BLOOFINZ - Gulf of Mexico

Plankton food webs in the oligotrophic Gulf of Mexico spawning grounds of Atlantic bluefin tuna

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We used linear inverse ecosystem modeling techniques to assimilate data from extensive Lagrangian field experiments into a mass-balance constrained food web for the Gulf of Mexico open-ocean ecosystem. This region is highly oligotrophic, yet Atlantic bluefin tuna (ABT) travel long distances from feeding grounds in the North Atlantic to spawn there. Our results show extensive nutrient regeneration fueling primary productivity (mostly by cyanobacteria and other picophytoplankton) in the upper euphotic zone. The food web is dominated by the microbial loop (>70% of net primary productivity is respired by heterotrophic bacteria and protists that feed on them). By contrast, herbivorous food web pathways from phytoplankton to metazoan zooplankton process <10% of the net primary production in the mixed layer. Nevertheless, ABT larvae feed preferentially on podonid cladocerans and other suspension-feeding zooplankton, which in turn derive much of their nutrition from nano- and micro-phytoplankton (mixotrophic flagellates, and to a lesser extent, diatoms). This allows ABT larvae to maintain a comparatively low trophic level (~4.2 for preflexion and postflexion larvae), which increases trophic transfer from phytoplankton to larval fish.

KEYWORDS: plankton ecology; marine food web; calanoid copepods; larval fish; nitrogen cycle
INTRODUCTION

The open-ocean Gulf of Mexico (GoM) is a nutrient-poor, low-plankton-biomass region (Biggs and Ressler, 2001; Muller-Karger et al., 2015; Damien et al., 2018; Shropshire et al., 2020). Nevertheless, it is an important region for the spawning and larval development of many commercially important fishes (Lindo-Atichati et al., 2012; Rooker et al., 2012; Rooker et al., 2013; Kitchens and Rooker, 2014; Cornic et al., 2018). The western stock of Atlantic bluefin tuna (ABT) travel long distances from feeding grounds throughout the North Atlantic to spawning grounds in the oligotrophic GoM, implying that some characteristics of this region enhance larval success (Rooker et al., 2007; Teo et al., 2007; Rodriguez-Ezpeleta et al., 2019). One strong possibility is that the low abundances of potential predators on eggs and larvae and the reduced competition for prey in this food-poor region are a prerequisite for pelagic larvae to survive to maturity (Biggs, 1992; Muhling et al., 2017; Laiz-Carrion et al., 2019; Shropshire et al., this issue), but local enrichment processes such as fronts and eddies may also provide areas of higher productivity (Bakun, 2006), increasing the chances of larval survival (Bakun and Broad, 2003; Ciannelli et al., 2015). ABT larvae could then exploit the available food resources which can transfer biomass originated from microbial loops to tuna larvae despite the low mean primary productivity. Nevertheless, it remains unclear how ABT and other GoM larval fishes manage to obtain sufficient nutrition during their critical first-feeding period. Discerning the structure of GoM planktonic food webs is crucial to answering such questions.

ABT larvae are selective feeders that rely disproportionately on specific prey taxa, including calanoid and pycnoclastomatoid copepods, cladocerans and appendicularians (Llopiz et al., 2010; Llopiz et al., 2015; Tilley et al., 2016; Uriarte et al., 2019; Shiroza et al., this issue). These prey items, however, have distinctly different trophic and ecological roles (Landry et al., 2019). Appendicularians are filter-feeding pelagic tunicates with fine meshes that give them access to some of the smallest cyanobacteria in the ocean (Allerdige, 1976; Gorsky and Fenaux, 1998). Pycnoclastomatoid copepods, by contrast, are predators of other metazoan zooplankton and hence likely feed comparatively high on the food chain (Turner, 1986). Cladocerans and calanoid copepods are often omnivorous filter feeders, although calanoid copepods can fill multiple trophic roles within the planktonic food web, including as predators on other metazoans (Uye and Kayano, 1994; Mauchline, 1998; Katechakis and Stibor, 2004; Bode et al., 2015).

Elucidating the linkages between larval fish, their prey and the base of the planktonic food web is crucial to predicting climate change impacts on larval survival (Landry et al., 2019). Different phytoplankton groups (e.g. Prochlorococcus, Trichodesmium, diatoms and mixotrophic nanoflagellates) will respond differently to warming, acidification and increased stratification in the oligotrophic ocean (Rost et al., 2008; Flombaum et al., 2013; Flynn et al., 2013; Barton et al., 2016; Hong et al., 2017). These variable responses originate not only from different physiological responses to stressors but also due to the fundamentally different relationships between these groups and the limiting nutrient, light or temperature conditions. For instance: Trichodesmium and other diazotrophs (N2-fixing phytoplankton) are not nitrogen limited, Prochlorococcus is adapted to utilizing recycled nitrogen available at low concentrations in oligotrophic regions, and nanoflagellates may rely partially on phagotrophic behavior (mixotrophy) to alleviate nutrient stress (Scalan and Post, 2008; Zehr, 2011; Stoecker et al., 2017). The pathways that connect different nutrient sources (upwelling, lateral advection, recycled production and diazotrophy) through phytoplankton and zooplankton to larval fishes will determine how these organisms respond to climate change.

Here, we use linear inverse ecosystem models (LIEMs) as a data synthesis tool to constrain pelagic food webs of the oligotrophic GoM. We utilize results from field experiments designed to investigate the open-ocean GoM ecosystem from nutrients to fish (Gerard et al., this issue). LIEM allows us to incorporate diverse ecosystem measurements (e.g. primary productivity, protistan grazing rates, copepod δ15N and larval ABT gut contents) into a mass-balance constrained ecosystem model. We use the results to address four distinct questions: What is the trophic level (TL) of larval ABT? What is the trophic efficiency of food chains leading to larval ABT? Which phytoplankton groups ultimately support secondary production by larval ABT? What nitrogen sources support the specific food web pathways utilized by larval ABT?

METHODS

In situ measurements

Our data are derived from two cruises in ABT spawning grounds in April–May 2017 and 2018 as part of the Bluefin Larvae in Oligotrophic Ocean Foodwebs: Investigating Nutrients to Zooplankton in the Gulf of Mexico (BLOOFINZ–GoM) Project (Table I). During these cruises, we conducted regional zooplankton sampling.
### Table 1: Rate, biomass and δ¹⁵N measurements used as inputs to the inverse model

| Rate measurements | Units | C1       | C5       | Source                  |
|-------------------|-------|----------|----------|-------------------------|
| NPP (shallow)     | mmol N m⁻² d⁻¹ | 2.23 ± 0.13 | 3.08 ± 0.14 | Yingling et al. (this issue) |
| NPP (deep)        | mmol N m⁻² d⁻¹ | 1.64 ± 0.07 | 1.34 ± 0.05 | Yingling et al. (this issue) |
| f-ratio (shallow) | mmol N m⁻² d⁻¹ | 0.06 ± 0.04 | 0.14 ± 0.04 | Yingling et al. (this issue) |
| f-ratio (deep)    | mmol N m⁻² d⁻¹ | 0.44 ± 0.3  | 0.09 ± 0.02 | Yingling et al. (this issue) |
| Protistan grazing rate (shallow) | mmol N m⁻² d⁻¹ | 1.72 ± 0.6  | 2.58 ± 0.14 | Yingling et al. (this issue) |
| Protistan grazing rate (deep)    | mmol N m⁻² d⁻¹ | 1.44 ± 0.48 | 0.66 ± 0.18 | Yingling et al. (this issue) |
| Picophyto NPP (shallow) | mmol N m⁻² d⁻¹ | 1.15 ± 0.21 | 2.38 ± 0.37 | Landry et al. (this issue) |
| Picophyto NPP (deep) | mmol N m⁻² d⁻¹ | 0.78 ± 0.15 | 1.02 ± 0.17 | Landry et al. (this issue) |
| Flagellate NPP (shallow) | mmol N m⁻² d⁻¹ | 1.01 ± 0.37 | 0.69 ± 0.17 | Landry et al. (this issue) |
| Flagellate NPP (deep) | mmol N m⁻² d⁻¹ | 0.83 ± 0.35 | 0.32 ± 0.13 | Landry et al. (this issue) |
| Diatom NPP (shallow) | mmol N m⁻² d⁻¹ | 0.08 ± 0.03 | 0.01 ± 0 | Landry et al. (this issue) |
| Diatom NPP (deep) | mmol N m⁻² d⁻¹ | 0.02 ± 0.02 | 0 ± 0 | Landry et al. (this issue) |
| Picophyto mortality (shallow) | mmol N m⁻² d⁻¹ | 0.71 ± 0.32 | 1.6 ± 0.12 | Landry et al. (this issue) |
| Picophyto mortality (deep) | mmol N m⁻² d⁻¹ | 0.43 ± 0.07 | 0.34 ± 0.09 | Landry et al. (this issue) |
| Flagellate mortality (shallow) | mmol N m⁻² d⁻¹ | 0.86 ± 0.26 | 0.19 ± 0.11 | Landry et al. (this issue) |
| Flagellate mortality (deep) | mmol N m⁻² d⁻¹ | 0.6 ± 0.17 | 0.04 ± 0.02 | Landry et al. (this issue) |
| Diatom mortality (shallow) | mmol N m⁻² d⁻¹ | 0.04 ± 0.02 | 0.01 ± 0 | Landry et al. (this issue) |
| Diatom mortality (deep) | mmol N m⁻² d⁻¹ | 0.01 ± 0.01 | 0 ± 0 | Landry et al. (this issue) |
| NVM mesozoo grazing | mmol N m⁻² d⁻¹ | 0.39 ± 0.1 | 0.52 ± 0.1 | Landry and Swalethorp (this issue) |
| VM mesozoo grazing | mmol N m⁻² d⁻¹ | 0.03 ± 0.06 | 0.15 ± 0.08 | Landry and Swalethorp (this issue) |
| SedTrap flux (shallow) | mmol N m⁻² d⁻¹ | 1.53 ± 0.55 | 1.08 ± 0.07 | Stukel et al. (this issue) |
| SedTrap flux (deep) | mmol N m⁻² d⁻¹ | 0.46 ± 0.02 | 0.87 ± 0.18 | Stukel et al. (this issue) |
| Chl sinking (shallow) | mmol N m⁻² d⁻¹ | 0.02 ± 0.02 | 0.03 ± 0.02 | Stukel et al. (this issue) |
| Chl sinking (deep) | mmol N m⁻² d⁻¹ | 0.02 ± 0.01 | 0.05 ± 0.04 | Stukel et al. (this issue) |
| Fecal pellet sinking (shallow) | mmol N m⁻² d⁻¹ | 0.03 ± 0.03 | 0.02 ± 0.02 | Stukel et al. (this issue) |
| Fecal pellet sinking (deep) | mmol N m⁻² d⁻¹ | 0.13 ± 0.06 | 0.25 ± 0.2 | Stukel et al. (this issue) |
| Microozo to prefix | mmol N m⁻² d⁻¹ | 0.34 ± 0.16 | 0 ± 0 | Shiroza et al. (this issue) |
| Microzo to postflex | mmol N m⁻² d⁻¹ | 5.45 ± 0.88 | 0 ± 0 | Shiroza et al. (this issue) |
| Appendicularian to prefix | mmol N m⁻² d⁻¹ | 0.77 ± 0.32 | 0.04 ± 0.03 | Shiroza et al. (this issue) |
| Appendicularian to postflex | mmol N m⁻² d⁻¹ | 6.75 ± 0.47 | 1.88 ± 0.2 | Shiroza et al. (this issue) |
| Cladoceran to prefix | mmol N m⁻² d⁻¹ | 0.16 ± 0.16 | 0.78 ± 0.25 | Shiroza et al. (this issue) |
| Cladoceran to postflex | mmol N m⁻² d⁻¹ | 23.43 ± 2.26 | 26.47 ± 2.78 | Shiroza et al. (this issue) |
| Calanoids to prefix | mmol N m⁻² d⁻¹ | 3.61 ± 1.34 | 1.87 ± 0.35 | Shiroza et al. (this issue) |
| Calanoids to postflex | mmol N m⁻² d⁻¹ | 83.62 ± 0.96 | 13.49 ± 0.32 | Shiroza et al. (this issue) |
| Poecilastomatoids to prefix | mmol N m⁻² d⁻¹ | 0 ± 0.01 | 0 ± 0 | Shiroza et al. (this issue) |
| Poecilastomatoids to postflex | mmol N m⁻² d⁻¹ | 1.74 ± 2.63 | 0.89 ± 0.03 | Shiroza et al. (this issue) |

