMS using a Costech Elemental Analyser interfaced with Thermo Finnigan Delta Plus XP (Natural Environment Research Council, Life Sciences Mass Spectrometry Facility, SUERC, East Kilbride, United Kingdom). Two laboratory standards were analysed for every ten samples in each analytical sequence. These alternated between paired alanine standards, differing in δ13C and δ15N, and an internal laboratory gelatin standard. Sulfur was analysed by Iso-Analytical using a SERCON Elemental Analyser coupled to a Europa Scientific 20–20 IRMS. Laboratory standards of barium sulphate (two sets of differing δ34S) and silver sulphide were used for calibration and drift correction. An internal standard of whole baleen was used for quality control (n = 28, 16.34% ± SD 0.21). Stable isotope ratios were expressed in delta (δ) notation as parts per thousand/permil (%). All internal standards are traceable to the following international standards: v-PDB (Pee Dee Belemnite), AIR (atmospheric nitrogen) and NBS-127 (barium sulphate), IAEA-S-1 (silver sulphide) and IAEA-SO-5 (barium sulphate). An external reference material of freeze dried and ground deep-sea fish white muscle (Antipodina rostrata) was also analysed (δ13C, n = 24, −18.94% ± SD 0.09; δ15N, n = 24, 13.11% ± SD 0.38; δ34S, n = 30, 18.20%, ± SD 0.59).

Data Analysis

Data were assessed for normality using a Shapiro-Wilk test before statistical tests examining spatial patterns in trophic structure and species stable isotope values. Homogeneity, or otherwise, of variances is ecologically informative, for example in identifying distinct energy sources at the base of the food web [36]. Inter-site differences in trophic structure were examined using a Fligner-Killeen test for homogeneity of variance to assess differences in the spread of the mean stable isotope values of each species. Inter-site differences in species were analysed using a one-way ANOVA followed by Tukey’s honest significant difference (HSD) when variance was homogeneous among sites. Welch’s ANOVA followed by t-tests were used when there was heterogeneity of variance among sites because it uses adjusted degrees of freedom to protect against Type I errors when variances are unequal [37]. A Bonferroni correction (p = 0.05/n) was used for multiple comparisons. When data were not normally distributed, a two sample Wilcoxon test was used. All statistics were preformed in R version 12.13.1 [38].

Results

Dissolved Inorganic Carbon Stable Isotope Values

Mean (± SD) δ13C_DIC of high temperature and diffuse flow fluids are summarised in Table 1. δ13C_DIC of high temperature samples collected at two E2 locations were −4.7% (±0.0) (max temperature 351.0°C) and −2.5% (±0.1) (max temperature 323.0°C). At E9N, δ13C_DIC from separate orifices of the same chimney structure were −4.6% (±0.0) (max temperature 380.2°C) and −4.5% (±0.0) (max temperature 357.0°C). No high temperature fluids were collected from E9S for δ13C_DIC analysis because the pressure was too high within the titanium samples to safely and accurately collect a representative sample. Diffuse flow samples from amongst Kiwa sp. and anemones, at E2, had δ13C_DIC values of 0.8% (±0.1) (max temperature 19.9°C) and 0.2% (±0.2) (max temperature 3.5°C), respectively. A single diffuse flow sample collected from amongst an aggregation of Kiwa sp. at E9N had a δ13C_DIC value of 1.3% (±0.1) (max temperature 12.6°C). At E9S, diffuse flow samples from amongst Kiwa sp. had a δ13C_DIC value of 0.9% (±0.1) (max temperature 19.9°C), while a sample taken from a mixed aggregation of Kiwa sp. and peltoспорoid gastropods had δ13C_DIC value of 0.1% (±0.1) (max temperature 5.0°C).

Comparison between Vent and Benthic Non-vent Macro-consumers at E9N

At E9N, mean δ13C and δ15N values of vent fauna overlapped with non-vent benthic fauna (Welch’s t-test, δ13C DF = 10.59, t = 0.66, p = 0.52; Welch’s t-test, δ15N DF = 10.42, t = −0.30, p = 0.76; Fig. 2, Tables 2 & 3) while mean δ34S values differed between non-vent benthic fauna and vent fauna (Welch’s t-test, DF = 12.36, t = −9.08, p < 0.01) (Fig. 3, Tables 2 & 3).

