FEATURE ARTICLE

Zooplanktivory is a key process for fish production on a coastal artificial reef

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ABSTRACT: Artificial reefs continue to be deployed in coastal areas to enhance local fisheries. An important factor influencing the success of artificial reefs may be the provision of refuge for zooplanktivorous fishes, which use artificial reefs as a base to forage the surrounding zooplankton. A numerical model was developed to quantify this trophic pathway on a designed coastal artificial reef, using field-parameterised data for zooplankton biomass, current velocity, and the consumption rate and abundance of a reef-resident zooplanktivorous fish (Atypichthys strigatus). The model estimated that this species consumed ~2.9 kg (1.0 g m⁻³) of zooplankton per day on this artificial reef, which represents only 0.35% of the total zooplankton biomass. The ability of this artificial reef to support ~130 kg standing stock of this species suggests that the zooplankton pathway is a reliable mechanism for fish production. A second model explored the influence of reef size on zooplanktivorous fish densities and the supply of zooplankton required to sustain their consumption rate. As reef size increased, the ratio between the foraging volume and refuge volume declined, meaning that small reefs have lots of food and not much refuge, and large reefs can have lots of refuge but not enough food. This indicates that reef size can be manipulated to maximise fish abundance while avoiding food limitation. Reef size, shape, and orientation should be considered carefully during the planning of artificial reefs, as it can greatly influence the foraging of reef-resident zooplanktivorous fishes and thus influence the entire reef assemblage.

KEY WORDS: Artificial reefs · Zooplankton · Zooplanktivorous fish · Foraging halo · Atypichthys strigatus · Trophic ecology

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INTRODUCTION

Artificial reefs are often deployed in coastal marine areas to create fishing opportunities, promote biodiversity, and restore degraded habitats (Miller 2002, Claudet & Pelletier 2004, Seaman 2007). This is especially true of designed artificial reefs, which, unlike materials of opportunity (Seaman et al. 1989), are built and deployed to achieve specific environmental objectives (Pickering & Whitmarsh 1997, Baine 2001). Artificial reefs are popular fisheries enhancement tools because fish biomass often increases in surrounding waters after their deployment (Bombace et al. 1994, Charbonnel et al. 2002, Leitão 2013). Whether increases in fish biomass involve new production or simply attraction of existing biomass is debated (Bohnsack 1989, Grossman et al. 1997, Pow-
ers et al. 2003, Brickhill et al. 2005). It is generally agreed, however, that artificial reefs can provide habitat space and food resources for fishes (Peterson et al. 2003, Powers et al. 2003, Cresson et al. 2014), and it is this provision of refuge and food that is most likely to drive any production of fish biomass on these reefs (Charbonnel et al. 2002, Powers et al. 2003). Thus, quantitative research is needed to explore how artificial reefs influence fish production via these factors.

The capacity for artificial reefs to increase habitat space and the availability of food is exemplified by the trophic link between zooplankton and reef-resident zooplanktivorous fishes. Zooplanktivores can be a dominant feeding guild in a variety of ecological contexts (Hobson & Chess 1976, 1978, Ebeling et al. 1980, Edgar et al. 2014) and can be extremely abundant on coastal artificial reefs (Scott et al. 2015). Their capacity to continuously access zooplankton supplied by prevailing coastal currents indicates that increasing the provision of reef habitat may allow for increased production of zooplanktivorous fishes. Zooplanktivores are readily preyed upon by piscivorous species (Young et al. 2010) and thus require the refuge provided by artificial reefs to forage the surrounding zooplankton. This suggests that the productivity of artificial reefs, and their subsequent contribution to local fisheries, may be largely dependent on the direct and underappreciated trophic link between zooplankton and reef-resident zooplanktivorous fishes.

There is evidence to suggest that a balance of predation risk and foraging success influences the association of fish with reefs (Frazer & Lindberg 1994, Biesinger et al. 2011, 2013), specifically the maximum distance from refuge habitat that prey fish will forage (Biesinger et al. 2011). The distance that reef-resident zooplanktivorous fish will forage from refuge determines the total volume of water surrounding a reef that is available to be foraged, and it is likely that this distance is largely independent of reef size (Scott et al. 2015). Thus, reef size (i.e. refuge habitat size) would not scale linearly with the surrounding foraging volume, which has interesting implications for food availability and reef-resident zooplanktivore density. The clear spatial boundaries of artificial reefs make them ideal sites to investigate this influence of reef size on the dynamics of reef-resident zooplanktivorous fishes.

The goal of this study was to explore the contribution of zooplanktivory to the production of fish biomass on artificial reefs and the management implications of this ecological process. Specifically, this study aimed to (1) describe the diet and habitat use of an abundant zooplanktivorous fish on a designed coastal artificial reef, (2) estimate the depletion of zooplankton due to predation by zooplanktivorous fish on this reef, and (3) illustrate the influence of artificial reef size on foraging volume and food availability for resident zooplanktivores.

**MATERIALS AND METHODS**

**Location**

This study was conducted at a coastal artificial reef that was deployed in October 2011, located 1.2 km off the coast of Sydney, NSW, Australia (33°50.797’ S, 151°17.988’ E; Fig. 1). The reef was designed to increase fishing opportunities for recreational anglers, is 15 by 12 m and 12 m tall, including 8 m high spires (Fig. 2). It is made of steel, and is located at a bottom depth of 38 m (Scott et al. 2015). The nearest natural reef of any significance is located ~600 m inshore from the artificial reef. Nine sampling trips to collect fish and zooplankton data were made from February to August 2014.

**Dietary analysis of *Atypichthys strigatus***

The fish species chosen for this study was mado *A. strigatus* (Günther). They are small reef-resident zooplanktivores common to the temperate reefs of southeastern Australia (Glasby & Kingsford 1994, Fowler & Booth 2013) and are one of the most abundant fishes found on this study’s artificial reef (NSW Department of Primary Industries 2013, Scott et al. 2015). Fifty-five *A. strigatus* were sampled from the reef using hook-and-line fishing, frozen on collection, and later

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**Fig 1. Location in Australia of the coastal artificial reef used in this study (circle)**
dissected with their stomachs stored in 5% formaldehyde. The diet of *A. strigatus* was determined by gravimetric analyses of stomach contents (Hyslop 1980). Total prey biomass and diet composition by mass were quantified for each individual. Stomach contents were sorted into broad taxonomic groups, and groups were dried separately at 60°C (Man & Hodgkiss 1977) to determine their respective dry weights (Berg 1979). The dry weight ratio of prey groups to total stomach contents was used to determine the corresponding wet biomass of individual prey groups, as wet and dry weights are highly correlated (Glenn & Ward 1968). Wet weight values were subsequently used to calculate the proportion of the *A. strigatus* diet comprised of zooplankton.