### Biomass and other measurements

| Temperature (0-50°C) | °C | 24.31 | 24.44 | CTD |
| Temperature (50-120°C) | °C | 22.14 | 21.68 | CTD |
| Temperature (120-300°C) | °C | 16.41 | 16.46 | CTD |
| HerbNVM biomass (shallow) | μmol N m⁻² | 23.24 | 45.63 | Shiroza et al. (this issue) |
| App biomass (shallow) | μmol N m⁻² | 0.22 | 0.32 | Shiroza et al. (this issue) |
| Clad biomass (shallow) | μmol N m⁻² | 0.06 | 0.12 | Shiroza et al. (this issue) |
| NVM Cal biomass (shallow) | μmol N m⁻² | 103.09 | 126.89 | Shiroza et al. (this issue) |
| Chaeto biomass (shallow) | μmol N m⁻² | 107.04 | 130.13 | Shiroza et al. (this issue) |
| Pocil biomass (shallow) | μmol N m⁻² | 28.65 | 18.53 | Shiroza et al. (this issue) |
| Preflex biomass (shallow) | μmol N m⁻² | 0.06 | 1.18 | Shiroza et al. (this issue) |
| Postflex biomass (shallow) | μmol N m⁻² | 0.62 | 1.91 | Shiroza et al. (this issue) |
| HerbNVM biomass | μmol N m⁻² | 88.62 | 14.86 | Shiroza et al. (this issue) |
| vmCal biomass | μmol N m⁻² | 78.29 | 14.86 | Shiroza et al. (this issue) |
| Cyano biomass (shallow) | mmol N m⁻² | 9.39 | 18.39 | Selph et al. (this issue) |
| Tricho biomass (shallow) | μmol N m⁻² | 27.23 | 0.97 | Selph et al. (this issue) |
| Diatom biomass (shallow) | mmol N m⁻² | 0.13 | 0.08 | Selph et al. (this issue) |
| Flag biomass (shallow) | mmol N m⁻² | 4.84 | 2.74 | Selph et al. (this issue) |
| Cyano biomass (deep) | mmol N m⁻² | 8.22 | 6.77 | Selph et al. (this issue) |
| Tricho biomass (deep) | μmol N m⁻² | 0.77 | 0.11 | Selph et al. (this issue) |
| Diatom biomass (deep) | mmol N m⁻² | 0.12 | 0.04 | Selph et al. (this issue) |
| Flag biomass (deep) | mmol N m⁻² | 6.97 | 3.63 | Selph et al. (this issue) |
| HerbNVM size | μg C ind⁻¹ | 1.35 | 1.35 | Shiroza et al. (this issue) |
| App size | μg C ind⁻¹ | 0.07 | 0.07 | Shiroza et al. (this issue) |
| Clad size | μg C ind⁻¹ | 0.68 | 0.68 | Shiroza et al. (this issue) |
| NVM Cal size | μg C ind⁻¹ | 4.44 | 4.44 | Shiroza et al. (this issue) |

Continued
surveys, which were guided partly by the Bluefin Tuna Index (Domingues et al., 2016), to identify contrasting open-ocean water parcels with and without high abundances of ABT larvae (Gerard et al., this issue). We then conducted 3- to 5-day Lagrangian experiments (hereafter ‘cycles’), while following satellite-enabled drift arrays with 3- to 1-m holey-sock drogues centered at 15 m depth, which allowed us to follow patches of mixed-layer water (Landry et al., 2009; Stukel et al., 2015). Five experimental cycles were conducted; in this study, however, we focus only on two experimental cycles with high larval ABT abundance—hereafter, Cycle 1 (C1) from the 2017 cruise and Cycle 5 (C5) from the 2018 cruise.

During each cycle, we conducted daily profiles with a CTD-Niskin rosette to measure temperature, salinity and density and to collect samples for chlorophyll a measurements (acidification method; Strickland and Parsons, 1972), phytoplankton pigment analyses (high-pressure liquid chromatography), picophytoplankton and heterotrophic bacteria enumeration (by flow cytometry; Selph et al., 2016; Selph et al., this issue), nano- and micro-phytoplankton biomass (Taylor and Landry, 2018; Selph et al., this issue), Trichodesmium biomass (Selph et al., this issue), nutrients (nitrate and ammonium; Knapp et al., this issue), dissolved organic nitrogen (DON; Knapp et al., this issue), particulate organic nitrogen (PON; Stukel et al., this issue) and δ15N of nitrate, DON and PON (Knapp et al., this issue; Stukel et al., this issue).

We also conducted a suite of daily in situ rate measurements that were incubated in mesh bags affixed at six depths spanning the euphotic zone on one of the floating arrays. These measurements included nitrate uptake (Yingling et al., this issue), net primary production (NPP) (Yingling et al., this issue) and group-specific phytoplankton growth and mortality due to protistan grazing (Landry et al., 2016; Landry et al., this issue). All in situ incubations were conducted for 24 h at natural light and temperature conditions. We also conducted shorter (4–6 h) shipboard incubations for nitrate and ammonium uptake (Yingling et al., this issue).

Twice per day (midday and midnight), we conducted oblique net tows through the euphotic zone to collect mesozooplankton that were analyzed for carbon, nitrogen, isotopes and gut pigment content (Landry and Swalethorp, this issue). Gut pigment contents were analyzed as in Décima et al. (2016) to estimate grazing rates (Landry and Swalethorp, this issue). ABT larvae were sampled frequently by standard double oblique tows (~8 tows d−1) with a 90-cm square bongo net (500-μm mesh) mounted with flowmeters to a depth of 25 m to ensure that we remained inside ABT habitat. Individual tuna larvae (2055 larvae, ranging from 3 to 9 mm length) were sorted onboard, and the identified ABT were liquid

| Units C1 | C5 | Source |
|---------|----|--------|
| Chaeto size | μg C ind⁻¹ | 20.83 | 20.83 | Shiroza et al. (this issue) |
| Poecili size | μg C ind⁻¹ | 5.33 | 5.33 | Shiroza et al. (this issue) |
| Preflex size | μg C ind⁻¹ | 83.71 | 83.71 | Shiroza et al. (this issue) |
| Postflex size | μg C ind⁻¹ | 179.75 | 179.75 | Shiroza et al. (this issue) |
| HerbVBM size | μg C ind⁻¹ | 4.44 | 4.44 | Shiroza et al. (this issue) |
| vmCal size | μg C ind⁻¹ | 4.44 | 4.44 | Shiroza et al. (this issue) |
| Maximum upwelling rate (shallow) | μmol N m⁻² d⁻¹ | 0.09 | 0.09 | Kelly et al. (in review) |
| Maximum upwelling rate (deep) | μmol N m⁻² d⁻¹ | 366.79 | 1543.33 | Kelly et al. (in review) |
| Maximum lateral advection of PON | mmol N m⁻² d⁻¹ | 3.22 | 3.22 | Kelly et al. (in review) |
| Maximum lateral advection of DON | mmol N m⁻² d⁻¹ | 1.56 | 1.56 | Kelly et al. (in review) |

δ15N values

- Upwelled nitrate: δ15N AIR (‰) 3.20 2.90 Knapp et al. (this issue)
- Preflex ABT: δ15N AIR (‰) 4.63 7.50 Swalethorp et al. (unpub.)
- Postflex ABT: δ15N AIR (‰) 4.21 6.16 Swalethorp et al. (unpub.)
- Shallow SedTrap: δ15N AIR (‰) 2.90 3.80 Stukel et al. (this issue)
- Deep SedTrap: δ15N AIR (‰) 4.89 4.55 Stukel et al. (this issue)
- Shallow DON: δ15N AIR (‰) 3.37 3.27 Knapp et al. (this issue)
- Deep DON: δ15N AIR (‰) 3.31 3.39 Knapp et al. (this issue)
- Shallow PON: δ15N AIR (‰) 1.44 2.66 Stukel et al. (this issue)
- Deep PON: δ15N AIR (‰) 1.80 1.63 Stukel et al. (this issue)
- Appendicularian: δ15N AIR (‰) 2.42 5.12 Swalethorp et al. (unpub.)
- Calanoid copepods: δ15N AIR (‰) 3.12 4.67 Swalethorp et al. (unpub.)
- Chaetognaths: δ15N AIR (‰) 5.70 7.58 Swalethorp et al. (unpub.)
- HerbVBM: δ15N AIR (‰) 4.73 5.88 Swalethorp et al. (unpub.)
- HerbNVMM: δ15N AIR (‰) 3.22 3.98 Swalethorp et al. (unpub.)
- Poeciliostomatoids: δ15N AIR (‰) 1.48 6.29 Swalethorp et al. (unpub.)
- Cladocerans: δ15N AIR (‰) 1.48 5.16 Swalethorp et al. (unpub.)

| Units C1 | C5 | Source |
|---------|----|--------|
| Deep PON | mmol N m⁻² d⁻¹ | 1.56 | 1.56 | Kelly et al. (in review) |
nitrogen- or ethanol-preserved for further analysis of abundance, body size, dry weight, gut content, otolith-based age and isotopic measurements (Laiz-Carrion et al., 2015; García et al., 2017; Malca et al., 2017; Laiz-Carrion et al., 2019; Malca et al., in prep.; Shiroza et al., this issue).

