2 3 **Regional-scale** variability benthic in the of response 4 macroinvertebrate assemblages to a marine heatwave 5 6 Dan A, Smale^{1,2*}, Thomas Wernberg² and Mathew A. Vanderklift³ 7 8 ¹Marine Biological Association of the United Kingdom, The Laboratory, Citadel Hill, 9 Plymouth, PL1 2PB, United Kingdom 10 ²UWA Oceans Institute and School of Plant Biology, University of Western Australia, 35 11 Stirling Highway, Crawley 6009 WA, Australia 12 ³ CSIRO Oceans & Atmosphere, Floreat, Western Australia 6014, Australia 13 14 15

Research article to Marine Ecology Progress Series - accepted 04/02/2017

16

1

17 Running head: extreme warming event and temperate reef fauna

1 ABSTRACT: Extreme climatic events are predicted to increase in severity as a consequence 2 of anthropogenic climate change. In marine ecosystems, the importance of marine heatwaves (MHWs) – discrete periods of anomalously high sea temperatures - is gaining recognition. In 3 4 2011, the highest-magnitude MHW ever recorded impacted the west coast of Australia (southeast Indian Ocean). The MHW was associated with widespread mortality of habitat-5 6 forming species, including corals and kelps, and structural changes in assemblages of macroalgae and fish. However, the responses of benthic macroinvertebrate assemblages have 7 8 not yet been fully documented. Here, we resurveyed 2 subtidal habitat types (reef 'flats' and 9 'slopes') at 4 locations (spanning >800 km of coastline and >6° of latitude) during the period 1999-2015 to examine the impacts of the 2011 MHW on herbivorous macroinvertebrates (i.e. 10 sea urchins, gastropod molluscs). Responses to the MHW varied with latitude; at our warmest 11 12 study location abundances were severely depleted whereas no effects were detected at the 13 coolest location. Across the entire study region subtle but significant shifts in assemblage structure were observed due to decreased abundances of more southerly-distributed species 14 15 (i.e. 'cool' affinity) and increased abundances of several more northerly-distributed species (i.e. 'warm' affinity). The 2011 MHW has had profound effects on the marine biota off the 16 west coast of Australia, across multiple trophic levels and taxonomic groups. Here, as in many 17 other regions, contemporary warming events are superimposed onto gradual warming trends, 18 increasing the likelihood of abrupt changes in ecosystem structure and functioning. 19

KEY WORDS: Benthic herbivores; Extreme climatic events; Leeuwin Current; Mobile
macro-invertebrates; Southeast Indian Ocean; Southwest Australia; Species distributions;
Temperature variability

1 INTRODUCTION

Climatic variability, including the frequency and magnitude of extreme climatic events, is 2 predicted to increase as a direct consequence of anthropogenic climate change (Meehl & 3 4 Tebaldi 2004, Rahmstorf & Coumou 2011, IPCC 2012). Natural climate variability is now superimposed onto decadal warming trends in most regions, increasing the likelihood of 5 discrete climatic events becoming 'extreme' or 'anomalous' (Hansen et al. 2012, Trenberth 6 7 2012). Such events, which include heatwaves, droughts, storms and floods, can affect both terrestrial and marine ecosystems and cause high mortality (Garrabou et al. 2009, Marba & 8 9 Duarte 2010) deleterious impacts on populations (Van De Pol et al. 2010, Smale & Wernberg 2013) and a reconfiguration of communities (Thibault & Brown 2008, Wernberg et al. 2013, 10 Wernberg et al. 2016). 11

Marine heatwaves (MHWs) are broadly defined as discrete prolonged periods when sea 12 13 temperatures are anomalously high relative to long-term records (Hobday et al. 2016). MHWs, which can be caused by a range of oceanographic and atmospheric processes, are gaining 14 recognition as widespread and potent drivers of change in marine ecosystems (Wernberg et al. 15 2013, Hobday et al. 2016). Indeed, the number of days of anomalously high seawater 16 temperatures has significantly increased along 30% of the world's coastlines in the last 30 years 17 (Lima & Wethey 2012), while several 'high profile' warming events have had far-reaching 18 ecological impacts (Hobday et al. 2016). For example, the European meteorological heatwaves 19 of 2003 and 2006 elevated seawater temperatures in the Mediterranean Sea, which in turn 20 caused widespread mortality, shifts in species' distributions and declines in local marine 21 biodiversity (Garrabou et al. 2009, Lejeusne et al. 2009, Marba & Duarte 2010). More recently, 22 the El Niño-driven warming event of 2015-2016 has devastated coral reefs at a global scale 23 (Normile 2016). It is clear that prolonged periods of extremely high seawater temperatures 24

affect processes across all biological scales, from genes (Bergmann et al. 2010) to organisms
 (Diaz-Almela et al. 2007) to ecosystems (Wernberg et al. 2016).

In the austral summer of 2010/2011, the southeast Indian Ocean experienced an extreme 3 warming event, during which seawater temperatures were the highest on record (~30 years for 4 satellite-derived SSTs and ~140 years for reconstructed SSTs, see Wernberg et al. 2013). 5 6 During the MHW, warming anomalies of 2-4°C persisted for around two months across >2000 km of temperate and subtropical coastline (Feng et al. 2013, Pearce & Feng 2013, Wernberg et 7 8 al. 2013). At the MHW's peak in late February/March 2011, warming anomalies of up to 5°C were observed at multiple coastal locations (Rose et al. 2012, Feng et al. 2013, Pearce & Feng 9 2013). The MHW was associated with unusually strong La Niña conditions, which increased 10 the flow of the region's main ocean current (the Leeuwin Current, 'LC') and the transfer of 11 tropical warm water polewards, and was superimposed onto a decadal scale warming trend in 12 the southeast Indian Ocean (Pearce & Feng 2007). 13

14 The MHW had wide-ranging consequences for marine ecosystems along the western coastline of Australia, which is a global hotspot of marine diversity and endemism (Tittensor et al. 2010, 15 Bennett et al. 2015a). Unprecedented rates of coral bleaching and mortality were recorded 16 across >1000 km of tropical and subtropical coastline (Moore et al. 2012, Depczynski et al. 17 2013), including at high latitude locations that are historically resistant to bleaching events 18 19 (Smale & Wernberg 2012). Significant declines in the abundance and geographical extent of habitat-forming macroalgae were observed (Smale & Wernberg 2013, Wernberg et al. 2013, 20 21 Wernberg et al. 2016), as were changes in fish abundances and species composition (Wernberg 22 et al. 2013, Bennett et al. 2015b, Wernberg et al. 2016) and mass mortalities of commerciallyimportant finfish and shellfish (Pearce et al. 2011, Caputi et al. 2016). 'Warm-temperate' 23 locations situated within the tropical-temperate transition zone were profoundly impacted, as 24 25 they suffered widespread loss of cool-water adapted habitat-forming species (kelps and large fucoids), which were unable to cope with the extreme temperatures experienced during the
 MHW (Smale & Wernberg 2013, Wernberg et al. 2013, Wernberg et al. 2016).