Intra- and Inter-site Differences in Community Trophodynamics

Eleven, ten and seven species were collected at E2, E9N and E9S respectively for stable isotope analysis (Table 3). The ranges of mean δ13C values of the vent fauna differed amongst the three sites (Fig. 2, ANOVA, df = 2, F = 8.67, p = 0.03). At E2, the narrowest δ13C range (−29.9% to −19.0%) was observed, whereas at E9N and E9S δ13C ranged from −31.4% to −9.9% and −30.0% to −10.5%, respectively (Fig. 2). Across the three sites Peltoспорoida sp. had the lowest values while Kiwa sp. had the highest δ13C values (Fig. 2, Table 3), and Lepetodrilus sp., Vulcanoepus sp., Panamarcus sp. and Colossendeis spp. all had intermediate δ13C values (Fig. 2, Table 3). However, there was no overall difference in mean δ13C values among sites for the combined data across species (Welch’s ANOVA, DF = 2, F = 0.59, p = 0.56). The range and mean δ34S values (Fig. 3, Table 3) did not differ among sites (Fligner-Killeen test, DF = 2, χ2 = 0.84, p = 0.65; ANOVA, DF = 2, 26, F = 1.94, p = 0.16), however Kiwa sp. had the lowest δ34S at E2 and E9S while Lepetodrilus sp. had the lowest δ34S values at E9N (Fig. 3, Table 3). The highest vent fauna δ34S values were in Panamarcus sp. (E2), Vulcanoepus sp. (E9N) and Seiscusura spp. (E9S) (Fig. 3, Table 3). Neither the range nor the mean δ34S values differed among sites (Fligner-Killeen test, DF = 2, χ2 = 0.40, p = 0.83; ANOVA, DF = 2, 26, F = 1.19, p = 0.31). The progamm gastropods at E2 and E9S had the lowest δ34S values while Peltoспорoida sp. had the lowest values at E9N (Fig. 2, Table 3).

Table 1. δ13C values of dissolved inorganic carbon (DIC) sampled from high temperature and diffuse flow venting from the E2 and E9 ridge segments of the East Scotia Ridge, Southern Ocean.

| Site | Temperature (°C) | δ13C DIC |
|------|-----------------|----------|
| E2   | 351.0           | −4.7 (0.0) |
|      | 323.0           | −2.5 (0.1) |
|      | 19.9            | 0.8 (0.1) |
|      | 3.5             | 0.2 (0.2) |
| E9N  | 380.2           | −4.7 (0.0) |
|      | 357.0           | −4.7 (0.0) |
|      | 12.6            | 1.5 (0.1) |
| E9S  | 19.9            | 0.9 (0.1) |
|      | 5.0             | 0.1 (0.1) |

Standard deviations are in parentheses. doi:10.1371/journal.pone.0065553.t001
Figure 2. $\delta^{13}C$ and $\delta^{15}N$ values of macroconsumers collected from the East Scotia Ridge, Southern Ocean. The values represent means (± standard deviations) for hydrothermal vent and non-vent macroconsumers from the three sample sites: (a) E2, (b) E9N and (c) E9S. Dashed vertical lines represent potential ranges of $\delta^{13}C$ values indicative of carbon sources sustaining macroconsumers at the ESR: triple dashed line represents the Calvin-Benson-Bassham (CBB) cycle utilising form I RuBisCO, double dashed line represents the reductive tricarboxylic acid (rTCA) cycle, mixed carbon sources occur between the triple and double dashed line and the continuous dashed line represents the approximate $\delta^{13}C$ values of the dissolved inorganic carbon from the diffuse flow areas.

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The stichasterid sp. consistently had the highest $\delta^{15}$N values relative to the other vent fauna at each site (Fig. 2, Table 3).