Nitrogen inputs to and outputs from the euphotic zone were constrained using sediment traps, Thorpe-scale analyses and remote-sensing products of lateral PON transport. Surface-tethered drifting sediment traps were used to collect sinking PON, chlorophyll and phaeopigments at 50 m depth, near the base of the euphotic zone (~120 m), and beneath the euphotic zone (200 m) (Stukel et al., this issue). We used Thorpe-scale analyses and nitrate concentration profiles to constrain vertical eddy diffusivity and upward nitrate flux (Gargett and Garner, 2008; Kelly et al., in review). We combined day–night differences in mesozooplankton biomass with allometric ammonium–excretion relationships to quantify active transport by diel vertical migrants (Ikeda, 1985; Landry and Swalethorp, this issue). We also quantified lateral transport of organic matter into the oligotrophic GoM using two independent approaches: a combination of remote-sensing-derived estimates of currents with remote-sensing-derived particulate carbon and a biogeochemical model developed for the open-ocean GoM (Shropshire et al., 2020; Kelly et al., in review).

Food web structure

Our food web structure was specifically designed to address the variability in trophic pathways within GoM foodwebs that channel energy toward the prey of ABT larvae (either efficiently or inefficiently) or to the multiple plankton taxa that are not suitable prey for the ABT larvae (Fig. 1). The model includes three inorganic N classes \( \text{NO}_3^- \), \( \text{NH}_4^+ \) and \( \text{N}_2 \) and three non-living organic matter pools [dissolved organic matter (DOM), small detritus and large detritus]. It includes four phytoplankton: \textit{Trichodesmium}, picophytoplankton (assumed to be potentially diazotrophic), diatoms and mixotrophic flagellates. It also includes heterotrophic bacteria, heterotrophic nanoflagellates and microzooplankton. Six suspension-feeding mesozooplankton are included: appendicularians (the only suspension feeders capable of feeding on cyanobacteria and heterotrophic bacteria), vertically migrating calanoid copepods, non-vertically migrating calanoid copepods, cladocerans, other non-vertically migrating herbivorous suspension feeders and other vertically migrating herbivorous suspension feeders. It includes two small predatory mesozooplankton: chaetognaths and poecilostomatoid copepods. It also includes four ‘higher TLs’ that serve as closure terms in the model: preflexion ABT larvae, postflexion ABT larvae, other planktivorous fish and predatory gelatinous zooplankton (e.g. ctenophores and cnidarians). ABT are assumed to feed on microzooplankton, appendicularians, cladocerans, non-vertically migrating calanoid copepods and poecilostomatoid copepods. Piscivory is not included in the model because field results showed that ichthyoplankton are not important prey to the 3–9-mm larvae (Shiroza et al., this issue). However, piscivory should be added if the model is used for larger larvae or in regions with higher ichthyoplankton densities. Other trophic pathways are determined based on known predator–prey relationships. Because ABT larvae feed only in the mixed layer, we include two layers in the model: upper euphotic zone (0–50 m) and deep euphotic zone (50–100 m on C1; 50–85 m on C5). All model compartments are identical, except that ABT larvae only exist in the upper euphotic zone. The two layers are connected through upward flux of nitrate, downward flux of sinking particles and the motions of vertical-migratory taxa, which are assumed to freely migrate into and between the two layers during the day, but reside beneath the euphotic zone (i.e. outside the model) during the day. Inputs to the model include upwelled nitrate, diazotrophy and lateral advection of particulate organic matter (POM) and DOM. Closure terms include secondary production of higher TLs, sinking of large detritus, sinking of diatoms, sinking of mixotrophic flagellates and excretion from vertical migratory taxa beneath the euphotic zone. We assume Redfield stoichiometry for all model flows, which allows us to relate respiration to ammonium excretion. We thus use the term ‘respiration’ when relating respiratory or excretory fluxes to primary production and the term ‘excretion’ when discussing nutrient recycling. Supplementary Table SI (see online supplementary data at Journal of Plankton Research online) shows all model flows.

Inverse model solution

To constrain the flux of nitrogen through unmeasured ecosystem pathways, we used LIEM techniques (Vézina and Platt, 1988; van Oevelen et al., 2010) to specify mass-balance constraints that must be exactly fit by food-web solutions, approximate equations that quantify measured rates with associated measurement uncertainty and inequality constraints that represent \textit{a priori} acceptable ranges for different ecosystem properties (e.g. gross growth efficiency varies from 10 to 40%). We used a total of 44 mass-balance constraints, 80 approximate equalities and 533 inequality constraints. However, with 302 total unknown food-web flows,
Fig. 1. Food web structure. All major food web flows between living organism groups are shown. However, for visual simplicity, we omit production of NH$_4^+$, DOM and detritus by all living groups as well as consumption of detritus by protistan zooplankton and suspension-feeding metazoans. Stars indicate groups at the highest TLs, for which secondary production is a model closure term. The model has a two-layer structure (≈ mixed layer and deep euphotic zone) with all trophic components in both layers, except for larval ABT. For all model flows, see Supplementary Table SI (see online supplementary data at Journal of Plankton Research online). HBac, heterotrophic bacteria; SDet, small detritus; LDet, large (sinking) detritus; Tricho, Trichodesmium; Cyano, cyanobacteria; Flag, mixotrophic flagellates; Dtm, diatoms; MIC, microzooplankton; HNF, heterotrophic nanoflagellates; App, appendicularians; HerbVM, vertically migrating herbivorous suspension feeders; HerbNVM, non-vertically migrating herbivorous suspension feeders; Clad, cladocerans; nvmCal, non-vertically migrating calanoid copepods; vmCal, vertically migrating calanoid copepods; Chaeto, chaetognaths; Poecil, poecilostomatoid copepods; Preflex ABT, preflexion ABT; Postflex ABT, postflexion ABT; Gel, gelatinous predators (ctenophores and cnidarians); Plank Fish, planktivorous fish.

The system remains under-constrained. To objectively determine representative solutions (and confidence limits) for all flows, we used the Markov Chain Monte Carlo (MCMC) with $^{15}$N approach (Stukel et al., 2018a, b). The MCMC approach conducts a random walk through the solution space that is constrained to fit the exact equations and bounded by the inequality constraints (Kones et al., 2009; Soetaert et al., 2009; Van den Meersche et al., 2009). New solutions are accepted based on the relative misfits of the new and previous solutions with respect to the approximate equality measurements. The mean solution of the MCMC approach has been shown to more accurately recover withheld measurement constraints than the previously used L$_2$ minimum norm approach (Stukel et al., 2012; Saint-Béat et al., 2013). The MCMC+$^{15}$N approach used herein allows for the incorporation of non-linear constraints associated with unknown $\delta^{15}$N values for some organisms or non-living nitrogen pools in the ecosystem to further constrain the system. For additional details, see the online supplementary appendix (see online supplementary data at Journal of Plankton Research online).

Food web analyses

TLs for all zooplankton were computed as one plus the ingestion-weighted mean TL of prey ($\text{TL}_{\text{consumer}} = \sum \text{TL}_{\text{prey},i} \times F_{\text{prey},i \rightarrow \text{consumer}} / \sum F_{\text{prey},i \rightarrow \text{consumer}}$, where $\text{TL}_{\text{prey},i}$ is the TL of prey $i$, and $F_{\text{prey},i \rightarrow \text{consumer}}$ is the rate of feeding of the consumer on prey $i$. All phytoplankton were assumed to have a TL of 1, except mixotrophic flagellates, which had TL = (1 – $p_{\text{phag}}$) + $p_{\text{phag}}$(1 + $\text{TL}_{\text{prey}}$), where $p_{\text{phag}}$ is the proportion of their nitrogen derived from phagotrophy (rather than dissolved nutrient uptake). Heterotrophic bacteria were assumed to have a TL equal to 1 plus the TL of the organism producing the organic matter they utilized.

To quantify indirect nitrogen flows through the food web, we used indirect food web flow analysis (Hannon,
The normalized amount of nitrogen (direct and indirect) that any organism derives from any other organism (or non-living nitrogen pool) can be computed as \((I - G)^{-1}\), where \(I\) is the identity matrix and \(G\) is the normalized production matrix (i.e., a matrix giving the percentage of an organism’s nitrogen requirement derived from any other organism).

Following Stukel et al. (2012), we defined three major food web pathways that describe energy and nutrient fluxes from the base of the food web: the herbivorous food chain, the multivorous food chain, and the microbial loop. (i) The herbivorous food chain = the sum of direct nitrogen flux from phytoplankton to metazoan zooplankton. (ii) The multivorous food chain = the sum of nitrogen flux that reaches metazoan zooplankton after passing through protistan grazers. (iii) The microbial loop = the sum of bacterial respiration and the fraction of protistan respiration that was supported by bacterial production. Results for each parameter are presented as means and 95% confidence intervals (CIs).

RESULTS

Model performance

The LIEM demonstrates close agreement with field measurements. The square root mean squared error (SRMSE), which can be thought of as the average number of standard errors that model estimates were from the measurements, was 1.17 for C1 if we consider only the field rate measurements and 1.53 for all approximate equality equations (including the \(\delta^{15}N\) mass-balance equations). For C5, the equivalent values were 1.40 and 1.65. One of the largest model-data mismatches was for sinking flux from the shallow to the deep euphotic zone during C1. The model struggled to find solutions that matched observations showing 3-fold higher sinking nitrogen flux from the upper euphotic zone to the lower euphotic zone than out of the euphotic zone. The model also slightly overestimated the grazing of suspension-feeding zooplankton on phytoplankton during both cycles, although in this case, the model’s 95% CIs overlapped the measured values. The model accurately recovered the ingestion rates of larval ABT. Model solutions were also strongly constrained by the comparatively low \(\delta^{15}N\) of larval ABT (Table I). The model struggled to determine solution vectors that matched the comparatively low \(\delta^{15}N\) of larval ABT with the fairly similar measured \(\delta^{15}N\) of upwelled nitrate, sinking detritus and bulk-suspended organic matter, thus leading to model misfits in the \(\delta^{15}N\) mass-balance equations.

Food web dynamics

Food web dynamics broadly reflected those expected for an oligotrophic, recycling-dominant ecosystem. \(\text{NH}_4^+\) was the dominant source of nitrogen to phytoplankton in the shallow euphotic zone (mean = 84%; 95% CI = 70–94% for C1 and 83%; 73–93% for C5). \(\text{NO}_3^-\) uptake (13%, 4–25% for C1; 16%, 6–25% for C5) and \(\text{N}_2\) fixation (1.2%, 0.03–4.3% for C1; 1.2%, 0.03–4% for C5) were comparatively less important.
Nutrient utilization patterns were broadly similar in the vicinity of the deep chlorophyll maximum (>50 m depth), although they varied between the two cycles with nitrate becoming substantially more important in the deep euphotic zone during C1 (42%, 13–72% for C1) than during C5 (10%, 4–17% for C5). Total production was slightly higher in the shallow euphotic zone (2.4 mmol N m$^{-2}$ d$^{-1}$, 2.2–2.6 mmol N m$^{-2}$ d$^{-1}$ for C1; 3.5, 3.2–3.7 mmol N m$^{-2}$ d$^{-1}$ for C5) than in the deep euphotic zone (1.8 mmol N m$^{-2}$ d$^{-1}$, 1.7–1.9 mmol N m$^{-2}$ d$^{-1}$ for C1; 1.3, 1.4–1.6 mmol N m$^{-2}$ d$^{-1}$ for C5).