Mobile macroinvertebrates such as echinoderms and molluscs play key roles in the trophic 3 ecology of temperate Australia (Shepherd & Edgar 2013), yet the impact of the 2011 MHW on 4 their distributions and abundances has not yet been analysed in detail. Within the current 5 biogeographical context of southwestern Australia, mobile macro-invertebrates are fairly low 6 in diversity and abundance compared with many other temperate and polar ecosystems, and 7 8 exhibit highly patchy spatial distributions (Vanderklift & Kendrick 2004, Wernberg et al. 2008, Levitus et al. 2012, Azzarello et al. 2014, Smale & Wernberg 2014). Despite their relatively 9 low diversity and abundances, and an apparent lack of grazing 'fronts' and urchin 'barrens' in 10 the region, densities can be locally high (>8 large inds.m⁻², see Vanderklift & Kendrick 2004, 11 Azzarello et al. 2014) and mobile macroinvertebrates represent a conspicuous and 12 characteristic component of kelp forest communities (Vanderklift & Kendrick 2004, Wernberg 13 et al. 2008, Azzarello et al. 2014). Moreover, key species of sea urchins and molluscs may play 14 15 a critical role in the food web (Vanderklift et al. 2006, Lozano-Montes et al. 2011, MacArthur et al. 2011), linking primary productivity (e.g. drifting and attached macroalgae) to higher 16 trophic levels (e.g. lobsters, finfish). 17

Off southwest Australia, the reef-associated benthic macroinvertebrate fauna has a 18 19 predominantly temperate affinity (Vanderklift & Kendrick 2004); the most abundant sea urchin in the region is the purple sea urchin Heliocidaris erythrogramma (Valenciennes 1846, 20 21 hereafter 'Heliocidaris'), which is widely distributed across southern Australia (Keesing 2001, 22 Smale & Wernberg 2014). The sea urchin Phyllacanthus irregularis (Mortensen 1928, hereafter 'Phyllacanthus'), and the large turbinid gastropod Lunella torquatus (Gmelin 1791, 23 recently synonymised with Turbo torquatus, hereafter 'Lunella') are also common, widespread 24 25 and have cool-temperate affinities. The sea urchin Centrostephanus tenuispinus (Clarke 1914,

1 hereafter 'Centrostephanus') has a warm-temperate distribution (Wernberg et al. 2016), while 2 tropical warm-water species such as the sea urchin *Tripneustes gratilla* (Linneaus 1758, hereafter 'Tripneustes') and the cowry gastropod Monetaria caputserpentis have occasionally 3 4 been recorded within kelp forest communities in southwestern Australia (authors pers. obs.). The broad-scale affinities and geographical distributions of common macroinvertebrates 5 6 observed in previous surveys are shown in Table 1. As such, inter-specific variation in biogeographic and thermal affinities may make some populations more susceptible to 7 temperature variability, such as that experienced during the 2011 MHW. 8

Here, we conducted geographically extensive surveys and used historical data to test the 9 following hypotheses (1) that the MHW significantly altered the structure of macroinvertebrate 10 assemblages on subtidal reefs in southwest Australia. We also hypothesised (2) that the 11 abundances of more southerly-distributed cool-temperate species would be lower after the 12 MHW, especially at our warmest study locations where thermal physiological tolerances may 13 have been exceeded. Conversely, we predicted that (3) the abundances of more northerly-14 15 distributed warm-temperate/tropical macroinvertebrates would be higher after the MHW, particularly at our warmest study locations situated within the tropical-temperate transition 16 17 zone.

18 MATERIALS AND METHODS

19

20 Study region

21

The extensive coastline of southwestern Australia is characterized by widespread subtidal rocky reef habitat that supports highly productive, diverse and spatially extensive benthic communities, which are generally dominated by the kelp *Ecklonia radiata* in shallow waters (i.e. <30 m depth). We examined the abundances of benthic macroinvertebrates on kelpdominated rocky reef habitats within 4 locations off southwest Australia; Hamelin Bay (34.2°S,

1 115.0°E), Marmion Lagoon (31.8°S, 115.7°E), Jurien Bay (30.2S, 115.0°E) and Kalbarri 2 (27.4°S, 114.1°E). Adjacent locations were situated >200 km apart (Fig. 1) and the study encompassed ~6° latitude and ~800 km of southwest Australian coastline (southeast Indian 3 4 Ocean). All locations were moderately exposed to the oceanic swell systems that influence the ecology and geomorphology of the region (Searle & Semeniuk 1985, Smale et al. 2011). The 5 6 study locations encompassed a temperature gradient of ~3°C and fall within a larger regionalscale oceanic temperature gradient that characterizes the west coast of Australia (Fig. 1). 7 8 Average summer sea temperatures ranged from 20.3°C at Hamelin to 23.2°C at Kalbarri (see 9 Smale & Wernberg 2009, for detailed climatology of the region). The coastline is strongly influenced by the LC which originates in the Indo-Pacific and flows polewards along the coast, 10 before deviating eastwards into the Great Australian Bight (Pearce 1991, Smith et al. 1991). 11 12 The LC transports tropical (and subtropical) dispersal stages of marine flora and fauna and warm, nutrient-poor water polewards (Ayvazian & Hyndes 1995, Caputi et al. 1996, Smale & 13 Wernberg 2009). 14

15

16 Field surveys

For each location, existing data on the abundance of mobile invertebrates were collated from 17 published studies (Vanderklift & Kendrick 2004, Wernberg et al. 2008) and from authors' 18 19 unpublished surveys that used identical survey methods. These studies were used to identify sites within each location that could be resurveyed to assess the impacts of the MHW. Multiple 20 21 comparable study sites, >1 km apart from one another, were selected at random from a larger 22 possible pool for resurveying. All study sites were characterised by extensive limestone reef habitats, at 6-16 m depth, and supported benthic assemblages typical of the wider region (Smale 23 et al. 2010). Two habitat types were defined a priori; flat reef platforms (hereafter 'flats') and 24 25 vertical or steeply-sloping rock faces (hereafter 'slopes'). These habitat types were treated

1 separately because (i) they support distinct invertebrate assemblages (Vanderklift & Kendrick 2004), and (ii) the quantity of available data and the most suitable study sites for resurveying 2 differed between habitat types. For reef flats, 5 sites were selected from each of the 4 locations 3 4 for resurveying, whereas reef slopes were resurveyed at 3 sites within 3 locations (existing data were not available for reef slopes at Kalbarri). Before the MHW, sites were surveyed between 5 6 1 and 3 times between 1999 and 2006 (Table S1). After the MHW, new targeted surveys were conducted for the current study at all sites 3 times (in 2013, 2014 and 2015), with the exception 7 of some sites at Hamelin which were not surveyed in 2014/2015 (Table S1). All surveys were 8 9 conducted during the austral summer (full details provided in Table S1). Previous research in the region has shown that short-term variability (i.e. seasons to years) in invertebrate 10 assemblage structure is minimal and that densities of dominant macroinvertebrates are 11 generally stable over periods of months to years (Vanderklift & Kendrick 2004, Smale & 12 Wernberg 2014). For example, an examination of Heliocidaris abundances from 3 consecutive 13 pre-MHW survey years (between 1999 and 2001) on reef slopes at Marmion indicated minimal 14 15 inter-annual variability (Fig S1). There was no discernible intensification of localized anthropogenic stressors, such as increased pollution, sedimentation or harvesting, which may 16 have confounded any effects of the MHW, at any of the locations during the study period. 17 Human populations in nearby settlements are relatively small (with the exception of the Perth 18 Metropolitan Area adjacent to our sites at Marmion, which fall within a designated Marine 19 20 Park) and localized anthropogenic impacts that could potentially confound temperature effects were deemed to be minimal. 21

All mobile macroinvertebrates (>20 mm) within 5 replicate 5 x 1 m belt transects were counted on SCUBA (by the authors) at each study site. Transects were positioned haphazardly and placed >5 m apart from one another. In total, counts were obtained from 685 transects (395 completed before the MHW and 290 after the MHW) covering \sim 3425 m² of subtidal reef habitat (~1975 m² before the MHW and ~1450 m² after the MHW).