Spatial Differences in Macroconsumer Trophodynamics

_Vulcanolepas_ sp. exhibited spatial differences in $\delta^{13}$C, $\delta^{15}$N and $\delta^{34}$S but there was no consistent pattern in isotopic differences among sites (Table 4). Male and female _Kiwa_ sp. at E2 did not differ in $\delta^{13}$C but males were lower in $\delta^{15}$N and $\delta^{34}$S than females (Table 5). Male _Kiwa_ sp. showed spatial differences in each stable isotope (Table 4). $\delta^{13}$C of the males showed a greater range (Figner-Killeen test, DF = 2, $\chi^2 = 10.91, p<0.01$) and lower values at E2 than E9N and E9S (Table 3 & 4). The epibionts attached to the ventral surface of male _Kiwa_ sp. also exhibited a greater spread of $\delta^{13}$C values at E2 than E9S (F-test, DF = 4, 3, F = 244.46, $p<0.01$).

Figure 3. $\delta^{13}$C and $\delta^{34}$S values of macroconsumers collected from the East Scotia Ridge, Southern Ocean. The values represent means (± standard deviations) for hydrothermal vent and non-vent macroconsumers from the three sample sites: (a) E2, (b) E9N and (c) E9S. $\delta^{34}$S values between the triple and double dashed lines represent potential areas of isotopic mixing between chemosynthetic and photosynthetic food sources. doi:10.1371/journal.pone.0065553.g003
### Table 2. Mean $\delta^{13}$C, $\delta^{15}$N and $\delta^{34}$S values (%) of non-vent deep-sea fauna collected from the E2 and E9 ridge segments of the East Scotia Ridge, Southern Ocean. Standard deviations are in parentheses.

| Taxonomic group | Species | Site  | N  | $\delta^{13}$C | $\delta^{34}$S | $\delta^{15}$N |
|-----------------|---------|-------|----|---------------|---------------|---------------|
| Crustacea       | Decapoda | Nematocarcinus lanceopes | E9  | 5  | −24.2 (0.7) | 18.9 (0.5) | 8.1 (0.4) |
|                 |         | Euphausia superba | E9  | 3  | −27.4 (0.8) | 19.0 (0.1) | 2.7 (0.7) |
| Echinodermata    | Asteroidea | Freyella cf fragilissima | E2  | 3  | −22.4 (0.3) | 18.0 (0.8) | 10.2 (0.6) |
|                 |         | Freyella cf fragilissima | E9  | 2  | −21.8 (0.5) | 17.5 (0.2) | 10.8 (0.2) |
|                 | Holothuroidea | Holothuroidea sp. | E9  | 3  | −24.9 (0.0) | 18.3 (0.6) | 8.0 (0.4) |
|                 | Ophiuroidea | Ophiuroidea sp. | E9  | 3  | −23.7 (0.8) | 17.6 (0.9) | 9.0 (0.8) |
| Vertebrata      | Osteichthys | Zoarcidae sp. | E9  | 4  | −26.5 (1.7) | 15.7 (0.4) | 12.0 (1.8) |

### Table 3. Mean $\delta^{13}$C, $\delta^{15}$N and $\delta^{34}$S values (%) of hydrothermal vent fauna collected from the E2 and E9 ridge segments of the East Scotia Ridge, Southern Ocean.