**Density and foraging volume of *A. strigatus***

It was necessary to estimate the density of *A. strigatus* at the reef in order to estimate the total consumption of zooplankton, and this was conducted using underwater video surveys. Video surveys were done using cameras lowered to the reef from a boat (‘drop cameras’) during periods when the reef was not being fished. Five replicate drop cameras, each of 10 min duration, were done on separate days to sample the abundance of *A. strigatus*. Drop cameras could not survey inside the reef, so the density of *A. strigatus* within the reef was estimated using a remotely operated vehicle (ROV; Seamor Marine). Two ROV surveys of the reef were done, each for ~30 min in duration. The average abundance of *A. strigatus* was estimated from 80 still frames, randomly selected from all suitable drop camera and ROV footage. Abundance was converted to density by estimating the volume of water in each still frame, using camera field-of-view and the day-specific viewing distance (i.e. water visibility). The drop camera field-of-view (in both vertical and horizontal planes) was calculated in an ocean swimming pool by filming a known area of a vertical surface from a known distance, while the ROV field-of-view was estimated using the ROV’s laser scaler. Viewing distance was calculated on each day by lowering a camera fixed above an array of black and white rods at known distances from the camera (J. A. Smith, W. K. Cornwell, M. B. Lowry, I. M. Suthers unpubl.). To refine the spatial distribution of *A. strigatus* around the reef, fish density was partitioned into 0.5 m distance bins (e.g. 0.5–1 m from the reef). This was done by estimating the distance of individual *A. strigatus* from the reef, achieved by relating their locations in each still frame to known distances between the reef’s structural features (e.g. vertical and horizontal beams). Likewise, the total volume of water in each snapshot was partitioned into distance bins to estimate bin-specific fish densities.

The reef volume and the surrounding foraging volume had to be determined to calculate total *A. strigatus* abundance from the above density estimates. Engineering diagrams were used to calculate the volume of water within the reef (hereafter ‘reef volume’), and the surrounding volume as distance from the reef increased (hereafter ‘foraging volume’; Fig. 2). Total reef volume and the volume of water held within 0.5 m foraging bins extending from the structure were combined with bin-specific density estimates to calculate *A. strigatus* abundance within each bin.

**Consumption by *A. strigatus***

Food consumption as a function of biomass \((Q/B \text{ yr}^{-1})\) was estimated using an empirical formula derived by Palomares & Pauly (1998), which is a common method to predict consumption rates of marine fishes (e.g. Hughes et al. 2014),

\[
\log_{10} \frac{Q}{B} = 7.964 - 0.204 \log_{10} W_a - 1.965 T + 0.083 AR + 0.532m + 0.398d
\]
where $W_\infty$ is the asymptotic weight of an individual (g), $T'$ is the mean water temperature (equal to $1000/\left(\text{T}^\circ\text{C} + 273.1\right)$), $AR$ is the aspect ratio of the caudal fin, and $h$ and $d$ are dummy variables relating to feeding preference; i.e. carnivore ($m = 0, d = 0$), detritivore ($m = 0, d = 1$), or herbivore ($m = 1, d = 0$). Both $m$ and $d$ were set to 0 due to $A$. strigatus being predominantly zooplanktivorous (Glasby & Kingsford 1994). Caudal fin aspect ratio is a species-specific morphometric variable described by:

$$AR = \frac{h^2}{s}$$ (2)

where $h$ and $s$ refer to the height (cm) and lateral surface area of the caudal fin (to the caudal peduncle; cm$^2$), respectively. $AR$ is a proxy of the metabolic functioning of fishes and generally increases as fish activity increases (Palomares & Pauly 1989). An average $AR$ was calculated from 53 $A$. strigatus across a broad size range. $W_\infty$ was calculated by converting asymptotic length ($L_\infty$; cm) to weight (g) using a standard length–weight relationship ($a = 0.03; b = 2.90$); $L_\infty$ was taken to be 25 cm (total length, TL) as reported by the Australian Museum. Water temperature was measured on each sampling day using a SBE 19-plus V2 SeaCAT Profiler CTD (Sea-Bird Electronics Inc.) at depths relevant to the reef (24–38 m).

It is known that the visual acuity of zooplanktivorous fishes affects patterns of prey size-selection by limiting their detection of small items (O’Brien 1979, Wankowski 1979, Li et al. 1985). Thus, an analysis of prey sizes in stomach contents of $A$. strigatus was done to quantify their pattern of prey size-selection.

The supply of zooplankton was measured using plankton tows between 50 and 200 m up-current from A. strigatus (Bai et al. 2014). Caudal velocity was measured on each sampling day using a mechanical flow meter attached to the reef, which measured average velocity at 10 min intervals from June–August 2013 and from December 2013–February 2014.

**Zooplankton supply and availability**

The supply of zooplankton was measured using plankton tows between 50 and 200 m up-current from the artificial reef. This was done up-current to ensure sampled zooplankton had not been exposed to consumption by this reef’s residents. Given the depth and comparatively small size of this reef, it was impractical to attempt a corresponding downstream sample of zooplankton as in Hamner et al. (1988), so the depletion of zooplankton could only be modelled. A 40 cm diameter, 100 μm mesh plankton net was towed horizontally from a boat for 4 min per tow, 15–20 m from the surface. The distance of each tow was calculated using a GPS, and each tow sampled ~25 m$^3$ of seawater. Three replicate tows were done per sampling day ($n = 27$). Plankton samples were preserved with 5% formaldehyde immediately after collection. Samples were inspected under a dissecting microscope to ensure they contained only zooplankton and were then sorted using a laser optical plankton counter (LOPC; Herman et al. 2004). The LOPC sorts particles into size classes and estimates total count and biomass (mg m$^{-3}$) per size class (Suthers et al. 2004).

