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April Cook  
*Nova Southeastern University*, acook1@nova.edu

Andrea Bernard  
*Nova Southeastern University*, andrbern@nova.edu

Kevin M. Boswell  
*Florida International University*

Heather Bracken-Grissom Dr.  
*Florida International University*, heather.bracken@gmail.com

Marta D'Elia  
*Florida International University*

*See next page for additional authors*

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Authors
Sergio DeRada
*Naval Research Laboratory at Stennis Space Center*

Cole Easson
*Nova Southeastern University; Middle Tennessee State University, ceasson@nova.edu*

David English
*University of South Florida*

Ron Eytan
*Texas A&M University at Galveston*

Tamara Frank
*Nova Southeastern University, tfrank1@nova.edu*

Chuanmin Hu
*University of South Florida*

Matt Johnston
*Nova Southeastern University, johnmatt@nova.edu*

Heather Judkins
*University of South Florida*

Chad Lembke
*University of South Florida*

Jose Lopez
*Nova Southeastern University, joslo@nova.edu*

Rosanna Milligan
*Nova Southeastern University, rboyle@nova.edu*

Jon A. Moore
*Florida Atlantic University*

Brad Penta
*Naval Research Laboratory at Stennis Space Center*

Nina Pruzinsky
*Nova Southeastern University, npruzinsky@gmail.com*

John A. Quinlan
*National Oceanic and Atmospheric Administration*

Travis M. Richards
*Texas A&M University, Galveston*
Isabel C. Romero  
*University of South Florida*

Mahmood S. Shivji  
*Nova Southeastern University, mahmood@nova.edu*

Michael Vecchione  
*National Museum of National History, Smithsonian Institute, Washington DC*

Max D. Weber  
*Texas A&M University, Galveston*

R.J. David Wells  
*Texas A&M University, Galveston*

Tracey Sutton  
*Nova Southeastern University, tsutton1@nova.edu*

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A Multidisciplinary Approach to Investigate Deep-Pelagic Ecosystem Dynamics in the Gulf of Mexico Following Deepwater Horizon

April B. Cook1*, Andrea M. Bernard2, Kevin M. Boswell2, Heather Bracken-Grissom2, Marta D’Elia3, Sergio deRada2, Cole G. Easson1,4, David English5, Ron I. Eytan6, Tamara Frank1, Chuanmin Hu5, Matthew W. Johnston1, Heather Judkins7, Chad Lembek5, Jose V. Lopez1, Rosanna J. Milligan1, Jon A. Moore5,6, Bradley Penta2, Nina M. Pruzinsky1, John A. Quinlan10, Travis M. Richards6, Isabel C. Romero5, Mahmood S. Shivji1, Michael Vecchione11, Max D. Weber4, R. J. David Wells6 and Tracey T. Sutton1

1 Guy Harvey Oceanographic Center, Halmos College of Arts and Sciences, Nova Southeastern University, Dania Beach, FL, United States, 2 Institute of Environment, Department of Biological Sciences, Florida International University – Biscayne Bay Campus, North Miami, FL, United States, 3 United States Naval Research Laboratory, Stennis Space Center, Washington DC, United States, 4 Department of Biology, Middle Tennessee State University, Murfreesboro, TN, United States, 5 College of Marine Science, University of South Florida, Tampa, FL, United States, 6 Department of Marine Biology, Texas A&M University at Galveston, Galveston, TX, United States, 7 Integrative Biology, University of South Florida St. Petersburg, St. Petersburg, FL, United States, 8 Wilkes Honors College, Florida Atlantic University, Jupiter, FL, United States, 9 Florida Atlantic University, Harbor Branch Oceanographic Institute, Fort Pierce, FL, United States, 10 Southeast Fisheries Science Center, National Oceanic and Atmospheric Administration, Miami, FL, United States, 11 NMFS National Systematics Laboratory, National Museum of Natural History, Washington, DC, United States

The pelagic Gulf of Mexico (GoM) is a complex system of dynamic physical oceanography (western boundary current, mesoscale eddies), high biological diversity, and community integration via diel vertical migration and lateral advection. Humans also heavily utilize this system, including its deep-sea components, for resource extraction, shipping, tourism, and other commercial activity. This utilization has had impacts, some with disastrous consequences. The Deepwater Horizon oil spill (DWHOS) occurred at a depth of ∼1500 m (Macondo wellhead), creating a persistent and toxic mixture of hydrocarbons and dispersant in the deep-pelagic (water column below 200 m depth) habitat. In order to assess the impacts of the DWHOS on this habitat, two large-scale research programs, described herein, were designed and executed. These programs, ONSAP and DEEPEND, aimed to quantitatively characterize the oceanic ecosystem of the northern GoM and to establish a time-series with which natural and anthropogenic changes could be detected. The approach was multi-disciplinary in nature and included in situ sampling, acoustic sensing, water column profiling and sampling, satellite remote sensing, AUV sensing, numerical modeling, genetic sequencing, and biogeochemical analyses. The synergy of these methodologies has provided new and unprecedented perspectives of an oceanic ecosystem with respect to composition, connectivity, drivers, and variability.