1 Statistical analysis

2 Differences in invertebrate assemblage structure between pre-MHW and post-MHW surveys was examined with permutational multivariate analysis of variance (PERMANOVA, see 3 Anderson 2001), using PRIMER 6 software (Clarke & Warwick 2001) with the 4 PERMANOVA add-on (Anderson et al. 2008). As macroinvertebrate abundance values per 5 transect were often low, the five transects completed per site/year combination were first 6 pooled (to generate abundance values per 25 m^2) so that a single value was used for each site-7 8 year combination. Initially a 'global' analysis was performed on data from all locations, using an orthogonal model with the two factors 'location' (fixed factor) and 'MHW' (fixed factor); 9 each habitat type was analysed separately. Permutations were based on a similarity matrix 10 generated from Bray-Curtis similarity matrix of square-root transformed pooled densities 11 (4999 permutations under a reduced model). As highly-significant (P<0.001) interactions 12 between location and MHW were detected for both habitat types (Table S2, Fig. S2), separate 13 a priori planned contrasts for each location were conducted to test the prediction that years 14 15 following the MHW would be distinct from those before the MHW (using the same similarity matrix and data transformation as above, and 4999 unrestricted permutations). In all cases, 16 dummy variables (equal to the lowest transformed abundance value; '1') were included in the 17 similarity matrices to alleviate the overpowering influence of transects with zero abundance 18 values (Clarke & Warwick 2001). Where a significant difference was detected, a SIMPER 19 analysis was performed to determine which taxa contributed most to the observed dissimilarity. 20 PCO plots for each location and habitat type were constructed to examine multivariate 21 partitioning before and after the MHW. 22

Temporal trends in total abundance (TA), taxon richness (TR) and the abundances of dominant
species were examined with univariate permutation-based ANOVA (Anderson et al. 2008),
using the planned contrasts described above (all response variables exhibited a significant

Location x MHW interaction term in initial global analyses and so each location was analysed
 separately). Permutations were based on Euclidean distances between untransformed
 abundance data (using 4999 unrestricted permutations).

4 **RESULTS**

5 The marine heatwave of 2010/11

At all locations temperature anomalies between +2°C to +3°C persisted for two months (Fig.
1) and absolute sea surface temperatures in the region were the highest on record (Feng et al.
2013, Pearce & Feng 2013, Wernberg et al. 2013). In addition to the extreme warming observed
in 2010/2011, sea surface temperatures in early 2012 and (to a lesser extent) early 2013 were
also higher than the climatological mean for each location (Fig. 1).

11 Assemblage-level responses

On reef flats, the composition of mobile invertebrate assemblages was not obviously impacted 12 13 by the MHW at the coolest study locations (PCO plots showed no clear partitioning in composition before and after the MHW at Hamelin or Marmion: Fig. 2) but were clearly altered 14 by the MHW at the warmest locations (partitioning was evident at Jurien and major shifts in 15 16 composition occurred at Kalbarri: Fig. 2). Multivariate statistical tests supported inferences from visual inspections of PCO plots, as *a priori* planned contrasts indicated that pre- and post-17 MHW assemblages at Jurien and Kalbarri were significantly different (Table S3). SIMPER 18 19 analysis indicated that the observed dissimilarities at Jurien were principally related to lower post-MHW abundances of Heliocidaris and Lunella and higher abundances of 20 Centrostephanus (Table S4). At Kalbarri, the gastropods Lunella, Dicathais orbita and 21 Astralium spp., which were not recorded after the MHW, were the principal contributors to the 22 observed dissimilarities between pre and post-MHW assemblages (Table S4). 23

1 On reef slopes, PCO plots indicated that assemblages at Hamelin showed no obvious 2 differences in composition before and after the MHW, whereas partitioning between pre- and post-MHW assemblages was observed at Marmion and Jurien (Fig. 3). These observations 3 4 were supported by PERMANOVA, as pre and post-MHW assemblages at Hamelin were statistically similar but significant differences in composition were observed at Marmion and 5 6 Jurien (Table S5). SIMPER analysis indicated that differences at Marmion were related to higher post-MHW abundances of Heliocidaris and Centrostephanus and lower abundances of 7 the sea star Petricia vernicina (Table S6). At Jurien, the sea urchins Centrostephanus, 8 9 Tripneustes (both with higher abundances post-MHW) and Phyllacanthus (lower abundances post-MHW) were the principal contributors to the observed dissimilarities between pre and 10 11 post-MHW assemblages (Table S6).

On reef flats total abundance (TA) and taxon richness (TR) did not vary significantly between 12 pre and post-MHW years at the 3 highest latitude locations (Hamelin, Marmion and Jurien, see 13 Fig 4). At Kalbarri, however, TA and TR were significantly lower after the MHW (Table S7) 14 15 to the extent that not a single macroinvertebrate individual was observed in any of the 75 post-MHW transects (Fig. 4), which covered a habitat area of $\sim 375 \text{ m}^2$. On average at Kalbarri, TA 16 decreased from 2.3 ± 0.8 to 0 inds.25 m⁻² and TR decreased from 1.3 ± 0.3 to 0 spp.25 m⁻² (Fig. 17 4). On reef slopes, TA did not differ significantly before and after the MHW at any location 18 but TR was significantly lower at Jurien after the MHW (Table S8), decreasing from 5.0 ± 0.6 19 to 2.9 ± 0.3 spp.25 m⁻² (Fig. 4). 20

21 **Population-level responses**

On reef flats, *Heliocidaris* was the most abundant macroinvertebrate, reaching a maximum average abundance of 8.8 ± 3.0 inds. $25m^{-2}$ at Jurien (Fig. 5). The MHW had no statistically significant effect on *Heliocidaris* abundances at Hamelin, Marmion and Jurien, and it was not recorded at Kalbarri during any survey year (Fig. 5, Table S9). The two most common gastropod taxa, *Lunella* and *Astralium* spp. (*Astralium tentorium* Thiele 1930 and *Astralium squamiferum* Koch 1844) were recorded at all locations before the MHW but were not recorded
at the warmest location, Kalbarri, after the MHW (Fig. 5). This marked post-MHW decline
was statistically significant for *Lunella* but not for *Astralium* (Fig. 5, Table S9).