**Model 1: Zooplankton depletion**

A mathematical model was developed to estimate the depletion of zooplankton caused by $A$. strigatus predation at the coastal artificial reef in this study. This model predicted the depletion of each size class of zooplankton by incorporating zooplankton biomass, current velocity, and the consumption rate, density, and foraging volumes of $A$. strigatus. A Monte Carlo simulation was used to include parameter variation in the model, and this is detailed in ‘Monte Carlo simulation and sensitivity analysis’. See Table 1 for parameter descriptions and values.

Zooplankton depletion by $A$. strigatus at this artificial reef was calculated as:

$$Dep_i = \frac{C_i}{Bb_i}$$ (3)

where $Dep_i$ is the proportional depletion of zooplankton biomass from size class $i$ ($i = 9$), and $Bb_i$ (mg m$^{-3}$) is the total biomass of zooplankton within size class $i$ before predation by $A$. strigatus. $Bb_i$ was also replaced by $Ba_i$ (Eq. S2 in the Supplement) to calculate the depletion of only the zooplankton bio-
Table 1. All model parameters (Sens.: * = parameters included in a sensitivity model), their mean values, and the distributions and associated variance used in the Monte Carlo analysis. Normal (N), lognormal (ln N), and beta sampling distributions (Beta) were used, and numbers within brackets represent each distribution’s sampling parameters, as defined using R (R Core Team 2015) notation. na: parameters for which uncertainty was not necessary or not logical. Figs. S2 & S3 and Eq. (S1) are in the Supplement.

<table>
<thead>
<tr>
<th>Name</th>
<th>Sens.</th>
<th>Description</th>
<th>Mean distribution</th>
<th>Sampling number</th>
<th>Equation</th>
</tr>
</thead>
<tbody>
<tr>
<td>AR</td>
<td></td>
<td>Caudal fin aspect ratio</td>
<td>2.29</td>
<td>N (2.29, 0.47)</td>
<td>1, 3, 13</td>
</tr>
<tr>
<td>$B_l$</td>
<td>*</td>
<td>Amount of $B_{bun}$ that is available to be foraged by $A. strigatus$ at the reef based on observed prey size-selection</td>
<td>149.0 mg m$^{-3}$</td>
<td>na</td>
<td>16</td>
</tr>
<tr>
<td>$B_{ai}$</td>
<td></td>
<td>Biomass zooplankton within size class i before predation at the reef that is available to $A. strigatus$ given the observed pattern of prey size-selection</td>
<td>See Fig. S3B</td>
<td>$B_b \times Sel_i$ (see Eq. S1)</td>
<td>3</td>
</tr>
<tr>
<td>$B_{bi}$</td>
<td></td>
<td>Biomass of zooplankton within size class i before predation at the reef</td>
<td>See Fig. S3B</td>
<td>In $N(\mu, \sigma)$ (see Supplement)</td>
<td>3</td>
</tr>
<tr>
<td>$B_{b_{min}}$</td>
<td></td>
<td>Minimum environmentally available biomass of zooplankton before predation at the reef</td>
<td>197.4 mg m$^{-3}$</td>
<td>na</td>
<td>16</td>
</tr>
<tr>
<td>$D_{max}$</td>
<td>*</td>
<td>Maximum distance that the zooplanktivore will forage from shelter under the risk of predation</td>
<td>3.36 m</td>
<td>$N(3.36, 0.48)^a$</td>
<td>11</td>
</tr>
<tr>
<td>$D_{maxo}$</td>
<td></td>
<td>Maximum observed distance of $A. strigatus$ from the reef</td>
<td>4 m</td>
<td>na</td>
<td>7</td>
</tr>
<tr>
<td>$D_v$</td>
<td></td>
<td>Mean viewing distance (i.e. water visibility)</td>
<td>10 m</td>
<td>$N(10,1.5)$</td>
<td>12</td>
</tr>
<tr>
<td>$D_{m}^{maxo}$</td>
<td>*</td>
<td>Maximum within reef density of $A. strigatus$ observed at the reef</td>
<td>5.72 $A. strigatus$ m$^{-3}$</td>
<td>na</td>
<td>8</td>
</tr>
<tr>
<td>$D_{m}^{p}$</td>
<td>*</td>
<td>Density of $A. strigatus$ within foraging bin p at the reef</td>
<td>See ‘Results’; Fig. 4</td>
<td>Sampled from raw data, see Fig. 4</td>
<td>5</td>
</tr>
<tr>
<td>$H$</td>
<td>*</td>
<td>Reef height</td>
<td>4 m</td>
<td>na</td>
<td>9, 11</td>
</tr>
<tr>
<td>$L_o$</td>
<td></td>
<td>Observed maximum length of the reef</td>
<td>15 m</td>
<td>na</td>
<td>7</td>
</tr>
<tr>
<td>$P_{con}$</td>
<td>*</td>
<td>Proportion of the total zooplankton supply allocated to the target zooplanktivore for consumption</td>
<td>0.25</td>
<td>Beta (18.5, 55.5)</td>
<td>16</td>
</tr>
<tr>
<td>$P_i$</td>
<td></td>
<td>Proportion of zooplankton size class i observed in the stomachs of $A. strigatus$</td>
<td>See Fig. S3A</td>
<td>na</td>
<td>6</td>
</tr>
<tr>
<td>$P_{zoo}$</td>
<td>*</td>
<td>Observed mean proportion of the $A. strigatus$ diet made up of zooplankton</td>
<td>0.93</td>
<td>Beta (23.3, 1.75)</td>
<td>6, 19</td>
</tr>
<tr>
<td>$Q/B$</td>
<td>*</td>
<td>Consumption rate of $A. strigatus$ as a function of biomass per year</td>
<td>8.88</td>
<td>$N(8.88, 0.91)^a$</td>
<td>1, 6, 19</td>
</tr>
<tr>
<td>$S_{speed}$</td>
<td></td>
<td>Swim speed of a common predator of zooplanktivorous fishes (herein Seriola lalandi)</td>
<td>5.53 m s$^{-1}$</td>
<td>$N(5.53, 0.6)$</td>
<td>12</td>
</tr>
<tr>
<td>$S_{pey}$</td>
<td></td>
<td>Swim speed of a reef-resident zooplanktivore (herein $A. strigatus$)</td>
<td>1.29 m s$^{-1}$</td>
<td>$N(1.29, 0.17)$</td>
<td>12</td>
</tr>
<tr>
<td>$T$</td>
<td></td>
<td>Mean water temperature observed at the reef</td>
<td>19.83°C</td>
<td>$N(19.83, 1)$</td>
<td>1</td>
</tr>
<tr>
<td>$V_p$</td>
<td></td>
<td>Volume of foraging bin p at the reef</td>
<td>See Fig. S2</td>
<td>na</td>
<td>5</td>
</tr>
<tr>
<td>$Vel_{min}$</td>
<td>*</td>
<td>Minimum current velocity observed at the reef</td>
<td>0.042 m s$^{-1}$</td>
<td>na</td>
<td>18</td>
</tr>
<tr>
<td>$Vel_{max}$</td>
<td>*</td>
<td>Mean current velocity observed at the reef</td>
<td>0.091 m s$^{-1}$</td>
<td>In $N(\ln(0.091), 0.4)$</td>
<td>7</td>
</tr>
<tr>
<td>$W_t$</td>
<td></td>
<td>Total observed volume occupied by $A. strigatus$ at the reef</td>
<td>2879 m$^3$</td>
<td>na</td>
<td>4</td>
</tr>
<tr>
<td>$W$</td>
<td>*</td>
<td>Mean individual weight of $A. strigatus$ at the reef</td>
<td>33900 mg</td>
<td>$N(33900, 1600)$</td>
<td>6, 19</td>
</tr>
<tr>
<td>$W_a$</td>
<td></td>
<td>Asymptotic individual weight of $A. strigatus$</td>
<td>283 g</td>
<td>na</td>
<td>1</td>
</tr>
</tbody>
</table>