| Taxonomic group | E2 $\delta^{13}$C | $\delta^{34}$S | $\delta^{15}$N | E9N $\delta^{13}$C | $\delta^{34}$S | $\delta^{15}$N | E9S $\delta^{13}$C | $\delta^{34}$S | $\delta^{15}$N |
|-----------------|------------------|---------------|---------------|------------------|---------------|---------------|------------------|---------------|---------------|
| Cirripedia      | Vulcanolepas sp. | 22 – 21.1 (0.6) | 8.2 (1.0) | 6.3 (0.7) | 23 – 26.9 (0.8) | 11.0 (0.8) | 9.0 (0.4) | 23 – 22.1 (0.8) | 5.4 (1.1) | 6.4 (0.6) |
| Decapoda        | Kiwa sp. female | 20 – 19.4 (1.5) | 3.9 (1.3) | 8.2 (0.5) | 0 – – – | 0 – – – | 22 – 10.6 (0.9) | 4.0 (0.7) | 8.9 (0.5) | 30 – 10.7 (0.6) | 2.4 (0.9) | 9.1 (0.6) |
| Pycnogonida     | Sericosura spp. | 6 – 24.7 (0.9) | 11.9 (0.4) | 8.5 (1.3) | 9 – 30.9 (0.5) | 6.8 (0.8) | 7.7 (0.5) | 2 – 27.2 (0.3) | 14.9 (0.3) | 9.1 (0.1) |
|                 | Colossendeis cf concedis | 0 – – – | 6 – 23.3 (2.5) | 10.9 (1.4) | 12.1 (0.5) | 0 – – – | 22 – 10.6 (0.9) | 4.0 (0.7) | 8.9 (0.5) | 30 – 10.7 (0.6) | 2.4 (0.9) | 9.1 (0.6) |
| Anthozoa        | cf Actinostola sp. 1 | 0 – – – | 4 – 14.7 (3.5) | 10.3 (0.4) | 9.9 (0.3) | 0 – – – | 22 – 10.6 (0.9) | 4.0 (0.7) | 8.9 (0.5) | 30 – 10.7 (0.6) | 2.4 (0.9) | 9.1 (0.6) |
|                 | Pacmanactis sp. | 5 – 23.8 (0.2) | 14.9 (0.7) | 7.1 (0.7) | 0 – – – | 0 – – – | 22 – 10.6 (0.9) | 4.0 (0.7) | 8.9 (0.5) | 30 – 10.7 (0.6) | 2.4 (0.9) | 9.1 (0.6) |
|                 | cf Marianactis sp. | 5 – 23.7 (0.3) | 14.0 (2.4) | 7.2 (1.8) | 0 – – – | 0 – – – | 22 – 10.6 (0.9) | 4.0 (0.7) | 8.9 (0.5) | 30 – 10.7 (0.6) | 2.4 (0.9) | 9.1 (0.6) |
| Asteroidea      | Stichasteridae sp. | 1 – 20.2 | 11.3 | 12.3 | 5 – 12.2 (0.6) | 10.0 (0.9) | 12.4 (0.4) | 5 – 14.7 (1.6) | 11.8 (2.6) | 13.4 (0.6) |
| Gastropoda      | Peltopsioidea sp. | 19 – 30.1 (0.6) | 6.0 (0.6) | 5.4 (0.4) | 22 – 31.2 (0.4) | 3.7 (0.5) | 5.8 (0.6) | 15 – 30.1 (0.5) | 4.7 (1.1) | 5.9 (0.7) |
|                 | Peltopsioidea sp (≤7 mm) | 4 – 23.9 (0.7) | 7.4 (2.0) | 6.8 (0.5) | 5 – 29.6 (2.7) | 4.2 (0.2) | 6.4 (1.3) | 0 – – – | – – – | – – – |
|                 | Provannidae sp. 1 | 1 – 26.5 | 8.0 | 4.2 | 0 – – – | 0 – – – | 22 – 31.2 (0.4) | 3.7 (0.5) | 5.8 (0.6) | 15 – 30.1 (0.5) | 4.7 (1.1) | 5.9 (0.7) |
|                 | P–rovannidae sp. 2 | 0 – – – | 0 – – – | 0 – – – | 4 – 21.8 (1.4) | 5.9 (0.8) | 4.0 (0.5) | 22 – 31.2 (0.4) | 3.7 (0.5) | 5.8 (0.6) | 15 – 30.1 (0.5) | 4.7 (1.1) | 5.9 (0.7) |
| Cladorhizidae   | Cladorhiza sp. | 5 – 26.1 (0.4) | 14.7 (1.4) | 8.7 (0.4) | 0 – – – | 0 – – – | 5 – 9.9 (0.3) | 6.6 (0.2) | 5.2 (0.8) |
| Potential food sources | Particulate suspended material | 3 – 23.2 (5.4) | 10.0 (1.1) | 0.1 (4.9) | 0 – – – | 0 – – – | 5 – 9.9 (0.3) | 6.6 (0.2) | 5.2 (0.8) |
|                 | Rock scrapings | 0 – – – | 1 – 23.2 | 0.8 | 2.4 | 1 – 31.1 | 1.9 |

Standard deviations are in parentheses and - indicates no data.

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Other species of ESR vent macroconsumers, which had for bathymodiolid mussels, vesicomyid clams and other biogeographical vent provinces include some species of single strain of endosymbiotic gamma-Proteobacteria living in RuBisCO form I at all three locations. Molluscs containing a, 13C and 15N values living bacteria [14,23] so organic carbon fixed via other carbon fixation pathways cannot be ruled out as part of their assimilated diet.