On reef slopes, *Heliocidaris* was again the most common macroinvertebrate and did not 5 respond significantly to the MHW (Fig. 6, Table S10). At Jurien, the abundance of the pencil 6 urchin Phyllacanthus was significantly lower after the MHW (Fig. 6, Table S10). The most 7 8 striking observation was the marked increases in the abundance of *Centrostephanus* at both Marmion and Jurien (Fig. 6). Following the MHW, the average abundance of *Centrostephanus* 9 was ~15 times higher at Jurien and also significantly higher at Marmion, increasing from 10 complete absence in transects before the MHW to an average abundance of 1.9 ± 0.7 inds. $25m^2$ 11 following the event (Fig. 6, Table S10). A marked but statistically non-significant increase in 12 Centrostephanus abundance was also observed at Hamelin (Fig. 6, Table S10). The collector 13 urchin *Tripneustes* was not recorded in any pre-MHW transect but was recorded at some sites 14 15 at Jurien after the MHW, in 2013 and 2014 (Fig. 6). However, the planned contrast between pre- and post-MHW years was not statistically significant because of high variability between 16 sites and years (Tripneustes was recorded at 2 sites in 2013, 1 site in 2014 and was absent in 17 2015), indicating that patterns were variable between sites and years. 18

Temporal shifts in the relative abundances of sea urchin species on reef slopes (i.e. the percentage of all sea urchin individuals represented by each species, with all 3 sites per location pooled) were also examined (Fig. 7). This analysis showed a consistent clear pattern of higher relative abundance of *Centrostephanus* since the MHW at all 3 study locations (Fig. 7). For example, at Jurien before the MHW *Centrostephanus* represented 3.5% of all sea urchins recorded, yet by 2015 (4 years after the MHW) *Centrostephanus* represented 90.1% of all sea urchin individuals. A similar trend was observed at Hamelin and Marmion (Fig. 7). Furthermore, at Jurien the relative contributions of *Heliocidaris* and *Phyllacanthus* individuals to the sea urchin assemblage was markedly lower following the MHW, partly as a consequence of the higher abundances of *Tripneustes* (temporarily) and *Centrostephanus* (Fig. 7). The relative abundances of *Heliocidaris* and *Phyllacanthus* at the other study locations were more variable between years and showed no clear trend (Fig. 7).

6

7 **DICUSSSION**

8

The 2011 MHW was extreme in terms of magnitude, duration and spatial extent (Feng et al. 9 2013, Wernberg et al. 2013). Our data unequivocally demonstrate that the MHW significantly 10 altered the composition of benthic macroinvertebrate assemblages on subtidal reefs in 11 12 southwest Australia, with the magnitude of impact inversely related to latitude (i.e. the warmest locations were the hardest hit). At the coolest study location, Hamelin, the composition of 13 invertebrate assemblages on reef flats and slopes did not differ between pre and post-MHW 14 15 years. At the mid-latitude locations, Marmion and Jurien, changes in the relative abundances of macroinvertebrate taxa led to significant alterations in species composition (on reef slopes 16 at Marmion and on both habitat types at Jurien), whereas major shifts in species composition 17 were observed at the lowest latitude location, Kalbarri. This aligns with the responses of fish 18 and macroalgae assemblages (Wernberg et al. 2013), which were impacted by the MHW at a 19 20 'warm' location (Jurien) but not at a 'cool' location (Hamelin). These data support our first 21 hypothesis, that macroinvertebrate assemblage structure in southwest Australia was altered by 22 the 2011 MHW, although responses varied considerably between locations.

23

The most striking observation of the current study was the decimation of all benthic macroinvertebrates at Kalbarri, which were completely absent after the MHW. The most common mobile invertebrates at Kalbarri before the MHW were all large gastropods (the turban shells *Lunella* and *Astralium* spp. and the carnivorous muricid *Dicathais orbita* Gmelin

1 1791) which, although not abundant, were commonly recorded before the MHW, being 2 ubiquitous in all surveys prior to 2011. These species exhibit temperate distributions spanning southern Australia, having likely evolved under cool, climatically-stable Tethyan conditions 3 4 (Williams 2007). Kalbarri is situated towards the equatorward limit of these species' distributions and, although thermal tolerances for these species are unknown, it is very likely 5 6 that extreme temperatures experienced during the MHW had direct adverse physiological effects and induced high mortality rates. During the MHW, there was 99% mortality of the 7 8 commercially-important gastropod Halliotis roei (Gray 1826, 'Roe's abalone') on inshore reefs 9 at Kalbarri, which represents the equatorward limit of this species' distribution (Caputi et al. 2016). The mass die-offs at Kalbarri were associated with 30°C temperatures, discoloured 10 water and (probably) depleted oxygen levels (Pearce et al. 2011), and deleterious impacts of 11 12 warming on *H. roei* populations further south in the Perth Metropolitan Area (i.e. near 13 Marmion) were also observed (Caputi et al. 2016). Lunella also decreased in abundance after the MHW at Marmion and Jurien (although not significantly), providing further support for the 14 15 susceptibility of range edge populations to extreme warming.

16

17 In addition to direct thermal stress, it is possible that the indirect effects of loss of habitat and food also affected invertebrate populations at Kalbarri and, to a lesser extent, Jurien. The MHW 18 had direct adverse effects on habitat-forming seaweeds such as the dominant kelp Ecklonia 19 radiata, which resulted in a 30-40% decline in total canopy cover in Jurien (Wernberg et al. 20 2013) and the extirpation of the large fucoid Scytothalia dorycarpa at its range edge (Smale & 21 Wernberg 2013). At Kalbarri, habitat structure was dramatically impacted by the MHW, as the 22 spatial coverage of the canopy-forming kelp Ecklonia radiata decreased from ~75% of the 23 reef's surface (Wernberg et al. 2010) to complete absence after the MHW (Wernberg et al. 24 2016). It has been shown that *Lunella* has a high affinity with kelp cover, and that sharp 25 declines in Lunella abundance are associated with loss of kelp during ENSO events on the East 26

1 coast of Australia (Ettinger-Epstein & Kingsford 2008). Although Lunella and Astralium spp. 2 do not primarily feed on kelp, preferring to consume filamentous or foliose algae (Ettinger-Epstein & Kingsford 2008, Wernberg et al. 2008), canopy-forming macroalgae represent a 3 4 critical resource through shelter provision (Ettinger-Epstein & Kingsford 2008). As such, loss of structural habitat may have indirectly impacted gastropod abundance, as loss of macroalgal 5 6 shelter can induce behavioural responses and increase vulnerability of invertebrates to predators (Ettinger-Epstein & Kingsford 2008, Stoner 2009). Moreover, drifting kelp 7 fragments are an important food source for benthic macroinvertebrates in the region 8 9 (Vanderklift & Wernberg 2010), so a decline in food availability may have affected the ecological performance of individuals and the structure of populations. 10

11

12 The post-MHW surveys were conducted 2-4 years after the 2011 event, with invertebrate 13 populations at Kalbarri showing no signs of recovery following apparent widespread mortality. This is unsurprising, as recovery of the gastropod populations formerly recorded at Kalbarri 14 15 will likely be hampered for 2 reasons; (1) the larval duration for some of these species is presumed to be fairly short (i.e. days to weeks) (Phillips 1969, Joll 1980), so that post-MHW 16 recovery will depend on proximity to source populations, and (2) source populations are likely 17 to be located at higher latitudes, downstream of the main poleward-flowing ocean current, 18 thereby reducing the likelihood of larval transport into impacted locations (Caputi et al. 1996). 19 20 As Kalbarri is situated towards the equatorward distribution limits for these cool-water species, it is possible that the MHW has induced poleward range contractions (as with a temperate 21 seaweed, see Smale & Wernberg 2013, Wernberg et al. 2016), although additional surveys are 22 needed to confirm species' range shifts. In addition to gastropods at Kalbarri, abundances of 23 the pencil urchin *Phyllacanthus* were markedly lower on reef slope habitats at Jurien after the 24 MHW, which was a major contributor to the observed dissimilarity between pre and post-25 MHW assemblages. Phyllacanthus has a cool-water temperate distribution with its 26

equatorward range edge estimated at the Abrolhos Islands, ~28.5°S (Marsh 1994). As such, its 1 2 absence at Kalbarri and post-MHW decline at Jurien may be related to recent warming, but thermal tolerance experiments are needed to examine this further. However, we observed no 3 4 clear trend in the abundance of *Heliocidaris*, which has a similar temperate distribution to Phyllacanthus, therefore indicating inter-specific variability in responses to the MHW (as has 5 6 been shown for seaweed and fish, see Wernberg et al. 2013). In summary, our second hypothesis, which predicted lower post-MHW abundances of more southerly distributed cool-7 8 water species, was partially supported as cool-water species at Kalbarri were decimated by the 9 MHW and some, but not all, cool water species exhibited a response at Jurien.