*aDistributions calculated from component models rather than raw data

\[ C_i = \frac{\sum_{p=1}^{9} (C_{in_i} \times N_p)}{V_{t_o}} \]  

where $V_{t_o}$ was the total volume (m$^3$) occupied by $A. strigatus$ at the reef (within-reef volume plus surrounding foraging volume), calculated using the maximum observed distance of $A. strigatus$ from the reef ($D_{maxo}$) during video surveys. $C_{in_i}$ (mg $A. strigatus^{-1}$ period$^{-1}$) is the consumption of zooplankton size that is within the size range available to $A. strigatus$ predation. $C_i$ is the consumption of zooplankton size class i by $A. strigatus$ in the duration that the zooplankton supply is exposed to predation. This duration was determined by current velocity and is termed a 'period' (Eq. 7). Thus, $C_i$ has the unit mg m$^{-3}$ period$^{-1}$ and is the sum of the consumption in the nine 0.5 m-wide foraging volume bins (p), calculated as:
class \( i \) by an individual \( A. \text{strigatus} \) per period, and \( N_p \) is the total abundance of \( A. \text{strigatus} \) within foraging volume bin \( p \), calculated as:

\[
N_p = \text{Den}_p \times V_p
\]

where \( \text{Den}_p \) is the density of \( A. \text{strigatus} \) within foraging bin \( p \) \((\text{A. strigatus m}^{-3})\), and \( V_p \) is the volume of foraging bin \( p \) \((\text{m}^3)\). \( C_{in} \) was calculated:

\[
C_{in} = \frac{Q \cdot B}{365 \times W \times P_{zoopl} \times P_i \times Per}
\]

where \( W \) (mg) is the mean individual weight of \( A. \text{strigatus} \) at the reef \((n = 55)\), \( P_{zoopl} \) is the observed mean proportion of the \( A. \text{strigatus} \) diet made up of zooplankton, \( P_i \) is the proportion of zooplankton size class \( i \) observed in the stomachs of \( A. \text{strigatus} \) \((n = 55)\), and \( Per \) is the mean time \((\text{s})\) taken for water to move completely through the foraging volume of \( A. \text{strigatus} \) (a ‘period’), calculated as:

\[
Per = \frac{(L_o + 2 \times D_{\text{max}})}{Vl_o}
\]

where \( L_o \) is the observed maximum length of the reef \((\text{m})\); \( D_{\text{max}} \) is the maximum distance that \( A. \text{strigatus} \) were observed from the reef \((\text{m})\); and \( Vel_o \) is the mean current velocity observed at the reef \((\text{m s}^{-1})\).

**Model 2: Reef size**

A general model was developed to identify how the size of an artificial reef influences the availability of food relative to the availability of habitat for reef-resident zooplanktivorous fish such as \( A. \text{strigatus} \) (Fig. 3). Specifically, this model was used to identify how artificial reef size influences the ratio of the total zooplankton supply (food supplied) to the required consumption of a reef’s maximum zooplanktivore population (food required). This model focused on the relationship between reef size and foraging volume, which is non-linear because the maximum foraging distance \( D_{\text{max}} \) for reef residents using the reef as a refuge is a constant that is generally independent of reef size (Biesinger et al. 2011). This is demonstrated in Fig. 3, in which per-capita food availability is highest when reef volume is small and foraging volume relative to reef volume (the \( Vf: Vr \) ratio) is large (Fig. 3A). Refuge is most abundant when reef volume is large, but foraging volume relative to reef volume is small and food limitation will, at some point, limit the density of fish (Fig. 3B). Model 2 was also designed to explore management implications associated with artificial reef design, given that reef size is relative to reef construction cost.

This general model can apply to any reef-resident zooplanktivorous fishes, i.e. those that use the reef structure for refuge and forage the surrounding pelagic environment, although the model depends on species-specific traits such as average body size and swim speed. The zooplankton biomass and current velocity input values were selected to reflect the harshest ecological conditions observed during the survey period (i.e. lowest density of available zooplankton and slowest observed current velocity) in order to conservatively estimate output values. To create the most general model possible, it was assumed that the internal reef volume was not part of the foraging volume (whereas fish could forage for zooplankton inside the reef in Model 1). This is true for many modern concrete artificial reefs, which, unlike the artificial reef in this study, are not necessarily designed to encourage water flow through the structure. Like Model 1, this model used a Monte Carlo simulation to include parameter variation in the model, and this is detailed in the following section. See Table 1 for parameter descriptions and values.