Vent macroconsumers inhabiting the hottest areas of the hydrothermal vent tolerable to metazoan life, including rimicarid shrimps, polychaetes Alvinella spp. and Riftia pachyptila and some alvinococchid gastropods, tend to assimilate rTCA-fixed carbon from their diet [11,41,42] and have 13C values ≥−16‰ [14,25,43,44]. As 13C values is approximately 1‰ at the ESR sites, vent macroconsumers utilising carbon fixed via the rTCA cycle would have had 13C values ≥−13‰, assuming a −2‰ to −14‰ net fractionation between the inorganic substrate and organic product catalysed by the enzymes involved in the rTCA cycle [29,30]. Kiwa sp. living at E9N and E9S, along with its epibionts, had 13C values that were −12‰ and are found in areas close to discharging vent fluids [22]. This potentially indicates the epibions living on Kiwa sp. ventral setae were fixing carbon via the rTCA cycle. Kiwa sp. was also 15N-enriched by between 3.8‰ and 4.2‰ relative to its epibions, suggesting the epibionts were an important food source. A similar episymbiotic relationship between the ESR kiwiid is therefore hypothesised to that of Kiwa puravida, for which lipid, stable isotope and behavioural analyses indicate the harvesting of epibiont bacteria [43]. Stichasteridae sp. (−13‰) and cf. Actinostola sp. (−14‰) also appeared to be assimilating carbon indicative of the rTCA cycle at E9N and E9S.

Several vent macroconsumers fell within the range of 13C values indicative of mixed carbon sources. Those within the 13C −22‰ to −15‰ range may consume free-living bacteria or are predators or scavengers that utilise a number of trophic pathways. At the ESR hydrothermal vents, Lepetodrilus sp., Provannidae sp. 2, Vulcanelepas sp., Kiwa sp., Stichasteridae sp. and Colossendeis sp. fell into this range at one or more sites. Related species of Lepetodrilus sp., Provannidae sp. 2 and Vulcanelepas sp. are all thought to consume free-living bacteria at other vents sites [14,23]. Such feeding can result in consuming heterogeneous bacterial communities, which have multiple pathways for carbon fixation and elemental cycling [9,46,47]. The biological cycling of carbon is very complex at hydrothermal vents because of the multiple single carbon substrates for carbon fixation (e.g. CO2, CH4, CO), spatial variability in the 13C value of the substrate and various microbial primary producers associated with different carbon fixation pathways [7,11,40]. Furthermore, the incorporation of photosynthetic derived carbon as particulate or dissolved organic matter is possible and may provide some nutrition to vent macroconsumers [12,13]. Therefore, complex isotopic mixes of food sources are available to these species.

### Discussion

This study described the trophic structure of a new vent biogeographical province recently discovered on the ESR in the Southern Ocean [21]. In addressing this aim, the study shared the challenges of preceding work in characterising energy sources, separating isotopic overlap and mixing of energy sources, and following energy sources into subsequent predator-prey relationships. However, the tri-isotope approach and integration of both vent chemistry and microbiology, here, provided a more holistic understanding of vent trophic ecology at within- and amongst-vent field scales.

### Intra-site Trophic Interactions and Energy Sources

Scarcity of Δ13C estimates between inorganic carbon and cellular biomass for primary producers at hydrothermal vents [27,30] makes interpretation of the origin of organic carbon fixed within the hydrothermal vent system and assimilated by macroconsumers tentative for species not within a symbiotic or known within the hydrothermal vent system and assimilated by macroconsumers [27,30] makes interpretation of the origin of organic carbon fixed within the hydrothermal vent system [27,30]. Pelagia sp. containing an endosymbiotic gamma-Proteobacteria (K. Zwigirmaier unpublished data) and is within the Δ13C range expected for carbon fixed via RuBisCO form I at all three locations. Molluscs containing a single strain of endosymbiotic gamma-Proteobacteria living in other biogeographical vent provinces include some species of bathyomodiolid mussels, vesicomyid clams and Fimbria gastrapod, all of which have Δ13C values between −37‰ and −27‰ [39,40]. Other species of ESR vent macroconsumers, which had Δ13C values <−22‰ included Vulcanelepas sp. (E2 and E9S), Sericosa sp., E2 anemones and Lepetodrilus sp. These species consume free-living bacteria [14,23] so organic carbon fixed via other carbon fixation pathways cannot be ruled out as part of their assimilated diet.