Most primary production was from picophytoplankton (54%, 40–68% for C1; 79%, 68–90% for C5) and flagellates (42%, 28–55% for C1; 21%, 14–28% for C5) in the shallow euphotic zone, as suggested by the field data. Diatoms were comparatively less important (4.9%, 3.2–6.8% for C1; 0.4%, 0.3–0.5% for C5), while *Trichodesmium* production was negligible. The relative proportions of each group were fairly similar at the deep chlorophyll maximum. Mixotrophic flagellates derived 18% (C1) and 24% (C5) of their nitrogen from phagotrophy in the shallow euphotic zone (and slightly more in the deep euphotic zone). They consumed more heterotrophic bacteria than cyanobacteria.

Phytoplankton mortality was dominated by protistan grazing. These zooplankton (including mixotrophic flagellates) consumed 64% (49–79%) of phytoplankton production during C1 and 54% (47–61%) during C5. Metazoan zooplankton consumed a lower portion of phytoplankton production (20%, 14–26% for C1; 23%, 16–30% for C5), although they consumed more of the production of diatoms than protists did. Suspension-feeding metazoans also relied heavily on protistan zooplankton as dietary sources. This was reflected in trophic positions that averaged greater than 3.0 for all metazoans except appendicularians (Fig. 3a–d). In the upper euphotic zone, predatory zooplankton (poecilostomatoid copepods, chaetognaths and gelatinous predators) had particularly high trophic positions of 4.4, 4.4 and 4.7, respectively, for C1 (Fig. 3a) and 4.3, 4.3 and 4.6 for C5 (Fig. 3c). Their mean trophic positions in the deep euphotic zone were similar (4.3, 4.3 and 4.6 for C1, Fig. 3b; and 4.1, 4.1 and 4.4 for C5, Fig. 3d, for poecilostomatoid copepods, chaetognaths and gelatinous predators, respectively).

Quantification of major food web pathways showed that the GoM euphotic zone is dominated by the microbial loop (Fig. 4). The microbial loop (defined as respiration by heterotrophic bacteria and the proportion of protistan respiration supported by bacterial production) processed 70% (51–90%) of NPP in the shallow euphotic zone (Fig. 4a) and 77% (61–91%) of the NPP in the deep euphotic zone (Fig. 4b) during C1, whereas during C5, it used 71%; 58–84% in the upper euphotic zone (Fig. 4c) and 81%; 65–96% in the lower euphotic zone (Fig. 4d). For comparison, the herbivorous and multivorous food chains were responsible for processing 7.2% and 46% of NPP, respectively, in the shallow euphotic zone (Fig. 4a), and 37 and 46%, respectively, in the deep euphotic zone, (Fig. 4b) during C1. During C5, the herbivorous and multivorous food chains were responsible for 9.8% and 70% of NPP in the shallow euphotic zone (Fig. 4c) and for 54% and 44% in the deep euphotic (Fig. 4d). The dominance of microbial loop pathways aligns with the importance of recycled NH$_4^+$ for phytoplankton production and conforms with an expectation of tight recycling in oligotrophic ecosystems with limited new nutrient supply. In the shallow euphotic zone, where recycling and the microbial loop were most important, DON production was substantial (2.0 and 2.9 mmol N m$^{-2}$ d$^{-1}$, for C1 and C5). Phytoplankton and protistan zooplankton had large roles in DON production (38 and 32%, respectively) during C1, with the remainder primarily coming from dissolution of detritus (9.8%) and mesozooplankton excretion (11%). During C5, phytoplankton exudation was responsible for 47% of DON production, while protists were responsible for 28% and metazoan zooplankton contributed 15% of DON production. Bacterial excretion was in turn responsible for 50% of NH$_4^+$ regeneration in the shallow euphotic zone during C1 and for 46% during C5, with protist excretion generating an additional 32% (C1) or 29% (C5) and mesozooplankton excretion producing 15% (C1) or 20% (C5) of the NH$_4^+$ used by phytoplankton.

**Larval Atlantic bluefin tuna in the GoM ecosystem**

As suggested by the gut content data, model results show that larval ABT feed predominantly on cladocerans and calanoid copepods, with a lesser role for microzooplankton, appendicularians and poecilostomatoid copepods in their diets (Fig. 5). Calanoid copepods comprised 76% of the diet of preflexion ABT (95% CI = 59–88%) during C1 and 69% (55–83%) during C5. Microzooplankton (6%; CI = 1–13%) during C1; 0.4%, 0.03–0.9% during C5), appendicularians (14%; 4–26% during C1; 1.6%, 0.2–3.7% during C5) and cladocerans (4%; 0.3–9% during C1; 29%; 14–42% during C5) were smaller contributors to the diets of preflexion ABT, while poecilostomatoid copepods were negligible contributors to preflexion ABT diets (<0.7% during both cycles) (see Fig. 5a and c). Although calanoid copepods were also the dominant dietary source for postflexion ABT during
C1 (62%; 59–66%), these larger larvae also relied substantially on cladocerans (23%; 19–26% during C1; 62, 57–67% during C5) (see Fig. 5b and d).

The prevalence of suspension-feeding zooplankton in the diets of both preflexion and postflexion ABT led to relatively low TLs for ABT larvae (Fig. 3a and c). Given the ecosystem structure used in the model (Fig. 1), larval ABT could potentially have a TL between 3 and 7. However, both preflexion and postflexion larvae had TLs on the low end of this range. Preflexion ABT had a TL of 4.2 (4.0–4.5) during C1 and 4.2 (3.9–4.5) during C5, while postflexion ABT had TL estimates of 4.2 (4.0–4.5) during C1 and 4.1 (3.8–4.5) during C5. Both developmental stages of ABT larvae thus had trophic positions averaging ∼0.6 of their maximum possible TL (Fig. 6) and only one trophic position higher than their theoretically lowest possible TL within the food web. The trophic positions of larval ABT were thus notably low relative to those if feeding on the longest possible food chains that the model allowed. Based on this metric, their trophic positions were also notably lower than many of the zooplankton and other fish in the model.

The food chains supporting larval ABT were diverse and relied on significant production of picophytoplankton, flagellates and diatoms (while the production of *Trichodesmium* was insignificant for ABT food chains).
Preflexion ABT excreted $1.0 (0.4–2.0) \text{nmol N m}^{-2} \text{d}^{-1}$ derived from the production of flagellates, $0.9 (0.4–1.8) \text{nmol N m}^{-2} \text{d}^{-1}$ from picophytoplankton and $0.16 (0.05–0.37) \text{nmol N m}^{-2} \text{d}^{-1}$ from diatoms during C1 (Fig. 7a). During C5, preflexion ABT excreted $0.37 (0.14–0.79), 0.57 (0.28–0.99)$ and $0.06 (0.005–0.21) \text{nmol N m}^{-2} \text{d}^{-1}$ from flagellates, picophytoplankton and diatoms, respectively (Fig. 7c). Postflexion larvae excreted $17 (8–29) \text{nmol N m}^{-2} \text{d}^{-1}$ from flagellates, $13 (6.7–23) \text{nmol N m}^{-2} \text{d}^{-1}$ from picophytoplankton and $4.1 (1.2–9.2) \text{nmol N m}^{-2} \text{d}^{-1}$ from diatoms during C1 (Fig. 7a) and $4.7 (1.8–9.7), 6.7 (3.4–12)$ and $1.8 (0.14–5.3) \text{nmol N m}^{-2} \text{d}^{-1}$ during C5 (Fig. 7c). These values were influenced in large part by the different production rates of each phytoplankton taxa (flagellate, picophytoplankton and diatom NPP in the shallow euphotic zone were $1.8, 1.7$ and $0.24 \text{mmol N m}^{-2} \text{d}^{-1}$ during C1 and $1.5, 4.2$ and $0.015 \text{mmol N m}^{-2} \text{d}^{-1}$ during C5). When normalized to phytoplankton NPP, it becomes clear that larval ABT rely disproportionately on the production of large phytoplankton (Fig. 7b and f), even though large phytoplankton production is low in absolute terms. Preflexion ABT respired $1.1 \times 10^{-4}\% \ (C1)$ and
Nitrogen cycle and support of the upper euphotic zone ecosystem

In nitrogen-limited ecosystems, such as the open-ocean GoM, the supply of new nitrogen can control overall ecosystem productivity. Our results suggest that in the upper euphotic zone where ABT larvae feed, the ecosystem is not substantially supported by locally upwelled nitrate (which supplied $4.4 \times 10^{-3}$ (2.1 $\times 10^{-3}$–8.4 $\times 10^{-3}$) mmol N m$^{-2}$ d$^{-1}$ to the upper euphotic zone during C1 and 4.3 $\times 10^{-3}$ (2.3 $\times 10^{-3}$–8.4 $\times 10^{-3}$) mmol N m$^{-2}$ d$^{-1}$ during C5) or by nitrogen fixation [which supplied 0.092 (4.3 $\times 10^{-3}$–0.32) mmol N m$^{-2}$ d$^{-1}$ to the upper euphotic zone during C1 and 0.06 (0.002–0.2%) mmol N m$^{-2}$ d$^{-1}$ during C5]. Rather, nitrogen entered the ecosystem primarily through the lateral advection of organic matter (PON lateral advection = 0.18, 0.007–0.51 mmol N m$^{-2}$ d$^{-1}$ during C1; 0.94, 0.30–1.6 mmol N m$^{-2}$ d$^{-1}$ during C5; DON lateral advection = 0.19, 0.007–0.45 mmol N m$^{-2}$ d$^{-1}$ during C1; 0.10, 0.002–0.37 mmol N m$^{-2}$ d$^{-1}$ during C5)] from more productive regions (likely from shelf-break regions in the northern GoM, Gerard et al., this issue). Indeed, ABT derived only 0.2% (0.004–0.7%) of their nitrogen from upwelled nitrate during C1 and 0.1% (0.003–0.46%) during C5 and 2.2% (0.2–7.6%) and 0.95% (0.08–3.3%) of their nitrogen from nitrogen fixation during C1 and C5, respectively. They derived 98% (92–99%) and 99% (97–99%) from lateral advection.

This laterally advected organic matter entered the planktonic food web through multiple pathways. DON was utilized by bacteria, which had a gross growth efficiency of 27% (20–30%) during C1 and 28% (24–30%) during C5 and hence converted 73% of the DON they utilized to NH$_4^+$ during C1 and 72% during C5. The suspended particles (which presumably comprised laterally advected PON) were consumed primarily by protistan grazers or were converted into DON (likely through the activity of particle-attached microbes that were not explicitly included in our model). This highlights the importance of the microbial food web in mediating and enhancing phytoplankton in oligotrophic regions.