10

11 Our third hypothesis – that the relative abundance of more northerly distributed 'warm-water' taxa would increase after the MHW – received some support as the abundance of *Tripneustes* 12 notably increased at some sites at Jurien for the years immediately following the MHW. 13 14 Tripneustes is a warm-water 'collector urchin' that is widely distributed across the Indo-Pacific and extended tropics, where it primarily feeds on drifting seagrass and macroalgae fragments 15 but can actively graze when per capita food supply is insufficient (Ogden et al. 1989, Valentine 16 & Edgar 2010). Tripneustes is notoriously 'boom and bust' and intense population outbreaks 17 have been observed in the temperate-tropical transition zone in eastern Australia, with 18 19 community-level consequences (Valentine & Edgar 2010). Tripneustes is common in subtropical waters off Western Australia and, although it has occasionally been recorded 20 21 further poleward than Jurien (e.g. at Rottnest Island, Vanderklift personal observation), it was 22 extremely rare at all study locations and was not recorded in any transects completed before 23 the MHW. We suggest that the higher abundances of Tripneustes at Jurien in 2013 and 2014 was a consequence of the MHW for the following reasons: (1) the enhanced poleward flow of 24 25 the LC during the MHW would have increased larval supply into cooler locations (Feng et al. 2013, Wernberg et al. 2013); (2) temperatures experienced during the MHW in the Jurien Bay 26

1 region (i.e. 26-27°C, see Wernberg et al. 2013) would have been favourable for the development of Tripneustes larvae (Sheppard Brennand et al. 2010); (3) Tripneustes larvae are 2 relatively long-lived and are generally released in the austral summer (Dworjanyn & Pirozzi 3 4 2008), which was co-incident with the MHW and the unusually early strengthening of the LC (Pearce & Feng, 2013); and (4) the cohort of sea urchins observed in 2013 were similar in size 5 6 (test diameters of ~10-12 cm, Smale pers. obs), which according to growth rate estimates (Bacolod & Dy 1986) would have coincided with a recruitment event during or soon after the 7 MHW. However, as no *Tripneustes* individuals were recorded in 2015 the proliferation of the 8 9 warmer-water species was short-lived, with no indication that the population at Jurien will persist. 10

11

12 The hypothesis received limited support as there was no evidence of a proliferation of warmaffinity macroinvertebrates at the northernmost location (Kalbarri), where reef habitats were 13 devoid of benthic macrofauna. On the other hand, the principal 'winner' of the ecological 14 15 disturbance was *Centrostephanus*, which increased in abundance by a factor of ~15 at Jurien to outnumber *Heliocidaris* to become the most abundant sea urchin, and significantly increased 16 in abundance at Marmion following the MHW. Centrostephanus can be described as a 'warm-17 temperate' species and its proliferation does therefore provide some support for the third 18 hypothesis. Although *Centrostephanus* exhibits a temperate distribution from South Australia 19 to mid-Western Australia, it is far more abundant towards the warm northern limit of its 20 distribution compared with the cooler southern parts of its range (Vanderklift & Kendrick 2004, 21 Wernberg et al. 2016). For example, an unpublished survey based on 90 habitat-scale transects 22 conducted across 18 similar reefs between South Australia (Adelaide) and Marmion in 2005/06 23 did not find a single Centrostephanus individual on these southern reefs (T. Wernberg 24 unpublished data). Moreover, its equatorward range edge extends further north than the other 25

sea urchin species (except *Tripneustes*), with the northernmost population recorded at ~25°S (Marsh 1994, GBIF record 137025088).

3

2

Very little is known about the biology of Centrostephanus tenuispinus, but its warm-temperate 4 congener on the east coast of Australia, Centrostephanus rodgersii, has been extensively 5 6 studied in recent years (see Byrne & Andrew 2013 for review). C. rodgersii has recently extended its range polewards into Tasmania, in response to increased temperature and a 7 8 strengthening of the East Australian Current, where it has overgrazed kelp forest habitat to 9 create widespread urchin barrens (Ling et al. 2008, Ling et al. 2009). C. rodgersii has longlived planktonic larvae that facilitates long distance dispersal (Byrne & Andrew 2013); it is 10 likely that *C. tenuispinus* also has the potential for widespread dispersal. Increasing abundance 11 12 of *C. tenuispinus* along the southwest coastline of Australia is congruent with the recent MHW 13 and the longer-term gradual warming trend in the region (Pearce & Feng 2007). Thermal 14 tolerance studies on C. rodgersii and Heliocidaris on the east coast of Australia would suggest 15 that their thermal windows for fertilisation, development, growth and survival do not differ markedly (Byrne et al. 2010, Wolfe et al. 2013, Pecorino et al. 2014). As such, it is currently 16 unclear whether recent warming has directly (through physiological stress) or indirectly 17 (through temperature-mediated competitive interactions) favoured C. tenuispinus over 18 Heliocidaris in southwest Australia, and focussed experimental work is needed. It is also not 19 20 known whether C. tenuispinus on the west coast has the potential to modify kelp forest habitat to the extent of its congener on the east coast of Australia. 21

22

In conclusion, the 2011 MHW has had profound effects on the marine biota along the southwest
coastline of Australia. Rapid changes in the abundance, distribution and condition of organisms
representing a wide range of taxonomic and trophic groups have been reported (Pearce et al.
2011, Moore et al. 2012, Smale & Wernberg 2012, Smale & Wernberg 2013, Wernberg et al.

1 2013, Caputi et al. 2016, Wernberg et al. 2016). This study has shown that the responses of mobile macroinvertebrates assemblages to the warming event varied dramatically with latitude, 2 with greatest impact at the northernmost study locations, even though the magnitude of the 3 4 warming anomaly was consistent along the latitudinal gradient. Recent modelling approaches based on species distributions and projected temperatures suggest that species' range 5 6 expansions will be more prevalent than range contractions under climate change scenarios (García Molinos et al. 2016). However, in this 'extreme' example of warming, we primarily 7 observed loss of cooler-water macroinvertebrate species and less evidence for an influx or 8 9 proliferation of warmer-water species over the timescale of observation. In contrast, observed responses of other taxonomic groups to MHWs, including seaweeds, fish and corals, have 10 included a proliferation of warm-water species (Wernberg et al. 2013, Wernberg et al. 2016). 11 12 Clearly, better understanding of variability between biogeographic regions and taxonomic groups is needed to improve predictions the effects of climate change on marine ecosystems. 13

Coastal marine ecosystems along the vast and varied coastline of temperate Australia have 14 15 responded to longer-term gradual ocean warming and concurrent stressors in complex and unpredictable ways (Wernberg et al. 2011). As short-term climatic variability is superimposed 16 onto longer-term gradual warming trends in southwest Australia as in many other regions 17 around the world, absolute temperatures may now reach unprecedented highs during extreme 18 climatic events (Trenberth 2012). This will increase the likelihood of ecological tipping points 19 being exceeded, triggering rapid phase-shifts in some regions and habitats (Wernberg et al. 20 2016). Only time will tell, but the 2011 MHW may well have triggered rapid 'tropicalization' 21 (see Vergés et al. 2014) along much of the coastline, by driving widespread loss of temperate 22 flora and fauna and creating opportunities for rapid colonisation by a warm-water biota. 23 Moreover, inter-specific variability in susceptibility and responses to warming trends and 24 events, as shown here, will cause a reshuffling of species and the emergence of novel 25 communities and ecosystems. 26