The maximum zooplanktivore density \( (\text{Den}_L; \text{fish m}^{-3}) \) that a specific reef of length \( L \) \((\text{length and width are assumed to be equal})\) is capable of supporting in its foraging volume \( (Vf_L; \text{m}^3) \) was calculated as:

\[
\text{Den}_L = \frac{N_L}{Vf_L}
\]

where \( N_L \) is the abundance of zooplanktivores at the artificial reef of length \( L \). \( N_L \) was calculated by assuming that all zooplanktivores must be able to...
simultaneously seek refuge within the reef (volume $V_{rL}$), and that the within-reef fish density is equal to the maximum within-reef fish density observed at this study’s artificial reef ($D_{enn,\text{max}}$). Thus, all artificial reefs in Model 2 have an equal maximum within-reef density, independent of reef size.

$V_{rL}$ ($m^3$) is the within volume of a reef of length $L$ ($m$), calculated as:

$$V_{rL} = L^2 \times H$$

where the simulated reef’s length ($L$) and width are considered equal, and $H$ is reef height ($m$). Reef height was equal to 4 m in this model and held constant because artificial reef units are often fixed in height. The height used is arbitrary and can be customised for specific reefs, but the general shapes of the relationships from Model 2 are independent of reef height.

$V_{fL}$ is the foraging volume ($m^3$) surrounding an artificial reef of length $L$, calculated as:

$$V_{fL} = V_{tL} - V_{rL}$$

where $V_{tL}$ ($m^3$) is the total volume of a reef of length $L$, including its surrounding foraging volume, calculated as:

$$V_{tL} = (L + D_{\text{max}} \times 2)^2 \times (H + D_{\text{max}})$$

where $D_{\text{max}}$ is the maximum distance ($m$) that a zooplanktivore will forage from shelter under the risk of predation. By using a generic function for $D_{\text{max}}$ (adapted from Biesinger et al. 2011), the model is applicable to a variety of reef-resident fishes and differing predator–prey relationships. $D_{\text{max}}$ was calculated as:

$$D_{\text{max}} = \frac{D_v}{S_{\text{pred}}} + 1$$

where $D_v$ is the mean viewing distance ($m$), $S_{\text{pred}}$ ($m \ s^{-1}$) is the swimming speed of a characteristic predator of zooplanktivorous fishes (for this model, the yellowtail kingfish Seriola lalandi), and $S_{\text{prey}}$ ($m \ s^{-1}$) is the swim speed of a zooplanktivore (for this model, A. strigatus). $S_{\text{pred}}$ and $S_{\text{prey}}$ were estimated using a derived relationship between swim speed, fish length and caudal fin aspect ratio (Sambilay 1990):

$$\log_{10}(S_a) = -0.828 + 0.6196\log_{10}(L) + 0.3478\log_{10}(AR) + 0.7621(Mo)$$

where $S_a$ is the absolute swim speed (km h$^{-1}$; converted to m $s^{-1}$ for use in Eq. 12), $L$ is the standard length of the fish (cm), $Mo$ is swimming mode (0 for sustained and 1 for burst speeds), and $AR$ is the aspect ratio of the caudal fin, calculated using Eq. 2 for A. strigatus ($n = 53$) and S. lalandi ($n = 8$). Burst speed (i.e. $Mo = 1$) was used as the swimming mode for both $S_{\text{pred}}$ and $S_{\text{prey}}$ based on the assumption that both species will maximise their swimming speeds during a predation event. The $S_{\text{pred}}$ value estimated for yellowtail kingfish using Eq. 13 agreed closely with empirical swim speed data for this species (S. Brodie unpubl. data).

Food availability was determined by dividing the total amount of zooplankton supplied to a reef’s foraging volume by the zooplankton required to sustain fish consumption (calculated using Eq. 1). A value $<1$ for this ratio shows that there is not enough zooplankton to support a population’s consumption. A value much larger than 1 suggests that there is more zooplankton available than can be consumed and food is unlikely to be a limiting factor. This ratio ($C_{\text{ratio}}$) is calculated as:

$$C_{\text{ratio}} = \frac{B_L}{C_t}$$

where $C_t$ is the total consumption of zooplankton by an individual zooplanktivore (mg fish$^{-1} \ d^{-1}$), and $B_L$ is the supply of zooplankton biomass to each zooplanktivore within the foraging volume of a reef of length $L$ (mg fish$^{-1} \ m^{-3}$), calculated as:

$$B_L = B_a \times V_{inL} \times R_L$$

where $V_{inL}$ ($m^3$ fish$^{-1}$) is the foraging volume available to an individual zooplanktivore, $R_L$ (periods $d^{-1}$) is the rate that the foraging volume is replaced by prevailing coastal currents, and $B_a$ is the available biomass of zooplankton allocated to an individual zooplanktivore for consumption (mg fish$^{-1} \ m^{-3}$), calculated as:

$$B_a = B_L \times P_{\text{con}}$$

where $P_{\text{con}}$ is the proportion of the total zooplankton supply allocated to the zooplanktivore for consumption, which is necessary to account for the requirements of other zooplanktivorous organisms on the reef (e.g. barnacles, ascidians, other fishes) and was given a mean value of 0.25 in this model based on relative biomass estimates (Scott et al. 2015, J. Smith unpubl. data). $B_L$ (mg $m^{-3}$) is the biomass of zooplankton capable of being foraged for by a zooplanktivore. $B_L$ was calculated as the amount of the minimum environmentally available biomass of zooplankton before predation observed at the artificial reef ($B_{\text{min}}$) that is available to A. strigatus, based on the observed prey size-selection.
\[ \text{Vin}_L = \frac{Vf_L}{N_L} \]  
where \( Vf_L \) (m\(^3\)) is the total foraging volume surrounding an artificial reef with a length \( L \) (Eq. 10), and \( N_L \) is the abundance of zooplanktivores within the same reef. \( R_L \) (periods d\(^{-1}\)) was calculated as:

\[ R_L = \frac{24 \times 60 \times 60}{\text{Per}} \]  
where \( \text{Per} \) is from Eq. 7, but replaces mean observed current velocity (Vel) with the minimum observed current velocity at this artificial reef (Vel\(_{\text{min}}\)) to conservatively estimate \( R_L \).

\( C_t \) was calculated as:

\[ C_t = \frac{Q/B \times W \times P_{\text{zoo}}}{365} \]  
where \( Q/B \) (yr\(^{-1}\)) is the consumption rate of a zooplanktivore as a function of biomass (Eq. 1), \( W \) (mg) is the mean weight of an individual zooplanktivore, and \( P_{\text{zoo}} \) is the observed mean proportion of a zooplanktivore's diet made up by zooplankton.