Indeed, even nitrate (which is often considered to be a ‘new’ nutrient in the euphotic zone) was primarily produced in situ by microbial activity (i.e. nitrification conducted by implicitly modeled ammonium-oxidizing bacteria). Modeled nitrification rates in the upper euphotic zone were 0.45 (0.15–0.83) mmol N m$^{-2}$ d$^{-1}$ during C1 and 0.08 (0.3–1.3) mmol N m$^{-2}$ d$^{-1}$ during C5. This equates to nitrification rates of 8.9 and 16 mmol N L$^{-1}$ d$^{-1}$ for C1 and C5, respectively. Notably, despite these low absolute nitrification rates, nitrate in the surface ocean was able to be regenerated every 2–3 days because nitrate concentrations were consistently

Figure 6. The ratio of the TL of different modeled zooplankton and fish to the TL they would have in the longest possible model food chain. The ‘other’ category includes all planktivorous fish and all zooplankton that are not larval ABT prey.
low in the upper euphotic zone. Considering that ammonium concentrations were \(\sim 90 \text{ nmol L}^{-1}\) during C1 and \(\sim 50 \text{ nmol L}^{-1}\) during C5, this suggests a specific ammonium-oxidation rate of 0.1 d\(^{-1}\) during C1 and 0.32 d\(^{-1}\) during C5. These results highlight the complexity of microbial and zooplankton linkages that support larval ABT in their oligotrophic nursery regions and suggest that the circulation of the GoM plays an important role in sustaining suitable conditions for larval growth.

DISCUSSION

The open-ocean GoM is an incredibly oligotrophic ecosystem with low productivity and a deep nitracline (Biggs, 1992; Gomez et al., 2018; Knapp et al., this issue; Yingling et al., this issue). Nevertheless, it is an important spawning ground for many migratory fish species, including multiple species of tuna, dolphinfish, sailfish and marlin (Rooker et al., 2012; Kitchens and Rooker, 2014; Cornic et al., 2018; Laiz-Carrión et al., 2019). It is also a region in which substantially depressed vertical mixing limits the phytoplankton productivity during ABT spawning periods (Gomez et al., 2018). If nutrient supply is indeed crucial for supporting these oligotrophic systems, predicted future warming and increased stratification could have deleterious impacts on taxa living in the mixed layer (Muhling et al., 2011; Liu et al., 2015; Muhling et al., 2015). Understanding how pelagic ecosystems and the larval fish they support will respond to climate change requires knowledge of the food web pathways that convert phytoplankton production into larval ABT.
into the preferred prey of different species (Landry et al., 2019).

We can hypothesize two potential ways in which an organism’s diet could make it well adapted to life in an oligotrophic region. First, it could feed preferentially on taxa that have either direct or indirect linkages to some of the most abundant primary producers in the ecosystem (e.g., cyanobacteria). For instance, a reliance on appendicularians would give larval fish access to a suspension feeder that can consume picophytoplankton directly (Gorsky and Fenaux, 1998; Lloptiz et al., 2010). Conversely, preference for calanoid copepods and podonid cladocerans would make a larval fish more dependent on the production of diatoms and other large phytoplankton. A second, but not mutually exclusive, hypothesis is that larval fish are more likely to thrive in oligotrophic ecosystems if they feed at a low trophic position, thus maximizing trophic transfer efficiency from phytoplankton to larvae regardless of the source of production.

Our results provide no evidence for the former hypothesis. Although diatom production only contributed to ~10% of ABT larval diets, it was a disproportionately high fraction of the ABT diet relative to the proportional role of diatoms to total NPP in the upper euphotic zone (~5% during C1, <1% during C5). Indeed, relative to a phytoplankton taxon’s productivity, the proportional contribution of each phytoplankton taxon to food-web pathways that support pre- and postflexion ABT larvae increased with increasing phytoplankter size from picophytoplankton to flagellates to diatoms (Fig. 7). The disproportionately large role of diatom-driven pathways was largely the result of the important role that podonid cladocerans played in ABT diets. Although they were only abundant in the water column during C5, our experiment conducted closest to the shelf break, podonids were consistently over-represented in ABT guts (Shiroza et al., this issue). Cladocerans are more frequently found in the coastal regions of the GoM, suggesting that they prey preferentially on large phytoplankton, as suggested by the LIEM and consistent with feeding studies (Kim et al., 1989; Katechakis and Stibor, 2004). Non-vertically migrating calanoid copepods, which formed another important component of ABT diets (because they were the most abundant mesozooplankton prey available to ABT), had a more diverse diet of diatoms, mixotrophic flagellates and heterotrophic protists. By contrast, while efficient pathways from cyanobacteria to ABT larvae can occur through appendicularians and microzooplankton, these taxa were not abundant in ABT guts. Appendicularians were rare in the water column, while microzooplankton were abundant but were seldom selected by ABT. The majority of cyanobacteria were consumed by heterotrophic nanoflagellates. These heterotrophic nanoflagellates had moderate gross growth efficiency in the model (30–36%) and were preyed upon by other protists (microzooplankton) and suspension-feeding metazoans. Cyanobacteria and heterotrophic nanoflagellates thus contributed disproportionately to the recycling pathways of the microbial loop, forming a largely distinct food web from the multivorous and herbivorous pathways, which mostly began with mixotrophic flagellates and diatoms and supported the production of larval ABT and other planktivorous fish. Despite distinct differences in prey selectivity with ontogeny (large cladocerans were much more important prey for postflexion larvae, Shiroza et al., this issue), our results show similar dependence on large phytoplankton for both larval stages.

Our results offer more support for the hypothesis that ABT larvae feed at a relatively low TL, maximizing the proportion of NPP available to them, and help explain how they survive in their oligotrophic spawning grounds (Fig. 6). The trophic position of ABT larvae (~4) is much closer to the minimum TL that our model allows (3: phytoplankton→prey→larvae) than to the maximum allowed TL (7: phytoplankton→bacteria→nanoflagellates→microzooplankton→suspension-feeders→carnivorous zooplankton→larvae). The low trophic position of ABT larvae is particularly striking, considering the relatively weak herbivorous food chain. Generally, planktivorous fish are more likely to be at a low TL in an ecosystem classically dominated by large phytoplankton and herbivorous mesozooplankton. However, the herbivorous food chain was responsible for only 7.2% (C1) or 9.8% (C5) of NPP processing in the shallow euphotic zone where the ABT larvae feed; the multivorous food chain processed 46% (C1) or 70% (C5) of NPP, while the microbial loop processed 70–71% of NPP on both cycles (Fig. 4, and note that the total exceeds 100% because NPP does not include phytoplankton DON production). The low trophic positions of ABT larvae were primarily due to two factors: (i) although total protistan secondary production was higher than total mesozooplankton secondary production, a comparatively small proportion of this secondary production made its way to larval tuna; most was dissipated as respiration in the microbial loop. Food chains supporting larval ABT were largely distinct from those involving the smallest class of heterotrophic protists. (ii) Both size classes of ABT larvae fed preferentially on podonid cladocerans, which fed lower in the food chain than other suspension-feeding taxa. Shiroza et al. (this issue) found selection for cladocerans to be an active process, further supporting the notion that ABT larvae are highly specialized at
maximizing trophic efficiency within their oligotrophic nurseries.

While the trophic position of ψ4 is low for a species known to preferentially feed on carnivorous copepods (poecilostomatoids) in a cyanobacteria- and microbial loop-driven ecosystem, we note that this is not actually a low TL relative to some other mass-balance constrained marine food web models. Many models based on ECOPATH software include only one (or zero) protistan trophic step and a single mesozooplankton group (Arreguin-Sanchez et al., 2004; Walters et al., 2008; Geers et al., 2016). These models constrain zooplankton to TLs 2 or 3; hence, the maximum allowed trophic position for planktivores is only 3 or 4. The additional complexity of our modeled ecosystem is a far more realistic depiction of natural food web complexity (Fig 1). Even so, our model allows only a maximum of two trophic steps within the protistan zooplankton (heterotrophic nanoflagellates and microzooplankton), which is an arbitrary limit, given the fluidity of protistan trophic interactions (Boenigk and Arndt, 2002; Pomeroy et al., 2007; Calbet, 2008; Caron et al., 2012; Sherr and Sherr, 2016). Some protists (e.g. pallium-feeding dinoflagellates) routinely feed at a 1:1 predator:prey size ratio, while others (e.g. ciliates) feed closer to a 10:1 predator:prey size ratio (Kiorboe, 2008; Fuchs and Franks, 2010). Some protists may consequently function at a higher trophic position than allowed by our model.

The BLOOFINZ–GoM study offers new insights to the physical dynamics of the GoM that support larval ABT. Kelly et al. (in review) analyzed vertical profiles of nitrate and buoyancy frequency from our cruises and concluded that exceedingly low amounts of nitrate were upwelled into the shallow euphotic zone where ABT spawn and their larvae grow. Instead, results from remote-sensing products and a 3D biogeochemical model provide compelling evidence that most nitrogen for export in the ABT habitat arrives via horizontal advection of organic matter. While our model constrains these inputs to be non-living organic matter (PON and DON), we note that a substantial proportion of this organic matter might be living plankton advected from more productive regions including the shelf-break region of the northern GoM and the Campeche Banks region north of the Yucatan Peninsula. Indeed, Gerard et al. (this issue) backtracked physical flows for the source of waters sampled in C1 and C5 to their origins 2–4 weeks previously along the shelf-slope margin in the northeastern GoM. Stukel et al. (this issue) found that 20% of particulate organic carbon in the upper euphotic zone was contained in living organisms. Landry and Swalethorp (this issue) further determined that (particularly during C5) predatory zooplankton likely relied on prey advected into our study region from more productive areas. Shropshire et al. (this issue) showed that ABT survival was also dependent on advection of prey from coastal areas and concluded that the most beneficial region for ABT spawning was near the shelf-break where prey are abundant for first-feeding larvae, but where offshore currents can transport larvae that survive the critical period to more oligotrophic regions before predation becomes a substantial source of mortality.

Our results show the importance of extensive recycling pathways for supporting phytoplankton production in this habitat. Despite the very low rates of vertical nitrate input and nitrogen fixation to the upper euphotic zone, sinking carbon flux from the upper euphotic zone was substantial (Stukel et al., this issue). This export, and indeed nearly all production in the upper euphotic zone, was supported by nutrients regenerated from PON through the activity of heterotrophic bacteria and protistan zooplankton. NH4+ was responsible for 85% of the production of phytoplankton in the upper euphotic zone, as is commonly the case in the mixed layer of oligotrophic, open-ocean regions (McCarthy et al., 1996; Lipschultz, 2001). However, in contrast to simple interpretations of nitrogen utilization, even NO3− was primarily generated autochthonously in the shallow euphotic zone and did not represent a ‘new’ form of nitrogen. The utility of nitrate as a tracer of ‘new’ production (Eppley and Peterson, 1979) has been extensively debated in the light of evidence of nitrification in shallow waters (Yool et al., 2007). The emerging consensus suggests that ammonium-oxidizing bacteria are not intrinsically light-inhibited (although some taxa may be), but rather they are often outcompeted in the euphotic zone by Prochlorococcus and other low-nutrient specialist phytoplankton (Smith et al., 2014; Wan et al., 2018; Xu et al., 2019). Our results do not contradict this view. Indeed, the LIEM suggests that phytoplankton utilize NH4+ more rapidly than ammonium-oxidizing bacteria. However, the low NO3− concentrations throughout the euphotic zone (Knapp et al., this issue) and exceedingly low NO3− flux (Kelly et al., in review) allow nitrification to dominate NO3− supply despite low absolute nitrification rates. Indeed, our estimate of the specific rate of ammonium oxidation necessary to support phytoplankton NO3− utilization (0.1–0.3 d−1) is near the median value for surface ocean ammonium oxidation in the synthesis of Yool et al. (2007). Notably, Clark et al. (2008) measured ammonium and nitrite oxidation rates in the oligotrophic regions of the Atlantic Ocean which were slightly lower than our LIEM-predicted values, and Bronk et al. (2014) measured substantially higher nitrification rates in the offshore regions of the West Florida Shelf. Nitrification rate measurements from other regions have been highly
variable, and there is not, as yet, a consensus on the relative importance of shallow nitrification to NO$_3^-$ supply in oligotrophic regions (Newell et al., 2013; Clark et al., 2016; Shiozaki et al., 2016).