1	Acknowledgements: We thank Scott Bennett, Tiffany Simpson, Emily Gates and Thibaut de		
2	Bettignies for assistance in the field. This work was funded by the Australian Research Council		
3	through grants to T.W and by a Marie Curie International Incoming Fellowship (within the		
4	seventh European Community Framework Programme) awarded to D.A.S., who is currently		
5	funded by the Natural Environment Research Council of the UK (IRF: NE/K008439/1).		
6			
7	LITERATURE CITED		
8			
9	Anderson MJ (2001) A new method for non-parametric multivariate analysis of variance.		
10	Austral Ecol 26:32-46		
11	Anderson MJ, Gorley RN, Clarke KR (2008) Permanova+ for primer: guide to software and		
12	statistical methods. PRIMER-E, Plymouth, UK		
13	Ayvazian SG, Hyndes GA (1995) Surf-zone fish assemblages in south-western Australia: do		
14	adjacent nearshore habitats and the warm Leeuwin Current influence the characteristics		
15	of the fish fauna? Mar Biol 122:527-536		
16	Azzarello J, Smale DA, Langlois TJ, Håkansson E (2014) Linking habitat characteristics to		
17	abundance patterns of canopy-forming macroalgae and sea urchins in southwest		
18	Australia. Mar Biol Res 10:682-693		
19	Bacolod PT, Dy DT (1986) Growth, recruitment pattern and mortality rate of the sea urchin,		
20	Tripneustes gratilla Linnaeus, in a seaweed farm at Danahon Reef, Central Philippines		
21	Phillipine Sci 23:1-14		

1	Bennett S, Wernberg T, Connell SD, Hobday AJ, Johnson CR, Poloczanska ES (2015a) The
2	'Great Southern Reef': social, ecological and economic value of Australia's neglected
3	kelp forests. Mar Freshwater Res 67:47-56
4	Bennett S, Wernberg T, Harvey ES, Santana-Garcon J, Saunders BJ (2015b) Tropical
5	herbivores provide resilience to a climate-mediated phase shift on temperate reefs. Ecol
6	Lett 18:714-723
7	Bergmann N, Winters G, Rauch G, Eizaguirre C, Gu J, Nelle P, Fricke B, Reusch TBH (2010)
8	Population-specificity of heat stress gene induction in northern and southern eelgrass
9	Zostera marina populations under simulated global warming. Mol Ecol 19:2870-2883
10	Byrne M, Andrew N (2013) Centrostephanus rodgersii. In: Lawrence JM (ed) Sea urchins:
11	biology and ecology. Elsevier
12	Byrne M, Soars NA, Ho MA, Wong E, McElroy D, Selvakumaraswamy P, Dworjanyn SA,
13	Davis AR (2010) Fertilization in a suite of coastal marine invertebrates from SE
14	Australia is robust to near-future ocean warming and acidification. Mar Biol 157:2061-
15	2069
16	Caputi N, Fletcher WJ, Pearce A, Chubb CF (1996) Effect of the Leeuwin Current on the
17	recruitment of fish and invertebrates along the Western Australian coast. Mar
18	Freshwater Res 47:147-155
19	Caputi N, Kangas M, Denham A, Feng M, Pearce A, Hetzel Y, Chandrapavan A (2016)
20	Management adaptation of invertebrate fisheries to an extreme marine heat wave event
21	at a global warming hot spot. Ecol Evol 6:3583-3593
22	Clarke KR, Warwick RM (2001) Change in marine communities: an approach to statistical
23	analysis and interpretation. PRIMER-E Ltd., Plymouth, UK

1	Depczynski M, Gilmour JP, Ridgway T, Barnes H, Heyward AJ, Holmes TH, Moore JAY,
2	Radford BT, Thomson DP, Tinkler P, Wilson SK (2013) Bleaching, coral mortality and
3	subsequent survivorship on a West Australian fringing reef. Coral Reefs 32:233-238
4	Diaz-Almela E, Marba N, Duarte CM (2007) Consequences of Mediterranean warming events
5	in seagrass (Posidonia oceanica) flowering records. Glob Change Biol 13:224-235
6	Dworjanyn SA, Pirozzi I (2008) Induction of settlement in the sea urchin Tripneustes gratilla
7	by macroalgae, biofilms and conspecifics: A role for bacteria? Aquaculture 274:268-
8	274
9	Ettinger-Epstein P, Kingsford MJ (2008) Effects of the El Niño southern oscillation on Turbo
10	torquatus (Gastropoda) and their kelp habitat. Austral Ecol 33:594-606
11	Feng M, McPhaden MJ, Xie S, Hafner J (2013) La Niña forces unprecedented Leeuwin Current
12	warming in 2011. Sci Rep 3:1277
13	García Molinos J, Halpern BS, Schoeman DS, Brown CJ, Kiessling W, Moore PJ, Pandolfi
14	JM, Poloczanska ES, Richardson AJ, Burrows MT (2016) Climate velocity and the
15	future global redistribution of marine biodiversity. Nature Clim Change 6:83-88
16	Garrabou J, Coma R, Bensoussan N, Bally M, Chevaldonne P, Cigliano M, Diaz D, Harmelin
17	JG, Gambi MC, Kersting DK, Ledoux JB, Lejeusne C, Linares C, Marschal C, Perez T,
18	Ribes M, Romano JC, Serrano E, Teixido N, Torrents O, Zabala M, Zuberer F, Cerrano
19	C (2009) Mass mortality in Northwestern Mediterranean rocky benthic communities:
20	effects of the 2003 heat wave. Glob Change Biol 15:1090-1103
21	Hansen J, Sato M, Ruedy R (2012) Perception of climate change. Proc Nat Acad Sci USA
22	109:E2415-E2423

1	Hobday AJ, Alexander LV, Perkins SE, Smale DA, Straub SC, Oliver EC, Benthuysen J,
2	Burrows MT, Donat MG, Feng M, Holbrook NJ, Moore PJ, Scannell HA, Sen Gupta
3	A, Wernberg T (2016) A hierarchical approach to defining marine heatwaves. Prog
4	Oceanogr 141:227-238
5	IPCC (2012) Managing the Risks of Extreme Events and Disasters to Advance Climate Change
6	Adaptation. A Special Report of Working Groups I and II of the Intergovernmental
7	Panel on Climate Change. Cambridge University Press, Cambridge, UK, and New
8	York, NY, USA
9	Joll L (1980) Reproductive biology of two species of Turbinidae (Mollusca : Gastropoda). Mar
10	Freshwater Res 31:319-336
11	Keesing JK (2001) The ecology of Heliocidaris erythrogramma. In: John ML (ed)
12	Developments in Aquaculture and Fisheries Science, Book Volume 32. Elsevier
13	Lejeusne C, Chevaldonne P, Pergent-Martini C, Boudouresque CF, Perez T (2009) Climate
14	change effects on a miniature ocean: the highly diverse, highly impacted Mediterranean
15	Sea. Trends Ecol Evol 25:250-260
16	Levitus S, Antonov JI, Boyer TP, Baranova OK, Garcia HE, Locarnini RA, Mishonov AV,
17	Reagan JR, Seidov D, Yarosh ES, Zweng MM (2012) World ocean heat content and
18	thermosteric sea level change (0-2000 m), 1955-2010. Geophys Res Lett 39:L10603
19	Lima FP, Wethey DS (2012) Three decades of high-resolution coastal sea surface temperatures
20	reveal more than warming. Nature Commun 3:704
21	Ling SD, Johnson CR, Frusher S, King CK (2008) Reproductive potential of a marine
22	ecosystem engineer at the edge of a newly expanded range Glob Change Biol 14:907-
23	915