| Species               | 13C  | 15N  | 15N  |
|-----------------------|------|------|------|
|                       | DF   | F    | p    | DF   | F    | p    | DF   | F    | p    | DF   | F    | p    |
| Vulcanolepas sp.      | 2    | 63   |     |      |      |      | 2    | 63   |     |      |      |      |
| Kiwa sp. male         | 2    | 31.36| 147.29|<0.01 |      |      | 2    | 63   | 176.16 |<0.01 |      |      |
| Sericosa sp.          | 2    | 15   | 215.00|<0.01 |      |      | 2    | 15   | 100.61 |<0.01 |      |      |
| Pelagia sp.           | 2    | 52   | 29.50|<0.01 |      |      | 2    | 52   | 49.26 |<0.01 |      |      |
| Lepetodrilus sp.      | 2    | 10   | 17.41|<0.01 |      |      | 2    | 10   | 31.99 |<0.01 |      |      |

Welch’s ANOVA with post hoc analysis by t-test with Bonferroni correction (p = 0.05/3 = 0.017). doi:10.1371/journal.pone.0065553.t004

| Species               | 13S  | 15N  | 15N  |
|-----------------------|------|------|------|
|                       | DF   | F    | p    | DF   | F    | p    | DF   | F    | p    |
| Vulcanolepas sp.      | 2    | 63   |     |      |      |      | 2    | 63   |     |      |      |      |
| Kiwa sp. male         | 2    | 31.36| 147.29|<0.01 |      |      | 2    | 63   | 176.16 |<0.01 |      |      |
| Sericosa sp.          | 2    | 15   | 215.00|<0.01 |      |      | 2    | 15   | 100.61 |<0.01 |      |      |
| Pelagia sp.           | 2    | 52   | 29.50|<0.01 |      |      | 2    | 52   | 49.26 |<0.01 |      |      |
| Lepetodrilus sp.      | 2    | 10   | 17.41|<0.01 |      |      | 2    | 10   | 31.99 |<0.01 |      |      |

Welch’s ANOVA with post hoc analysis by t-test with Bonferroni correction (p = 0.05/3 = 0.017). doi:10.1371/journal.pone.0065553.t005

Table 4. Results of ANOVA and post-hoc Tukey honest significant differences tests for the differences in stable isotope values of vent fauna among the three sites on the East Scotia Ridge.
The majority of ESR vent macroconsumers had δ34S values less than or equal to the 10‰ threshold, indicating chemosynthetic food sources [49]. Species exceeding the 10‰ value occurred mainly at E2 in the anemones *Pocanactis* sp. and cf *Marianactis* sp., the sponge *Cladorhiza* sp., the pycnogonids *C. elegans* and the stichasterid seastar along with *Sericosura* spp. and stichasterid seastar at E9S. All had δ34S values between 10‰ and 16‰. Mixing of epipelagic photosynthetic and hydrothermal vent chemosynthetic production sources at these sites cannot be ruled out.

Determining intra-site differences in food sources and trophic interactions using δ34S is challenging for macroconsumers with δ34S values <10‰ because the δ34S values of inorganic substrates and the net fractionation effect between inorganic substrates and products for primary producers and consumers are uncertain. At E9, δ34S appeared to increase from macroconsumers living closest to vent openings and within diffuse flow areas (i.e. *Kiwa* sp., *Peltospiroidea* sp. and *Leptodrilus* sp.) to those in the periphery (i.e. anemones, stichasterid seastars and *Colossendeis* spp.). It is unclear why an increase in δ34S occurred from the centre of the vent to the periphery: it may be the result of changes in sulfide speciation [50] or other sulfur sources with increasing distance from the vent opening [33], differences in levels of sulfide exposure [50], incorporation of epipelagic photosynthetic primary production or a combination of the above.