The importance of laterally advected organic matter for supporting oligotrophic communities in the GoM offers important insight into the physical characteristics that make the GoM an ideal spawning habitat for ABT. While previous studies have focused on the role of vertical mixing and upwelling, our results show that mixed layer productivity may be more directly tied to horizontal fluxes driven by the high mesoscale energy of the GoM. In the oligotrophic GoM, the Loop Current and the eddies that it sheds are prominent features enhancing circulation (Forristall et al., 1992; Oey et al., 2005; Schmitz et al., 2005). These features have the potential to fundamentally restructure open-ocean ecosystems, with warm-core eddies (including Loop Current Eddies) depressing the nutricline and primary production, while cold-core eddies increase open-ocean upwelling and productivity (Biggs and Müller-Karger, 1994). These altered nutrient supply and phytoplankton regimes lead to substantially higher zooplankton biomass in cold-core eddies (Wells et al., 2017). However, the relative importance of each eddy type, as well as the distinct gradient regions that form on their edges, on larval ABT remains a topic of active debate (Muhling et al., 2010; Domingues et al., 2016). Our results suggest that both eddy types can be important nitrogen sources to the upper euphotic zone since the high horizontal velocities along the eddy can transport living and non-living organic matter from high biomass regions to oligotrophic areas, especially when eddies impinge on coastal regions. Shropshire et al. (2020) also found substantial transport into our study region mediated by entrainment of plankton-rich waters from the Campeche Bank into the edges of the Loop Current. Notably, the larvae distribution in the major recognized ABT eastern stock spawning area, around the Balearic Islands in the western Mediterranean basin, is influenced from frontal zones resulting from the convergence of recent and resident Atlantic surface waters (Alemany et al., 2010; Muhling et al., 2017; Reglero et al., 2017). Such mesoscale features have been hypothesized to act as retention larval feeding areas, enhancing particle food concentrations and increasing the probability of survival of larvae that rely substantially on copepods and cladoceran prey during preflexion stage in this oligotrophic environment (Catalán et al., 2011; Uriarte et al., 2019). Horizontal flows associated with these features may also connect the nearby coastal region to oligotrophic nursery areas, a possibility that should be explored in future studies.

The potential importance of cross-shore fluxes to survival of first-feeding ABT suggests that determining the responses of pelagic food webs and ABT larvae to climate change will require characterizing changes in GoM circulation in response to future forcing along with the expected food web processes that regenerate nutrients and promote growth of larval ABT prey (Muhling et al., 2011; Liu et al., 2015). Our study offers insight into the processes allowing larval ABT to survive in a food-scarce environment. However, substantial additional research is needed to quantify the impacts of spatial and interannual variability, as well as secular change, on these ecosystems and threatened species.

**CONCLUSION**

ABT larvae develop in oligotrophic ecosystems, dominated by cyanobacteria and other small phytoplankton. The major trophic pathway through the microbial loop is highly inefficient, with most production lost to remineralized nutrients by bacteria and multi-step protistan grazing chains. Both pre- and postflexion larval ABT feed preferentially on less dominant pathways associated with herbivorous and multivorous food chains, without pronounced ontogenetic differences in food-web roles between pre- and postflexion stages, despite distinct changes in diet. Consequently, ABT larvae depend on the production of diatoms and mixotrophic flagellates that support herbivorous zooplankton, particularly calanoid copepods and cladocerans. Preferential utilization of these more direct trophic pathways allows the larvae to feed at relatively low TLs despite the fact that the taxa responsible for the majority of secondary production in the food web (bacteria and heterotrophic nanoflagellates) are not accessible to them as prey. Further research is needed to understand how these ecological interactions might be altered under different disturbance regimes.

**DATA AVAILABILITY**

Data utilized in this manuscript are available on NCCOS and BCO-DMO (https://www.bco-dmo.org/project/819488). MCMC+15N model code is available on Github: https://github.com/stukel-lab. The specific code used to run the BLOOFINZ-GoM inverse model, with set-up files and instructions for running it can be accessed at: https://github.com/mstukel/N15-LIM-BLOOFINZ-GoM

**SUPPLEMENTARY DATA**

Supplementary data can be found at *Journal of Plankton Research* online.
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REFERENCES

Alemany, F., Quintanilla, L., Veláz-Belchí, P., García, A., Cortés, D., Rodríguez, J. M., DE Puuelles, M. F., González-Pola, C. et al. (2010) Characterization of the spawning habitat of Atlantic bluefin tuna and related species in the Balaenica Sea (western Mediterranean). Prog. Oceans. 86, 21–38.

Allerdige, A. (1976) Field behavior and adaptive strategies of appendicularians (Chordata: Tunicata). Mar. Biol. 38, 29–39.

Arreguin-Sanchez, F., Zetina-Rejon, M., Manickchand-Heileman, S., Ramirez-Rodriguez, M. and Vidal, L. (2004) Simulated response to harvesting strategies in an exploited ecosystem in the southwestern Gulf of Mexico. Ecol. Model. 172, 421–432.

Bakun, A. (2006) Fronts and eddies as key structures in the habitat of marine fish larvae: opportunity, adaptive response and competitive advantage. Sci. Mar. 70, 105–122.

Bakun, A. and Broad, K. (2003) Environmental ‘loopholes’ and fish population dynamics: comparative pattern recognition with focus on El Niño effects in the Pacific. Fish. Oceanogr. 12, 458–473.

Barton, A. D., Irwin, A. J., Finkel, Z. V. and Stock, C. A. (2016) Anthropogenic climate change drives shift and shuffle in North El Niño effects in the Pacific. J. Geophys. Res. Oceans. 121, 7371–7384.

Biggs, D. C. (1992) Nutrients, plankton, and productivity in a warm-core ring in the western Gulf of Mexico. J. Geophys. Res. Oceans. 97, 2143–2154.

Biggs, D. C. and Muller-Karger, F. E. (1994) Ship and satellite observations of chlorophyll stocks in interacting cyclone-anticyclone eddy pairs in the western Gulf of Mexico. J. Geophys. Res. Oceans. 99, 7371–7384.

Biggs, D. C. and Ressler, P. H. (2001) Distribution and abundance of phytoplankton, zooplankton, ichthyoplankton, and micronekton in the deepwater Gulf of Mexico. Gulf Mex. Sci. 19, 2.

Bode, M., Hagen, W., Schukat, A., Teuber, L., Fonseca-Batista, D., Dehairs, F. and Auel, H. (2015) Feeding strategies of tropical and subtropical calanoid copepods throughout the eastern Atlantic Ocean - latitudinal and bathymetric aspects. Prog. Oceans. 138, 268–282.

Boenigk, J. and Arndt, H. (2002) Bacterivory by heterotrophic flagellates: community structure and feeding strategies. Anton. Lenuc. Int. J. G. 81, 465–480.

Bronk, D. A., Killeberg-Thoresen, L., Sipler, R. E., Mulholland, M. R., Roberts, Q. N., Bernhardt, P. W., Garrett, M., O’Neil, J. M. et al. (2014) Nitrogen uptake and regeneration (ammonium regeneration, nitrification and photoproduction) in waters of the West Florida Shelf prone to blooms of Karenia brevis. Harmful Algae. 38, 50–62.

Calbet, A. (2008) The trophic roles of microzooplankton in marine systems. ICES J. Mar. Sci. 65, 325–331.

Caron, D. A., Countway, P. D., Jones, A. C., Kim, D. Y. and Schnetzer, A. (2012) Marine protist diversity. Ann. Rev. Mar. Sci. 4, 467–493.

Catalán, I. A., Tejedor, A., Alemany, F. and Reglero, P. (2011) Trophic ecology of Atlantic bluefin tuna Thunnus thynnus larvae. J. Fish Biol., 78, 1545–1560.

Ciannelli, L., Bailey, K. and Olsen, E. M. (2015) Evolutionary and ecological constraints of fish spawning habitats. ICES J. Mar. Sci., 283–296.

Clark, D. R., Rees, A. P. and Joint, I. (2008) Ammonium regeneration and nitrification rates in the oligotrophic Atlantic Ocean: implications for new production estimates. Lanniol. Ocean., 53, 32–62.

Clark, D. R., Widicombe, C. E., Rees, A. P. and Woodward, E. M. S. (2016) The significance of nitrogen regeneration for new production within a filament of the Mauritanian upwelling system. Bioseasciences, 13, 2873–2888.

Cornic, M., Smith, B. L., Kitchens, L. L., Bremer, J. R. A. and Rooker, J. R. (2018) Abundance and habitat associations of tuna larvae in the surface water of the Gulf of Mexico. Hydrobiologia, 806, 29–46.

Damien, P., Pasqueron de Fommervault, O., Scheinbaum, J., Jouanno, J., Camacho-Ibar, V. F. and Duteil, O. (2018) Partitioning of the open waters of the Gulf of Mexico based on the seasonal and interannual variability of chlorophyll concentration. J. Geophys. Res. Oceans, 123, 2592–2614.

Décima, M., Landry, M. R., Stukel, M. R., Lopez-Lopez, L. and Krause, J. W. (2016) Mesozooplankton biomass and grazing in the Costa Rica Dome: amplifying variability through the plankton food web. J. Plankton Res., 38, 317–330.

Domínguez, R., Goni, G., Bringas, F., Munhing, B., Lindo-Aitchi, D. and Walter, J. (2016) Variability of preferred environmental conditions for Atlantic bluefin tuna (Thunnus thynnus) larvae in the Gulf of Mexico during 1993–2011. Fish. Oceanogr., 25, 320–336.

Eppley, R. W. and Peterson, B. J. (1979) Particular organic matter flux and planktonic new production in the deep ocean. Nature, 282, 677–680.

Flombaum, P., Gallegos, J. L., Gordillo, R. A., Rincon, J., Zahala, L. L., Jiao, N., Karl, D. M., Li, W. K. W. et al. (2013) Present and future global distributions of the marine cyanobacteria Prochlorococcus and Synechococcus. Proc. Natl. Acad. Sci. U. S. A., 110, 9024–9029.

Flynn, K. J., Stoecker, D. K., Mitra, A., Raven, J. A., Gilbert, P. M., Hansen, P. J., Graneli, E. and Burkholler, J. M. (2013) Misuse of the phytoplankton-zooplankton dichotomy: the need to assign organisms as mixotrophs within plankton functional types. J. Plankton Res., 35, 3–11.