### Monte Carlo simulation and sensitivity analysis

Both Model 1 and Model 2 contained parameters with uncertain or variable values, so a Monte Carlo simulation was used to incorporate this parameter variation into model outputs. Each model was iterated 5000 times, and the mean and variance calculated from these iterations. Each parameter’s sampling distribution was either a normal, lognormal, or beta distribution (Table 1). The lognormal was used when there was evidence the data were skewed, and the beta distribution was used for proportion data. Standard deviations were generated from raw data wherever possible, and standard errors were used when the goal was to generate error in estimating the population mean, rather than variation at the individual level.

A sensitivity analysis was done to quantify the relative importance of parameter variables in both Model 1 and Model 2. A subset of parameters was selected from each model (Table 1), avoiding nested or obviously collinear parameters. The Monte Carlo simulation for each model was then used to randomly sample each parameter by its mean, or its mean ± 10%. This was done 5000 times, and the linear model was fitted to the resulting dataset (Smith et al. 2012) using parameters standardised according to Kleijnen (1997). The response variable in Model 1 was the depletion (%) of total zooplankton biomass, and the response variable for Model 2 was the reef length (m) at which the food supplied:food required ratio = 1.

### RESULTS

#### Diet of Atypichthys strigatus

\textit{A. strigatus} fed predominantly on zooplankton (~93% of the diet; Table 2), and crustaceans dominated the identifiable biomass of observed stomach contents (Fig. S1 in the Supplement). The proportion of copepod biomass within the diet of \textit{A. strigatus} was found to exceed all other identifiable prey groups combined (Table 2). Non-zooplanktonic prey items were infrequent components of the diet (<17% occurrence). Much of the diet was too digested for identification, but diet studies of mado in similar environments support the dominance of zooplankton in their diet (Glasby & Kingsford 1994, H. T. Schilling, J. A. Smith, J. D. Everett, D. P. Harrison, I. M. Suthers unpubl.). The parameter \( P_{\text{zoo}} \) was varied to account for this uncertainty (Table 1) and took values between 60 and 100% during simulations. The total mean wet biomass of \textit{A. strigatus} stomach contents was 466.4 mg (±39.3 SE), or 0.013 mg food per mg \textit{A. strigatus} (±0.001 SE).

### Table 2. Mean proportion (±SE) by wet mass of prey items in the diet of \textit{Atypichthys strigatus} (n = 55)

<table>
<thead>
<tr>
<th>Prey category</th>
<th>Proportion of \textit{A. strigatus} diet (%)</th>
<th>Occurrence (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Copepod</td>
<td>0.160 (0.012)</td>
<td>100</td>
</tr>
<tr>
<td>Shrimp</td>
<td>0.032 (0.006)</td>
<td>65.5</td>
</tr>
<tr>
<td>Ostracod</td>
<td>0.010 (0.002)</td>
<td>52.7</td>
</tr>
<tr>
<td>Amphipod</td>
<td>0.002 (0.001)</td>
<td>10.9</td>
</tr>
<tr>
<td>Zoa</td>
<td>0.014 (0.004)</td>
<td>41.8</td>
</tr>
<tr>
<td>Nauplius</td>
<td>0.001 (0.001)</td>
<td>5.5</td>
</tr>
<tr>
<td>Gastropod</td>
<td>0.009 (0.005)</td>
<td>10.9</td>
</tr>
<tr>
<td>Mollusc</td>
<td>0.013 (0.009)</td>
<td>5.5</td>
</tr>
<tr>
<td>Plant material</td>
<td>0.010 (0.004)</td>
<td>16.4</td>
</tr>
<tr>
<td>Unidentifiable Crustacea</td>
<td>0.167 (0.011)</td>
<td>100</td>
</tr>
<tr>
<td>Unidentifiable</td>
<td>0.583 (0.017)</td>
<td>100</td>
</tr>
<tr>
<td>Total zooplankton</td>
<td>0.929* (0.022)</td>
<td>–</td>
</tr>
</tbody>
</table>

*Excludes unidentifiable material
Foraging volume and density of *A. strigatus*

The observed density of *A. strigatus* declined as distance from the reef increased (Fig. 4) and *A. strigatus* were not observed farther than 4 m from the study reef. The density observed within the reef (3.81 fish m$^{-3}$ ± 0.67 SE) greatly exceeded the mean density of all foraging bins surrounding the reef (0.73 fish m$^{-3}$). The foraging volume available to *A. strigatus* increased non-linearly with increasing distance from the reef (Fig. S2 in the Supplement). Reef volume was calculated as 663.6 m$^3$ and the total foraging volume (within reef plus surrounding volume to 4 m from reef) was 2879 m$^3$.

Consumption by *A. strigatus*

*A. strigatus* at the artificial reef were estimated to consume on average 8.88 times their biomass annually ($Q/B$), which equated to an average of 0.77 g (0.10 SD) *A. strigatus$^{-1}$d$^{-1}$. Caudal fin aspect ratio did not significantly correlate with fish size ($r = 0.21$, $p = 0.14$, $n = 53$), and therefore the mean AR of *A. strigatus* (2.29 ± 0.47 SD) was applied to the multiple regression model (Eq. 1). Mean individual body mass of *A. strigatus* was calculated as 33.9 g (1.6 SE; $n = 55$).

Zooplankton supply and availability

*A. strigatus* exhibited prey size-selection, and the most commonly selected prey size was 601–800 μm, despite the smallest size classes of zooplankton (<200 and 200–400 μm) being the most environmentally abundant (Fig. S3A in the Supplement). The abundance of zooplankton prey items in size classes larger than 601–800 μm declined in approximate proportion to the environmental availability (Fig. S3A).

The observed mean total biomass of zooplankton up-current of the artificial reef was 871 mg m$^{-3}$ (±168 SE; Fig. S3B). The observed size-selection of *A. strigatus* for zooplankton <200, 200–400 and 401–600 μm was determined to equal zero, 0.02 and 0.46, respectively, while selection for size classes ≥601–800 μm were assumed all equal to 1. Thus, the mean biomass of zooplankton available to *A. strigatus* at this artificial reef under the observed pattern of prey size-selection was 637 mg m$^{-3}$ (±109 SE; Fig. S3B), which represented 73% of the total zooplankton biomass.