1	Ling SD, Johnson CR, Frusher SD, Ridgway KR (2009) Overfishing reduces resilience of kelp
2	beds to climate-driven catastrophic phase shift. Proc Nat Acad Sci USA 106:22341-
3	22345
4	Lozano-Montes HM, Loneragan NR, Babcock RC, Jackson K (2011) Using trophic flows and
5	ecosystem structure to model the effects of fishing in the Jurien Bay Marine Park,
6	temperate Western Australia. Mar Freshwater Res 62:421-431
7	MacArthur LD, Phillips DL, Hyndes GA, Hanson CE, Vanderklift MA (2011) Habitat
8	surrounding patch reefs influences the diet and nutrition of the western rock lobster.
9	Mar Ecol Prog Ser 436:191-205
10	Marba N, Duarte CM (2010) Mediterranean warming triggers seagrass (Posidonia oceanica)
11	shoot mortality. Glob Change Biol 16:2366-2375
12	Marsh LM (1994) Echinoderms of the Houtman Abrolhos Islands, Western Australia and their
13	relationship to the Leeuwin Current. In: David B, Guille A, Feral J-P (eds) Echinoderms
14	through time: Proceedings of the 8th international echinoderm conference, Dijon,
15	France, 6-10 September 1993 A. A. Balkema Rotterdam
16	Meehl G, Tebaldi C (2004) More intense, more frequent, and longer lasting heat waves in the
17	21st Century. Science 305:994-997
18	Moore JAY, Bellchambers LM, Depczynski MR, Evans RD, Evans SN, Field SN, Friedman
19	KJ, Gilmour JP, Holmes TH, Middlebrook R, Radford BT, Ridgway T, Shedrawi G,
20	Taylor H, Thomson DP, Wilson SK (2012) Unprecedented mass bleaching and loss of
21	coral across 12° of latitude in Western Australia in 2010–11. PLoS ONE 7:e51807
22	Normile D (2016) El Niño's warmth devastating reefs worldwide. Science 352:15-16

1	Ogden NB, Ogden JC, Abbott IA (1989) Distribution, abundance and food of sea urchins on a	
2	leeward Hawaiian reef. Bull Mar Sci 45:539-549	
3	Pearce A, Feng M (2007) Observations of warming on the Western Australian continental	
4	shelf. Mar Freshwater Res 58:914-920	
5	Pearce A, Lenanton R, Jackson G, Moore J, Feng M, Gaughan D (2011) The "marine heat	
6	wave" off Western Australia during the summer of 2010/11. Fisheries Research Report	
7	No. 222. Department of Fisheries, Perth, Australia	
8	Pearce AF (1991) Eastern boundary currents of the southern hemisphere. J Roy Soc West Aust	
9	74:35-45	
10	Pearce AF, Feng M (2013) The rise and fall of the "marine heat wave" off Western Australia	
11	during the summer of 2010/2011. J Mar Syst 111–112:139-156	
12	Pecorino D, Barker MF, Dworjanyn SA, Byrne M, Lamare MD (2014) Impacts of near future	
13	sea surface pH and temperature conditions on fertilisation and embryonic development	
14	in Centrostephanus rodgersii from northern New Zealand and northern New South	
15	Wales, Australia. Mar Biol 161:101-110	
16	Phillips B (1969) The population of the whelk Dicathais aegrota in Western Australia. Mar	
17	Freshwater Res 20:225-266	
18	Rahmstorf S, Coumou D (2011) Increase of extreme events in a warming world. Proc Nat Acad	
19	Sci USA 108:17905-17909	
20	Rose T, Smale DA, Botting GC (2012) The 2011 marine heat wave in Cockburn Sound,	
21	southwest Australia. Ocean Sci 8:545-550	

1	Searle DJ, Semeniuk V (1985) The natural sectors of the inner Rottnest Shelf coast adjoining	
2	the Swan Coastal Plain. J Roy Soc West Aust 67:116-136	
3	Shepherd S, Edgar G (2013) Ecology of Australian temperate reefs: the unique South. CSIRO	
4	publishing, Clayton, Victoria, Australia	
5	Sheppard Brennand H, Soars N, Dworjanyn SA, Davis AR, Byrne M (2010) Impact of ocean	
6	warming and ocean acidification on larval development and calcification in the sea	
7	urchin Tripneustes gratilla PLoS ONE 5:e11372	
8	Smale DA, Kendrick GA, Wernberg T (2010) Assemblage turnover and taxonomic sufficiency	
9	of subtidal macroalgae at multiple spatial scales. J Exp Mar Biol Ecol 384:76-86	
10	Smale DA, Wernberg T (2009) Satellite-derived SST data as a proxy for water temperature in	
11	nearshore benthic ecology. Mar Ecol Prog Ser 387:27-37	
12	Smale DA, Wernberg T (2012) Ecological observations associated with an anomalous warming	
13	event at the Houtman Abrolhos Islands, Western Australia. Coral Reefs 31:441	
14	Smale DA, Wernberg T (2013) Extreme climatic event drives range contraction of a habitat-	
15	forming species. Proc Roy Soc B Biol Sci 280:20122829	
16	Smale DA, Wernberg T (2014) Population structure of the purple sea urchin Heliocidaris	
17	erythrogramma along a latitudinal gradient in southwest Australia. J Mar Biol Assoc	
18	UK 94:1033-1040	
19	Smale DA, Wernberg T, Vance T (2011) Community development on subtidal temperate reefs:	
20	the influences of wave energy and the stochastic recruitment of a dominant kelp. Mar	
21	Biol 158:1757–1766	

1	Smith RL, Huyer A, Godfrey JS, Church JA (1991) The Leeuwin Current off Western
2	Australia. J Phys Oceanogr 21:323-345
3	Stoner AW (2009) Habitat-mediated survival of newly settled red king crab in the presence of
4	a predatory fish: Role of habitat complexity and heterogeneity. J Exp Mar Biol Ecol
5	382:54-60
6	Thibault KM, Brown JH (2008) Impact of an extreme climatic event on community assembly
7	Proc Nat Acad Sci USA 105:3410-3415
8	Tittensor DP, Mora C, Jetz W, Lotze HK, Ricard D, Berghe EV, Worm B (2010) Global
9	patterns and predictors of marine biodiversity across taxa. Nature 466:1098-1101
10	Trenberth KE (2012) Framing the way to relate climate extremes to climate change. Climatic
11	Change 115:283-290
12	Valentine JP, Edgar GJ (2010) Impacts of a population outbreak of the urchin Tripneustes
13	gratilla amongst Lord Howe Island coral communities. Coral Reefs 29:399-410
14	Van De Pol M, Ens BJ, Heg D, Brouwer L, Krol J, Maier M, Exo K-M, Oosterbeek K, Lok T,
15	Eising CM, Koffijberg K (2010) Do changes in the frequency, magnitude and timing of
16	extreme climatic events threaten the population viability of coastal birds? J Appl Ecol
17	47:720-730
18	Vanderklift MA, Kendrick GA (2004) Variations in abundances of herbivorous invertebrates
19	in temperate subtidal rocky reef habitats. Mar Freshwater Res 55:93-103
20	Vanderklift MA, Kendrick GA, Smit AJ (2006) Differences in trophic position among
21	sympatric sea urchin species. Estuarine, Coastal and Shelf Science 66:291-297