Forristall, G. Z., Schaudt, K. J. and Cooper, C. K. (1992) Evolution and kinematics of a loop current eddy in the Gulf of Mexico during 1983. J. Geophys. Res. Oceans., 97, 2173–2184.
Fuchs, H. L. and Franks, P. J. S. (2010) Plankton community properties determined by nutrients and size-selective feeding. *Mar. Ecol. Prog. Ser.*, 413, 1–15.

García, A., Laiz-Carrión, R., Uriarte, A., Quintanilla, J. M., Morote, E., Rodríguez, J. M. and Alemay, F. (2017) Differentiated stable isotopes signatures between pre-and post-flexion larvae of Atlantic bluefin tuna (*Thunnus thynnus*) and of its associated tuna species of the Balearic Sea (NW Mediterranean). *Deep-Sea Res. II*, 140, 18–24.

Gargett, A. and Garner, T. (2008) Determining Thorpe scales from ship-lowered CTD density profiles. *J. Atmos. Oceanic Tech.*, 25, 1657–1670.

Geers, T. M., Pilkett, E. K. and Frisk, M. G. (2016) An original model of the northern Gulf of Mexico using Ecopath with Ecosim and its implications for the effects of fishing on ecosystem structure and maturity. *Deep-Sea Res. II*, 129, 319–331.

Gerard, T., Lamkin, J., Kelly, T. B., Knapp, A. N., Laiz-Carrión, R., Malca, E., Selph, K. E., Shiraza, A. et al. (this issue) Bluefin larvae in Oligotrophic Ocean Foodwebs, investigations of nutrients to zooplankton: overview of the BLOOFINZ-Gulf of Mexico program. *J. Plankton Res*.

Gomez, F. A., Lee, S.-K., Liu, Y., Hernandez, F. J. Jr., Muller-Karger, F. E. and Lamkin, J. T. (2018) Seasonal patterns in phytoplankton biomass across the northern and deep Gulf of Mexico: a numerical model study. *Biogeosciences*, 15, 3561–3576.

Gorsky, G. and Fenaux, R. (1998) The role of Appendicularia in marine food webs. In Bone (ed.), *The Biology of Pelagic Tunicates*, pp. 161–169.

Hannon, B. (1973) Structure of ecosystems. *J. Theor. Biol.*, 41, 535–546.

Hong, H., Shen, R., Zhang, F., Wen, Z., Chang, S., Lin, W., Kranz, S. A., Loo, Y-W. et al. (2017) The complex effects of ocean acidification on the prominent N2-fixing cyanobacterium *Trichodesmium*. *Science*, 356, 327–331.

Ikeda, T. (1985) Metabolic rates of epipelagic marine zooplankton as a function of body mass and temperature. *Mar. Biol.*, 85, 1–11.

Katechakis, A. and Stibor, H. (2004) Feeding selectivities of the marine cladocerans *Podon avoros*, *Podon intermedias* and *Eudore normannii*. *Mar. Biol.*, 145, 329–339.

Kelly, T. B., Landry, M. R., Selph, K. E., Knapp, A. N., Swalethorp, R. and Stukel, M. R. (in review) Lateral advection supports nitrogen export in the oligotrophic ecosystem of the open-ocean Gulf of Mexico. *Nat. Commum*.

Kim, S., Onbé, T. and Yoon, Y. (1989) Feeding habits of marine cladocerans in the Inland Sea of Japan. *Mar. Biol.*, 100, 313–318.

Kierboe, T. (2008) *A Mechanistic Approach to Plankton Ecology*, Vol., Princeton University Press, Princeton, NJ.

Kitchens, L. L. and Rooker, J. R. (2014) Habitat associations of dolphinfish larvae in the Gulf of Mexico. *Fish. Oceanogr.*, 23, 460–471.

Knapp, A. N., Thomas, R., Stukel, M. R., Kelly, T. B., Landry, M. R., Selph, K. E., Malca, E., Gerard, T. et al. (this issue) Constraining the sources of nitrogen fueling export production in the Gulf of Mexico using nitrogen isotope budgets. *J. Plankton Res*.

Körsch, J. K., Sonkaert, K., van Oevelen, D. and Owino, J. J. O. (2009) Are network indices robust indicators of food web functioning? A Monte Carlo approach. *Ecol. Model.*, 220, 370–382.

Laiz-Carrion, R., Gerard, T., Sucua, J. J., Malca, E., Uriarte, A., Quintanilla, J. M., Privoznik, S., Llopis, J. K. et al. (2019) Stable isotope analysis indicates resource partitioning and trophic niche overlap in larvae of four tuna species in the Gulf of Mexico. *Mar. Ecol. Prog. Ser.*, 619, 53–68.

Laiz-Carrión, R., Gerard, T., Uriarte, A., Malca, E., Quintanilla, J. M., Muhling, B. A., Alemany, F., Privoznik, S. L. et al. (2015) Trophic ecology of Atlantic bluefin tuna (*Thunnus thynnus*) larvae from the Gulf of Mexico and NW Mediterranean spawning grounds: a comparative stable isotope study. *Plos One*, 10, e0133406. doi:10.1371/journal.pone.0133406.

Landry, M. R., Beckley, L. E. and Muhling, B. A. (2019) Climate sensitivities and uncertainties in food-web pathways supporting larval bluefin tuna in subtropical oligotrophic oceans. *ICES J. Mar. Sci.*, 76, 359–369.

Landry, M. R., Ohman, M. D., Goericke, R., Stukel, M. R. and Tsyrlkevich, K. (2009) Lagrangian studies of phytoplankton growth and grazing relationships in a coastal upwelling ecosystem off Southern California. *Prog. Oceanogr.*, 83, 208–216.

Landry, M. R., Selph, K. E., Decima, M., Gutierrez-Rodriquez, A., Stukel, M. R., Taylor, A. G. and Pasulka, A. L. (2016) Phytoplankton production and grazing balances in the Costa Rica Dome. *J. Plankton Res.*, 38, 366–379.

Landry, M. R., Selph, K. E., Stukel, M. R., Swalethorp, R., Kelly, T. B., Beatty, J. and Quackenbush, C. R. (this issue) Microbial food web dynamics in the oceanic Gulf of Mexico. *J. Plankton Res*.

Landry, M. R. and Swalethorp, R. (this issue) Mesozooplankton biomass, grazing and trophic structure in the bluefin tuna spawning area of the oceanic Gulf of Mexico. *J. Plankton Res. doi: 10.1093/plankt/fbah008*.

Lindo-Avichat, D., Bringas, F., Goni, G., Muhling, B., Muller-Karger, F. E. and Habtes, S. (2012) Varying mesoscale structures influence larval fish distribution in the northern Gulf of Mexico. *Mar. Ecol. Prog. Ser.*, 463, 245–257.

Lipschultz, F. (2001) A time-series assessment of the nitrogen cycle at BATS. *Deep-Sea Res. II*, 48, 1897–1924.

Liu, Y., Lee, S.-K., Enfield, D. B., Muhling, B. A., Lamkin, J. T., Muller-Karger, F. E. and Roffer, M. A. (2015) Potential impact of climate change on the Itrras-Americas Sea: Part-1. A dynamic downscaling of the CMIP5 model projections. *J. Mar. Syst.*, 148, 56–69.

Llojize, J. K., Muhling, B. A. and Lamkin, J. T. (2015) Feeding dynamics of Atlantic bluefin tuna (*Thunnus thynnus*) larvae in the Gulf of Mexico. *Collet. Vol. Sci. Pap. ICAT 71*, 1710–1715.

Llojize, J. K., Richardson, D. E., Shiraza, A., Smith, S. L. and Cowen, R. K. (2010) Distinctions in the diets and distributions of larval tunas and the important role of appendicularians. *Lanmod. Oceanogr.*, 55, 983–996.

Malca, E., Muhling, B., Franks, J., García, A., Tilley, J., Gerard, T., Ingram, W. J. and Lamkin, J. T. (2017) The first larval age and growth curve for bluefin tuna (*Thunnus thynnus*) from the Gulf of Mexico: comparisons to the Straits of Florida, and the Balearic Sea (Mediterranean). *Fish. Res.*, 190, 24–33.

Malca, E., Swalethorp, R., Quintanilla, J. M., Laiz-Carrión, R., Shropshire, T. A., Kelly, T. B., Shiraza, A., Stukel, M. R. et al. (in prep.) Influence of habitat and food quality on growth rate of bluefin tuna larvae in the Gulf of Mexico. *J. Plankton Res.*, this issue.

Mauchline, J. (1998) *Adv. Mar. Biol.: The Biology of Calanoid Copepods*, Vol., Academic Press, San Diego.

McCarthy, J. J., Garside, C., Nevin, J. L. and Barber, R. T. (1996) New production along 140°W in the equatorial Pacific during and following the 1992 El Niño event. *Deep-Sea Res. II*, 43, 1085–1093.

Muhling, B. A., Lamkin, J. T., Alemany, F., García, A., Farley, J., Ingram, G. W., Berastegui, D. A., Reglero, P. et al. (2017) Reproduction and larval biology in tunas, and the importance of restricted area spawning grounds. *Rev. Fish. Biol. Fisheries*, 27, 697–732.
Muhling, B. A., Lamkin, J. T. and Roff, M. A. (2010) Predicting the occurrence of Atlantic bluefin tuna (Thunnus thynnus) larvae in the northern Gulf of Mexico: building a classification model from archival data. *Fish. Oceanogr.*, 19, 526–539.

Muhling, B. A., Lee, S.-K., Lamkin, J. T. and Liu, Y. (2011) Predicting the effects of climate change on bluefin tuna (*Thunnus thynnus*) spawning habitat in the Gulf of Mexico. *ICES J. Mar. Sci.*, 68, 1051–1062.

Muhling, B. A., Liu, Y., Lee, S.-K., Lamkin, J. T., Roff, M. A., Muller-Karger, F. and Walter, J. F. III (2015) Potential impact of climate change on the Intra-Americas Sea: part 2. Implications for Atlantic bluefin tuna and skipjack tuna adult and larval habitats. *J. Mar. Syst.*, 148, 1–13.

Muller-Karger, F. E., Smith, J. P., Werner, S., Chen, R., Roff, M., Liu, Y., Muhling, B., Lindo-Aitchati, D. et al. (2015) Natural variability of surface oceanographic conditions in the offshore Gulf of Mexico. *Prog. Oceanogr.*, 134, 54–76.

Newell, S. E., Fawcett, S. E. and Ward, B. B. (2013) Depth distribution of ammonia oxidation rates and ammonia-oxidizer community composition in the Sargasso Sea. *Limnol. Oceanogr.*, 58, 1491–1500.

Oey, L., Ezer, T. and Lee, H. (2005) Loop current, rings and related circulation in the Gulf of Mexico: a review of numerical models and future challenges. *Geophys. Monogr.-AGU*, 161, 31–56.

Punerea, L. R., Williams, P. J. I., Azam, F. and Hobbie, J. E. (2007) The microbial loop. *Oceanography*, 20, 28–33.

Reglero, P., Santos, M., Balbín, R., Laiz-Carrión, R., Alvarez-Berastegui, D., Ciannelli, L., Jiménez, E. and Alemán, F. (2017) Environmental and biological characteristics of Atlantic bluefin tuna and albacore spawning habitats based on their egg distributions. *Deep-Sea Res. II*, 140, 105–116.