Model 1: Zooplankton depletion

The *A. strigatus* population at this study’s coastal artificial reef was estimated to consume 2906 g (±425 SD) of zooplankton per day, which is approximately 1.0 g m$^{-3}$ of reef habitat (reef and foraging volumes combined) per day. This equates to an average depletion of 0.35% (±0.13 SD) of the total zooplankton biomass delivered to the artificial reef (Fig. 5), or 0.49% (±0.17 SD) depletion of the available zooplankton biomass. The size-specific analysis of zooplankton depletion revealed that size class 601–800 μm were most depleted by *A. strigatus* predation; equal to 0.94% (1521 g) of the total biomass of that size class (Fig. 5). Zooplankton within size classes 200–400 and >1600 μm were least depleted by *A. strigatus* predation (excluding size class <200 μm, which was not consumed by *A. strigatus*), equal to 0.04% of the total binned biomass or 30 g d$^{-1}$ and 0.02% of the total binned biomass or 20 g d$^{-1}$, respectively. There was considerable variation in these depletion estimates (Fig. 5) due to the variation and uncertainty in the foraging volume, the supply of
zooplankton density and consumption (Table 1). Sensitivity analysis showed that all these model facets are equally influential in the model (Fig. S4A in the Supplement).

**Model 2: Reef size**

As the modelled reef became larger, foraging volume per unit reef volume decreased, which resulted in: (1) higher possible densities of reef-resident zooplanktivorous fish in the surrounding water, due to the increased refuge; and (2) a corresponding decline in zooplankton availability (Fig. 6). Given minimum observed values for current velocity ($V_{\text{min}}$) and zooplankton biomass ($B_{\text{min}}$; Table 1), the zooplankton supplied to reefs larger than ~40 m in length (between 25 and 55 m) would not support the required consumption rate of the maximum density of the resident zooplanktivore $A. strigatus$ (Fig. 6). The biomass of zooplankton available for consumption on reefs <40 m in length increased exponentially with decreasing reef size. The maximum density of the resident zooplanktivore in the foraging volume increased asymptotically with reef size (Fig. 6). These results are for square reefs 4 m in height, and increasing this height would decrease the reef length at which food limitation begins. Changing reef shape would also change these relationships. As in Model 1, there was considerable uncertainty around the estimates due to variation and uncertainty in model parameters (Table 2). The sensitivity analysis likewise showed that all parameters are equally influential for determining the reef size at food limitation (Fig. S4B).

**DISCUSSION**

This study revealed that a coastal artificial reef can support a large biomass of zooplanktivorous fish, demonstrating the value of zooplankton to fish biomass on such reefs. The small distance from the reef that these fish were willing to forage suggests that this zooplankton could not be exploited without the refuge this artificial reef provides. Approximately 3800 individual $A. strigatus$ with a biomass of ~130 kg populated the reef and immediate foraging volume. Although this species consumed predominantly zooplankton, its total consumption depleted less than 0.5% of the prevailing supply of zooplankton to this reef. Even though this coastal artificial reef received a substantial supply of zoo-
plankton, this study showed that increasing the size of artificial reefs negatively affects the ability of resident zooplanktivorous fish to achieve their required daily food ration, which will eventually limit fish density. Until this limitation occurs, however, increasing reef size allows for higher densities of zooplanktivorous fish within the surrounding foraging volume, and this abundance of prey fish may create benefits for recreational fishers. This illustrates the ecological cost of an increasing reef size—there is more refuge volume for reef residents but less foraging volume to support their consumption. These findings suggest that zooplankton can contribute greatly to resident fish production on coastal artificial reefs but that the dynamics of this trophic link is dependent on reef size, which is a factor that could be exploited when designing and deploying artificial reefs.

**Zooplankton consumption by reef fish**

The zooplanktivore *A. strigatus* population was estimated to consume a large amount of zooplankton at this artificial reef (2.9 kg d\(^{-1}\) ± 0.5 SD), yet this was only a tiny proportion of the average total zooplankton biomass supplied to the reef (0.35%). This indicates that the pelagic environment is a vast source of energy that can be incorporated into coastal reef systems via consumption by zooplanktivorous fishes, although the importance of this link to fish production is likely to be proportional to a reef’s exposure to ocean currents (Hamner et al. 1988). The large proportion of zooplankton in the *A. strigatus* diet, and their high abundance and resident behaviour, suggest that the supply of food from the pelagic environment is a very important driver of this reef’s function.

The input of pelagic energy into reef ecosystems via zooplanktivory may be common and integral to the function of coastal and offshore reefs (Hamner et al. 1988, Bortone 1998, Yahel et al. 2005). This study has only considered one abundant species of zooplanktivore, but large abundances of other reef-associated zooplanktivorous fishes such as yellowtail scad *Trachurus novaezelandiae* have also been recorded (NSW Department of Primary Industries 2013, Scott et al. 2015). Hence, the consumption values in this study only reflect a part of the total energy that is transferred from the pelagic environment to this reef area via fish zooplanktivory. What is more, planktivory by sessile invertebrates can also be very large (Glynn 1973, Reiswig 1974, Ayukai 1995), so our results may only represent a fraction of the coastal reef biomass supported by zooplankton. The combined consumption of zooplankton by reef-associated fishes and invertebrates forms a ‘pelagic pathway’ of energy to artificial and natural reefs alike (Kingsford & MacDiarmid 1988, Cresson et al. 2014), and it is the synthesis of this energy and transfer to higher trophic levels that is one of the key processes that can underpin biomass production on artificial reefs (Lindberg 1997, Leitão 2013, Cresson et al. 2014).

**Influence of reef size on zooplanktivorous fish**

The foraging behavior of reef-resident planktivores such as *A. strigatus* can be generally described as foraging the volume surrounding a central reef or refuge, within a distance of the refuge that allows for retreat to avoid predation (Hamner et al. 1988, Motro et al. 2005, Biesinger et al. 2011). A maximum foraging distance (*D*\(_{\text{max}}\)) defines this foraging volume surrounding reef habitats. An interesting outcome of a fixed *D*\(_{\text{max}}\) is a non-linear relationship between reef size and foraging volume. As a result, when reef size increases the foraging volume surrounding a reef declines relative to the reef refuge volume. This non-linearity was found to have an important influence on the dynamics of reef-resident zooplanktivorous fish.