Stichasterid seastars, cf *Actinostola* sp. and *Colossendeis* spp. consistently had the highest δ15N values of all the ESR vent macroconsumers, which suggested they occupied the highest trophic positions of those predators sampled. Behavioural observations [22] and δ15N values indicated that *Kiwa* sp. is consumed by stichasterid seastar and cf *Actinostola* sp. 1 but only the stichasterid seastar had δ13C values indicative of a higher trophic position than *Kiwa* sp. In the case of *Colossendeis* spp., feeding on anemones occurs at the ESR vent sites [22] and at E2 all three stable isotopes indicated a strong predator-prey link. At E9N there was a large difference in δ13C and δ34S between cf *Actinostola* sp. 1 and the two species of *Colossendeis* as well as lower δ15N in these pycnogonids compared to cf *Actinostola* sp. 1. This suggests that at E9N the feeding incidents between cf *Actinostola* sp. 1 and *Colossendeis* spp. are either rare or stable isotopic values of *Colossendeis* spp. are strongly affected by isotopic mixing of different energy sources (δ13C and δ34S) and feeding over multiple trophic positions (δ15N).

It is evident from the ESR hydrothermal vent food webs that predators may have similar or lower δ13C values than their prey. Calculating trophic position assuming taxon specific nitrogen trophic discrimination factors [23] or applying the more universal value of 3.4‰ [12] was not undertaken within this study because they may have provided erroneous results. Establishing a suitable δ15N baseline is problematic because: the macroconsumer with the lowest δ15N differed among locations, is confounded by the use of different tissues (e.g. whole animals, muscle) to construct the food webs [32] and the observed high δ15N variability in potential food sources. Compound-specific amino acid stable isotope analysis may provide higher resolution information on the organic nitrogen compounds assimilated by vent macroconsumers because the isotopic values of different amino acids record trophic and basal source information [31,32]. Thus it may circumvent some of the limitations of bulk δ15N analysis and provide a better understanding of nitrogen cycling at hydrothermal vents.

**Spatial Patterns in Macroconsumer Trophodynamics**

Large spatial differences in δ13C values for *Kiwa* sp., *Stichasteridae* sp. and *Sericosura* spp. were attributed primarily to differences in carbon fixation pathways at the base of the food web, which is in turn transferred to higher trophic positions. δ15N values of *Kiwa* sp. differed by ~9‰ between E2 and E9S as did that of associated *Kiwa* sp. epibionts. Also, epsilon-Proteobacteria dominated the epibiont community at E9 with gamma-Proteobacteria largely absent, compared to a mix of gamma- and epsilon-Proteobacteria at E2 [K. Zwirglmaier unpublished data]. All epsilon-Proteobacteria to date use the rTCA cycle to fix carbon while gamma-Proteobacteria predominantly use the CBB cycle [11]. *Riftia pachyptila* has similar differences in δ13C among vent sites, but this is attributed to its endosymbionts shifting between rTCA and CBB cycles [53] rather than changes in the microbial community it consumes. Alvinocochid gastropods have δ13C values that differ by >20‰ among vent fields, which relates to whether epsilon- or gamma-Proteobacteria are the endosymbionts [54]. It is unclear why *Kiwa* sp. epibiont diversity is different between E2 and E9. At other hydrothermal vent locations differences in vent fluid chemical composition influences microbial communities [46,55] and it may be similar at the ESR vent fields. The difference in carbon fixation appeared to be transferred through *Kiwa* sp. to the predatory stichasterid seastar. Such a predator-prey interaction may also explain the large difference in δ13C values between E2 and E9N in *Sericosura* spp. At E2 *Sericosura* spp. were collected from amongst anemones that had δ13C values indicative of a mixed carbon source but at E9 they were collected from amongst peltiopiod gastropods dependent on CBB fixed carbon, although *Sericosura* spp. were not observed directly feeding on either anemones or *Peltospiroidea* sp.

**Table 5.** Results of t-tests for between-sites differences in stable isotope values of vent fauna at the East Scotia Ridge.