Rodríguez-Ezepeleta, N., Díaz-Areu, N., Walter, J. F. III, Richardson, D. E., Rooker, J. R., Notestad, L., Hanke, A. R., Franks, J. S. et al. (2019) Determining natal origin for improved management of Atlantic bluefin tuna. *Front. Ecol. Environ.*, 17, 439–444.

Rooker, J. R., Bremer, J. R. A., Block, B. A., Dewar, H., De Metrio, G., Corriero, A., Kraus, R. T., Prince, E. D. et al. (2007) Life history and stock structure of Atlantic bluefin tuna (*Thunnus thynnus*). *Res. Fish. Sci.*, 15, 265–310.

Rooker, J. R., Kitchens, L. L., Dance, M. A., Wells, R. D., Falterman, B. and Cornic, M. (2013) Spatial, temporal, and habitat-related variation in abundance of pelagic fishes in the Gulf of Mexico: potential implications of the Deepwater Horizon oil spill. *PLoS One*, 8, e76080.

Rooker, J. R., Simms, J. R., Wells, R. D., Holt, S. A., Holt, G. J., Graves, J. E. and Furey, N. B. (2012) Distribution and habitat associations of billfish and swordfish larvae across mesoscale features in the Gulf of Mexico. *PLoS One*, 7, e34180.

Rost, B., Zondervan, I. and Wolf-Gladrow, D. (2008) Sensitivity of phytoplankton to future changes in ocean carbonate chemistry: current knowledge, contradictions and research directions. *Mar. Ecol. Prog. Ser.*, 373, 227–237.

Saint-Béat, B., Vézina, A. F., Asmus, R., Asmus, H. and Niquil, N. (2013) The mean function provides robustness to linear inverse modelling flow estimation in food webs: a comparison of functions derived from statistics and ecological theories. *Ecol. Model.*, 258, 53–64.

Scanlan, D. J. and Post, A. E. (2008) Chapter 24 - aspects of marine cyanobacterial nitrogen physiology and connection to the nitrogen cycle. In Capone, D. G., Brouk, D. A., Mulholland, M. R. and Carpenter, E. J. (eds.), *Nitrogen in the Marine Environment*, 2nd edn, Academic Press, San Diego, pp. 1073–1095.

Schmitz, W., Biggs, D., Lugo-Fernandez, A., Oey, L. Y. and Sturges, W. (2005) A synopsis of the circulation in the Gulf of Mexico and on its continental margins. In *Circulation in the Gulf of Mexico: Observations and models*, pp. 11–29.

Selph, K. E., Landry, M. R., Taylor, A. G., Gutierrez-Rodriguez, A., Stukel, M. R., Wokuluk, J. and Pasulka, A. (2016) Phytoplankton production and taxon-specific growth rates in the Costa Rica Dome. *J. Plankton Res.*, 38, 199–215.

Selph, K. E., Swaethorps, R., Stukel, M. R., Kelly, T. B., Knapp, A. N., Fleming, K., Hernandez, T. and Landry, M. R. (this issue) Phytoplankton community composition and biomass in the oligotrophic Gulf of Mexico. *J. Plankton Res.*, doi: 10.1093/plankt/bfw006.

Sherr, E. B. and Sherr, B. F. (2016) Phagotrophic protist: central roles in microbial food webs. In Gilbert, P. M. and Kana, T. M. (eds.), *Aquatic Microbial Ecology and Biogeochemistry: A Dual Perspective*, Springer, Switzerland, pp. 3–12.

Shiozaki, T., Iijichi, M., Ijobe, K., Hashihama, F., Nakamura, K.-I., Ehama, M., Hayashizaki, K.-I., Takahashi, K. et al. (2016) Nitrification and its influence on biogeochemical cycles from the equatorial Pacific to the Arctic Ocean. *ISME J.*, 10, 2184–2197.

Shiroza, A., Malca, E., Lamkin, J., Gerard, T., Landry, M. R., Stukel, M. R., Laiz-Carrión, R. and Swaethorps, R. (this issue) Active prey selection in developing larvae of Atlantic bluefin tuna (*Thunnus thynnus*) in spawning grounds of the Gulf of Mexico. *J. Plankton Res.*

Shropshire, T. A., Morey, S. L., Chassaignet, E., Coles, V. J., Fiksen, O., Gerard, T., Malca, E., Laiz-Carrión, R. et al. (this issue) Trade-offs between risks of predation and starvation in larvae make the shelf break an optimal spawning location for Atlantic Bluefin tuna. *J. Plankton Res.*

Shropshire, T. A., Morey, S. L., Chassaignet, E. P., Bozec, A., Coles, V. J., Landry, M. R., Swaethorps, R., Zupile, G. et al. (2020) Quantifying spatiotemporal variability in zooplankton dynamics in the Gulf of Mexico with a physical-biogeochemical model. *Biogeosciences*, 17, 3385–3407.

Smith, J. M., Chavez, F. P. and Francis, C. A. (2014) Ammonium uptake by phytoplankton regulates nitrification in the sunlit ocean. *PLoS One*, 9, e108173.

Soetaert, K., Van den Meersche, K. and VAN Oevelen, D. (2009) limSolve: solving linear inverse models *R package version 1.5.4*. 

Stoecker, D. K., Hansen, P. J., Caron, D. A. and Mitra, A. (2017) Mixtography in the marine plankton. *Ann. Rev. Mar. Sci.*, 9, 311–335.

Strickland, J. D. and Parsons, T. R. (1972) *A Practical Handbook of Seawater Analysis*, Vol. 167, 2nd edn, Fisheries Research Board, Ottawa, Ontario, Canada.

Stukel, M. R., Décima, M. and Kelly, T. B. (2018a) A new approach for incorporating 15N isotopic data into linear inverse ecosystem models with Markov Chain Monte Carlo sampling. *PLoS One*, 13, e0199123.

Stukel, M. R., Décima, M., Landry, M. R. and Selph, K. E. (2018b) Nitrogen and isotope flows through the Costa Rica Dome upwelling ecosystem: the crucial mesozooplankton role in export flux. *Global Biogeochem. Cycles*, 32, 1815–1832.

Stukel, M. R., Kahrav, M., Benitez-Nelson, C. R., Decima, M., Goericke, R., Landry, M. R. and Ohman, M. D. (2015) Using Lagrangian-based process studies to test satellite algorithms of vertical carbon flux in the eastern North Pacific Ocean. *J. Geophys. Res. Oceans*, 120, 7208–7222.
Stukel, M. R., Kelly, T. B., Landry, M. R., Selph, K. E. and Swalethrop, R. (this issue) Sinking carbon, nitrogen, and pigment flux within and beneath the euphotic zone in the oligotrophic, open-ocean Gulf of Mexico. *J. Plankton Res.* doi: 10.1093/plankt/fbab001.

Stukel, M. R., Landry, M. R., Ohman, M. D., Goericke, R., Samo, T. and Benitez-Nelson, C. R. (2012) Do inverse ecosystem models accurately reconstruct plankton trophic flows? Comparing two solution methods using field data from the California Current. *J. Mar. Syst.*, **91**, 20–33.

Taylor, A. G. and Landry, M. R. (2018) Phytoplankton biomass and size structure across trophic gradients in the southern California Current and adjacent ocean ecosystems. *Mar. Ecol. Prog. Ser.*, **592**, 1–17.

Teo, S. L., Boustany, A., Dewar, H., Stokesbury, M. J., Weng, K. C., Beemer, S., Seitz, A. C., Farwell, C. J. et al. (2007) Annual migrations, diving behavior, and thermal biology of Atlantic bluefin tuna, *Thunnus thynnus*, on their Gulf of Mexico breeding grounds. *Mar. Biol.*, **151**, 1–18.

Tilley, J. D., Butler, C. M., Suárez-Morales, E., Franks, J. S., Hoffmayer, E. R., Gibson, D. P., Comyns, B. H., Ingram, G. W. Jr. et al. (2016) Feeding ecology of larval Atlantic bluefin tuna, *Thunnus thynnus*, from the central Gulf of Mexico. *Bull. Mar. Sci.*, **92**, 321–334.

Turner, J. T. (1986) Zooplankton feeding ecology: contents of fecal pellets of the cyclopoid copepods *Oncaea venusta*, *Corycaeus amazonicus*, *Oithona plumifera*, and *O. simplex* from the northern Gulf of Mexico. *Mar. Ecol.*, **7**, 289–302.

Uriarte, A., Johnstone, C., Laiz-Carrión, R., García, A., Llopiz, J. K., Shiroya, A., Quintanilla, J. M., Lozano-Peral, D. et al. (2019) Evidence of density-dependent cannibalism in the diet of wild Atlantic bluefin tuna larvae (*Thunnus thynnus*) of the Balearic Sea (NW-Mediterranean). *Fish. Res.*, **212**, 63–71.

Uye, S.-I. and Kayano, Y. (1994) Predatory feeding behavior of *T. thynnus* (Copepoda: Calanoida): life-stage differences and the predation impact on small planktonic crustaceans. *J. Crust. Biol.*, **14**, 473–483.

Van den Meersche, K., Soetaert, K. and Van Oevelen, D. (2009) `Sample()`: an R function for sampling linear inverse problems. *J. Stat. Softw., Code Snippets*, **30**, 1–15.

Van Oevelen, D., Van den Meersche, K., Meysman, F. J. R., Soetaert, K., Middelburg, J. J. and Vezina, A. F. (2010) Quantifying food web flows using linear inverse models. *Ecosystems*, **13**, 32–43.

Vézina, A. F. and Platt, T. (1988) Food web dynamics in the ocean. 1. Best estimates of flow networks using inverse methods. *Mar. Ecol. Prog. Ser.*, **42**, 269–287.

Walters, C., Martell, S. J., Christensen, V. and Mahmoudi, B. (2008) An Ecosim model for exploring Gulf of Mexico ecosystem management options: implications of including multistanza life-history models for policy predictions. *Bull. Mar. Sci.*, **83**, 251–271.

Wan, X. S., Sheng, H.-X., Dai, M., Zhang, Y., Shi, D., Trull, T. W., Zhu, Y., Lomas, M. W. et al. (2018) Ambient nitrate switches the ammonium consumption pathway in the eutrophic ocean. *Nat. Commun.*, **9**, 1–9.

Wells, R. D., Rooker, J. R., Quigg, A. and Wissel, B. (2017) Influence of mesoscale oceanographic features on pelagic food webs in the Gulf of Mexico. *Mar. Biol.*, **164**, 92.

Xu, M. N., Li, X., Shi, D., Zhang, Y., Dai, M., Huang, T., Gilbert, P. M. and Kao, S. J. (2019) Coupled effect of substrate and light on assimilation and oxidation of regenerated nitrogen in the eutrophic ocean. *Limnol. Oceanogr.*, **64**, 1270–1283.

Yool, A., Martin, A. P., Fernandez, C. and Clark, D. R. (2007) The significance of nitritification for oceanic new production. *Nature*, **447**, 999–1002.

Zehr, J. P. (2011) Nitrogen fixation by marine cyanobacteria. *Trends Microbiol.*, **19**, 162–173.