The availability of zooplankton to zooplanktivorous fish declined as reef size increased, according to the relationship between refuge volume and foraging volume. Although zooplanktivorous fish may not forage to the extent of *D*\(_{\text{max}}\) when their zooplankton food is abundantly available, this foraging limit becomes more important as reef size (and fish abundance) increases and the ratio of food supplied to food required becomes increasingly small. For the reef system modelled in this study, reefs greater than ~40 m in length have insufficient foraging volume to feed the maximum density of resident zooplanktivorous fish, during the periods of lowest current flow (Fig. 6). In other words, large reefs have refuge for a high abundance of fish but lack the foraging volume in which to feed them. This means that the trophic relationship between zooplankton and reef-resident zooplanktivorous fishes becomes increasingly inefficient with increasing reef size, and the density of zooplanktivores should decline due to food limitation. Studies have observed higher fish densities on smaller reefs than on larger reefs (Bohnsack et al. 1994, Jordan et al. 2005), which provides support for this relationship.

An upside of increasing reef size is the increased provision of refuge volume, which can allow higher densities of reef-resident zooplanktivorous fish within
the surrounding foraging volume. Again, this is due to the non-linear relationship between the refuge and foraging volumes and the assumption that maximum fish density in the refuge volume is largely independent of reef size. In the modelled scenarios in this study, maximum zooplanktivore density began to asymptote on reefs with a length or width of ~50 m (Fig. 6), but the actual density would probably decline given density-dependent responses to the declining abundance of food.

The Monte Carlo simulations provided insight into the considerable variation and uncertainty in model parameters. The variation in zooplankton depletion, for example, depends on real variation in parameters such as current speed, zooplankton biomass, and water visibility but also on uncertainty in our estimates of parameters such as mean mado density and mean proportion of zooplankton in the diet. The sensitivity analysis revealed that all parameters are similarly influential, but targeting research towards a better understanding of species-specific foraging volumes could be a priority. This includes quantifying diurnal patterns in foraging and movement, especially as much planktivory may be diurnal (Yahel et al. 2005), as these patterns could further influence zooplankton availability. Future models may also benefit from incorporating density-dependence into calculations of fish abundance. In Model 2, the abundance of zooplanktivorous fish on a given reef was determined by the refuge volume and a fixed fish density within that volume. The relationships between reef size and both food supply and fish density assume that this 'refuge fish density' remains constant, but it is likely this density will vary in response to density-dependent competition for food and/or refuge. This study focused on a single species using data from a single reef, and much can be gained by exploring foraging patterns across multiple reefs and for multiple species, including accounting for interactions between species (Klages et al. 2014).

Management implications

This study has shown that the size of artificial reefs can have a great influence on resident zooplanktivorous fish, such that large reefs can be food limited and small reefs can be considered refuge limited. Both refuge volume and foraging volume are drivers of fish abundance, but the non-linear relationship between them means that small reefs can have an abundance of food but little refuge, while large reefs can have lots of refuge but insufficient foraging volume to support all possible residents. This suggests that an optimum reef size exists that can successfully trade-off between food and refuge. Understanding the limitations of artificial reefs that are either too small or overly large is essential for designing reefs that effectively facilitate the important trophic link between zooplankton and reef-resident fishes. These limitations are particularly important, as the size of an artificial reef can represent its construction cost, and the size of individual units can vary greatly (Bohnsack et al. 1994, Cresson et al. 2014, Scott et al. 2015). Large reefs cost more but do not necessarily optimise the trophic transfer of energy from zooplankton to reef-resident zooplanktivorous fishes. A common alternative to the deployment of one large reef is to create multiple smaller reefs that combined have the same reef volume but have independent foraging volumes that do not overlap (Bortone et al. 1998, Brandt & Jackson 2013). This strategy would be beneficial as it is likely to result in a similar abundance of individual reef-resident zooplanktivorous fishes but a larger per-capita food supply.

The orientation of artificial reefs to prevailing currents, and the effect of this on the trophic transfer of energy from the pelagic environment to the reef assemblage, is an aspect of reef design and deployment that has been discussed only rarely (Pickering & Whitmarsh 1997). It has been suggested the long axis of an artificial reef could be orientated perpendicular to the prevailing current (Mathews 1981, Pickering & Whitmarsh 1997) to maximise the supply of zooplankton to all zooplanktivorous reef animals. The findings in this study highlight the real value of this idea for optimising the production of zooplanktivores on coastal reefs. Artificial reefs could realistically be very long, and support a high abundance of reef-resident zooplanktivores, provided that this long axis is perpendicular to a dominant current direction. The effect of reef shape and size on the local hydrology and subsequent delivery of zooplankton also deserves investigation.

The benefit of optimising the production of zooplanktivorous fish will depend on an artificial reef’s management objectives, which include providing fishing opportunities, habitat offsets, and coastal protection (Baine 2001). For reefs deployed to provide fisheries benefits, increasing the presence and production of targeted species is a key objective, and optimising the production of zooplanktivores may promote this by increasing the presence of piscivorous fish. Zooplanktivores not only provide significant organic matter for benthic reef assemblages
(Bray et al. 1981, 1986), but they are also preyed upon by piscivorous species (Bulman et al. 2001, Young et al. 2010) commonly targeted by anglers. One of the most common species observed on the reef in this study is the prized piscivore yellowtail kingfish Seriola lalandi (Scott et al. 2015). The transfer of energy from zooplankton to local fish production on artificial reefs across multiple trophic levels remains to be quantified (Grossman et al. 1997), but it is likely that zooplanktivory by resident fish is a dominant process contributing to a larger food web.

In conclusion, this study showed that the trophic link between zooplankton and zooplanktivorous fishes is an important avenue of energy for reef assemblages, and one that probably contributes much to fish production on coastal artificial reefs. This study also revealed the trade-off between food supply and habitat supply as reef size changes and that food limitation is probably a key process driving resident zooplanktivore production on larger reefs. Ways to facilitate the consumption of zooplankton by zooplanktivorous fishes should be considered when planning future reefs, including the manipulation of reef size and shape, as this trophic link may have the greatest potential for enhancing the production of fish biomass from artificial reefs.

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