| Species               | Comparison       | $\delta^{13}$C | DF | t   | p   | $\delta^{34}$S | DF | t   | p   | $\delta^{15}$N | DF | t   | p   |
|----------------------|------------------|----------------|----|-----|-----|----------------|----|-----|-----|----------------|----|-----|-----|
| *Kiwa* sp.           | E2 female v male | 36             | −0.50 | 0.62 | 36 | 2.23          | <0.05 | 36 | 5.13 | <0.01         |
| *Kiwa* sp. Epibions  | E2 v E9S         | 4.05           | −3.81 | <0.05 | 6.93 | 4.57          | <0.01 | 2  | na   | <0.05         |
| *Stichasteridae* sp. | E2 v E9N         | 4              | 28.92 | <0.01 | 4  | −3.29         | <0.05 | 4  | 5.14 | <0.01         |
| *Stichasteridae* sp.| E2 v E9S         | 4              | 7.28   | <0.01 | 4  | 0.385         | 0.59  | 4  | 5.94 | <0.01         |
| *Sericosura* sp.     | E9N v E9S        | 8              | 3.64   | <0.01 | 8  | −1.76         | 0.11  | 8  | −3.49 | <0.05         |

*Welch’s t-test, Wilcoxon test,*

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Relatively small differences in stable isotope values were observed among sites in Peltopsioideae sp., Lepidodrilus sp. and Valamolopas sp. To date, Peltopsioideae sp. contains a single strain of gamma-Proteobacteria endosymbiont (K. Zwirglmaier unpublished data), which means spatial differences in δ13C and δ34S are unlikely to be the result of differences in the type of endosymbiont [35]. The differences were potentially a result of site-specific variations in the δ13C.inc and inorganic δ34S values used by the endosymbionts during chemosauotrophy or physiological temperature-related effects on isotopic discrimination. Small differences among sites for the grazer Lepidodrilus sp. and suspension feeder Valamolopas sp. are harder to explain because of the various factors that are likely to influence their food source. δ13C values indicated these organisms consume a mixed diet of free-living microbes and particulate material. However, differences in δ13C values within sites may be related to the organism’s distribution within the vent field [56] and in turn the composition of the microbial community [46], the stable isotope values of the inorganic substrate used during chemosauotrophy [57] and temperature effects on trophic discrimination. Lepidodrilus sp. and Valamolopas sp. were collected from single points within each vent site and, therefore, it is not clear whether the difference in stable isotope values among sites is greater or less than that within sites.

Because of the snap-shot nature of this study, it is difficult to identify factors that caused the spatial differences in the Kioa sp. epibiont communities that resulted in a greater range of δ13C values at the E9 sites compared to E2. Higher concentrations of dissolved sulfides in vent fluids may favour the rTCA pathway resulting in increasing numbers of organisms with δ13C values greater than −16‰ [12]. On the ESR, E9 has higher hydrogen sulfide and lower chloride concentrations than E2 meaning that E9 has greater concentrations of available gases for microbial primary production due to phase separation [6,21]. Higher concentrations of reduced compounds and gases may be one of the drivers of the differences in trophic structure at the ESR vents. However, hydrothermal vent communities also undergo changes in community composition with age [58] and fluctuating hydrothermal activity [59], which will have an effect on trophic structure. As data presented here were obtained concurrently with the discovery of the new biogeographical province it is not possible to determine whether the communities at E2 and E9 represent different successional stages, are a product of varying chemistry or a mix of such processes.

Conclusion

Trophic structure differed substantially between the E2 and E9 vents fields, and only slightly between E9N and E9S. δ13C.inc of the end-member fluid and diffuse flow samples were similar among the sites but large differences in the δ13C values of some vent macroconsumers indicated spatial variations in the way microbes were fixing carbon at the base of the food chain. δ13C values >−13‰ at the E9N and E9S suggest that the relative contribution to the macroconsumer food web of carbon fixed via the rTCA cycle is likely to be greater than at E2. The greater range of δ34S values at E2 and E9S indicated a potentially greater influence of epipelagic photosynthetic primary production than at E9N. The greater contribution of rTCA fixed carbon at the E9 vent field may ultimately be related to differences in vent fluid, but more work is required to link vent fluid chemistry with microbial primary production and the related trophic structure at hydrothermal vents.

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Author Contributions

Conceived and designed the experiments: WDKR CJS BDW KZ JAH KL. NVCP. Performed the experiments: WDKR CJS KZ JAH KL. Analyzed the data: WDKR CJS BDW KZ JAH RARM NVCP. Contributed reagents/materials/analysis tools: WDKR CJS KZ JAH RARM. Wrote the paper: WDKR CJS BDW KZ JAH RARM NVCP. Participants on JC042: WDKR CJS KZ JAH KL.

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