1	Vanderklift MA, Wernberg T (2010) Stable isotopes reveal a consistent consumer-diet	
2	relationship across hundreds of kilometers. Mar Ecol Prog Ser 403:53-61	
3	Vergés A, Steinberg PD, Hay ME, Poore AGB, Campbell AH, Ballesteros E, Heck KL, Booth	
4	DJ, Coleman MA, Feary DA, Figueira W, Langlois T, Marzinelli EM, Mizerek T,	
5	Mumby PJ, Nakamura Y, Roughan M, van Sebille E, Gupta AS, Smale DA, Tomas F,	
6	Wernberg T, Wilson SK (2014) The tropicalization of temperate marine ecosystems:	
7	climate-mediated changes in herbivory and community phase shifts. Proc Roy Soc B	
8	281:20140846	
9	Wernberg T, Bennett S, Babcock RC, de Bettignies T, Cure K, Depczynski M, Dufois F,	
10	Fromont J, Fulton CJ, Hovey RK, Harvey ES, Holmes TH, Kendrick GA, Radford B,	
11	Santana-Garcon J, Saunders BJ, Smale DA, Thomsen MS, Tuckett CA, Tuya F,	
12	Vanderklift MA, Wilson S (2016) Climate-driven regime shift of a temperate marin	
13	ecosystem. Science 353:169-172	
14	Wernberg T, Russell BD, Moore PJ, Ling SD, Smale DA, Coleman M, Steinberg PD, Kendrick	
15	GA, Connell SD (2011) Impacts of climate change in a global hotspot for temperate	
16	marine biodiversity and ocean warming. J Exp Mar Biol Ecol 400:7-16	
17	Wernberg T, Smale DA, Tuya F, Thomsen MS, Langlois TJ, de Bettignies T, Bennett S,	
18	Rousseaux CS (2013) An extreme climatic event alters marine ecosystem structure in a	
19	global biodiversity hotspot. Nature Clim Change 3:78-82	
20	Wernberg T, Thomsen MS, Tuya F, Kendrick GA, Staehr PA, Toohey BD (2010) Decreasing	
21	resilience of kelp beds along a latitudinal temperature gradient: potential implications	

for a warmer future. Ecol Lett 13:685-694

1	Wernberg T, White M, Vanderklift MA (2008) Population structure of turbinid gastropods on			
2	wave-exposed subtidal reefs: effects of density, body size and algae on grazing			
3	behaviour. Mar Ecol Prog Ser 362:169-179			
4	Williams ST (2007) Origins and diversification of Indo-West Pacific marine fauna:			
5	evolutionary history and biogeography of turban shells (Gastropoda, Turbinidae). Bio			
6	J Linnean Soc 92:573-592			
7	Wolfe K, Dworjanyn SA, Byrne M (2013) Effects of ocean warming and acidification on			
8	survival, growth and skeletal development in the early benthic juvenile sea urchin			
9	(Heliocidaris erythrogramma). Glob Change Biol 19:2698-2707			
10				
11				

- 1 Table 1. Most common macroinvertebrate species recorded during surveys and their
- 2 affinities and broad-scale distributions within the southwest Australian study region. The
- 3 proportion of distribution records found either north or south of Jurien Bay $(30.3^{\circ}S)$ is
- 4 provided as an indication of the biogeographical affinities of each species (records were
- 5 downloaded from The Atlas of Living Australia <u>http://www.ala.org.au</u> on 30/01/2017).
- 6 Additional general information sourced from Edgar (1997). [#]Indicates few existing records
- 7 for that species and as such distributions should be treated with caution, but wider affinity
- 8 determined from best available knowledge.
- 9

Species	Taxa	Affinity and distribution
Centrostephanus	Sea	Warm; warm-temperate, northern range-limit at Shark Bay (25.3 °S).
tenuispinus	urchin	[#] 28.3% of 11 records north of 30.3°S. Family characteristic of tropical
		reefs
Tripneustes	Sea	Warm; tropical, northern Australia, 68.8% of 288 records north of
gratilla	urchin	30.3°S
Phyllacanthus	Sea	Cool; temperate, #70% of 20 records south of 30.3°S
irregularis	urchin	
Heliocidaris	Sea	Cool; temperate, 92.3% of 607 records south of 30.3°S.
erythrogramma	urchin	
Lunella	Gastropod	Cool; temperate, 87.4% of 372 records south of 30.3°S
torquatus	mollusc	
Astralium spp.	Gastropod	Cool; temperate, 94.9% of 431 records south of 30.3°S
(A. tentorium/ A.	mollusc	
squamiferum)		
Dicathais orbita	Gastropod	Cool; temperate, 90.9% of 2242 records south of 30.3°S
	mollusc	
Petricia	Sea star	Cool; temperate, 97.0% of 755 records south of 30.3°S
vernicina		

1 Figure legends

2

Fig 1. Map of southwest Australia indicating the Kalbarri (K), Jurien Bay (J), Marmion Lagoon
(M) and Hamelin Bay (H) study locations. The region is characterized by a well-defined
oceanic temperature gradient, represented here by average winter isotherms (in °C, 2005–07).
Temperature anomalies represent deviations from monthly long-term means (1981-2010),
derived from remotely-sensed satellite SSTs.

Fig. 2. PCO plots showing macroinvertebrate assemblage structure on reef flats before and after the MHW at each location. Ordinations are based on Bray-Curtis similarities of squareroot transformed abundance data. Dashed circle in Kalbarri plot indicates 16 overlapping samples (15 samples post-MHW and 1 sample pre-MHW) in which macroinvertebrates were absent.

Fig. 3. PCO plots showing macroinvertebrate assemblage structure on reef slopes before and
after the MHW at each location. Ordinations are based on Bray-Curtis similarities of squareroot transformed abundance data.

Fig. 4. Mean total abundance (left-hand plots) and taxon richness (right-hand plots) of all mobile macroinvertebrates on reef flats and slopes at each location before (blue bars) and after (red bars) the MHW (\pm SE). Values represent number of individuals/species per 25 m² sample area (i.e. 5 transects pooled per site). Significant differences before and after the MHW are indicated with an asterisk (at P>0.05, test results shown in Tables S6 and S7).

Fig. 5. Mean abundances (± SE) of dominant species on reef flats before (blue bars) and after
(red bars) the MHW. Significant differences before and after the MHW are indicated with an
asterisk (at P>0.05, test results shown in Table S8).

- 1 Fig. 6. Mean abundances (\pm SE) of dominant species on reef slopes before (blue bars) and after
- 2 (red bars) the MHW. Significant differences before and after the MHW are indicated with an
- 3 asterisk (at P>0.05, test results shown in Table S9)
- 4 Fig. 7. The relative abundances of sea urchins recorded on reef slopes during each survey year
- 5 at each location (average of 3 sites per location). The timing of the MHW is also shown.



- 5 Fig. 1









4 Fig. 3



4 Fig. 4







4 Fig. 5



- 4 Fig. 6