Keywords: micronekton, epipelagic, mesopelagic, bathypelagic, sampling, hydrography, acoustics, ecosystem structure
INTRODUCTION

Of the ecotypes of the Gulf of Mexico (GoM) affected by the Deepwater Horizon oil spill (DWHOS), the open-ocean pelagic ecotype was by far the largest. The spill began on April 20, 2010, about 66 km off the coast of Louisiana, at a depth ~1,500 m and continued for 87 days (Beyer et al., 2016). Some percentage of oil, other hydrocarbons, and injected dispersant reached the sea surface and seabed, whereas 100% occurred within the deep-pelagic domain (200 m depth to just above the seafloor). During the summer of 2010, a continuous plume of oil over 35 km in length was discovered at approximately 1,100 m depth (Camilli et al., 2010). This plume persisted for several months, prompting concern about the effects of the DWHOS on the meso- and bathypelagic (200–1,000 and >1,000 m depths, respectively; deep-pelagic, cumulatively) faunas. Deep-pelagic animals are known to vertically migrate to shallow, epipelagic (0–200 m depth) waters at night to feed (Sutton et al., 2020), a process which ostensibly increases exposure throughout the water column and connects the shallower and deeper parts of the oceanic GoM.

Gaining insight and understanding of pelagic ecosystems over time requires a multidisciplinary approach, given their complex physical (4-D, Lagrangian), biological, and ethological (vertically migratory) nature. Here we describe the sampling, sensing, and analysis methods of two major research programs, both aimed at characterizing effects, or potential effects, of the DWHOS on the epi-, meso-, and bathypelagic faunas of the northern GoM. The first program, ONSAP (Offshore Neptons Sampling and Analysis Program), was supported by the National Oceanic and Atmospheric Administration (NOAA) as part of the DWHOS Natural Resource Damage Assessment (NRDA) conducted in 2010–2015. This program encompassed in situ net sampling, water column profiling, and active acoustic sensing (Supplementary Tables 1,2) to address the question, “What could have been affected by the DWHOS in the deep-pelagic GoM?” The dearth/lack of pre-DWHOS data and the needs of the NRDA process required this initial approach. The second program was DEEPEND (DEEp Pelagic Nektos Dynamics), a research consortium supported by The Gulf of Mexico Research Initiative (GoMRI) from 2015 to 2020. This program, which added satellite remote sensing, AUV sensing, physical oceanographic numerical modeling, pelagic microbial ecology, genetic analysis, biogeochemical analysis, and trophic ecology (Supplementary Tables 3,4) was both a continuation and evolution of ONSAP. The additions to DEEPEND, when integrated with foundational information from ONSAP, addressed the questions, “What are the natural drivers of pelagic ecosystem structure in the GoM?” and “Did pelagic faunal abundance variations after DWHOS exceed this ‘natural envelope?’”

SURVEY APPROACH

The overall goal of the initial ONSAP project was to survey and quantify the deep-pelagic life forms living within or traveling through the area of the GoM affected by the oil spill (Frank et al., 2020; Sutton et al., 2020). Of interest was the water column fauna at the mesopelagic/bathypelagic interface, the depth stratum containing the deep hydrocarbon plume. The plume was discovered in areas surrounding the Macondo wellhead where the spill originated. To accomplish this goal, a multi-disciplinary approach was used. Acoustic profiles were collected to synoptically quantify organisms distributed throughout the water column. These can easily be repeated for comparisons across space and time. While a very useful tool, acoustics cannot discern between individual species nor can it detect many deep-pelagic organisms without swim bladders or air pockets. Discrete-depth midwater trawling was conducted to identify and quantify the organisms collected during both day and night to account for vertical migration. These results also help to ground truth the acoustic profiles. Environmental factors such as temperature, conductivity, and dissolved oxygen were collected from both trawl-mounted sensors and CTD rosette profiling from 0 to 1500 m depth. Details of survey design and methodologies are described further below.

When planning the DEEPEND program, several additional components were added to the survey approach to fill in data gaps and to expand on research objectives. A remote sensing and satellite imagery component was added to identify mesoscale oceanographic and riverine discharge features to inform planning and execution of field work (e.g., Androulidakis et al., 2019). A glider was deployed to collect oceanographic data that were assimilated in the ocean model which was used to establish DEEPEND cruise tracks. This multidisciplinary methodology, integrating physical oceanographic modeling, satellite observation, and in situ sensing, provided the spatiotemporal habitat context by which pelagic faunal composition, abundance and distribution were analyzed (i.e., biophysical coupling; Meinert et al., 2020; Milligan and Sutton, 2020; Pruzinsky et al., 2020).

A biogeochemical component was added to directly measure the amount of petrogenic contamination in animal organs, muscle tissues, and eggs using polycyclic aromatic hydrocarbons (PAHs) as a proxy (Romero et al., 2018, 2020). A molecular taxonomy component (DNA barcoding; Hebert et al., 2003) was added to help identify damaged, cryptic, and juvenile specimens where morphological characters do not yet exist or could not differentiate between species (e.g., Moore et al., 2020). The gene sequences analyzed are well established as robust markers for species identification of marine fishes (Ward et al., 2009) and invertebrates (Mantelatto et al., 2018). A population genomics component was added (double-digest Restriction Associated DNA sequencing; ddRADSeq) to study genetic diversity and connectivity of the GoM and adjacent water deep-pelagic fauna (Timm et al., 2020b). Genetic diversity and connectivity can be used as proxies to measure population health and resilience (Oliver et al., 2015). Over a time-series, these measures can show how diversity is maintained and restored in the face of anthropogenic and/or natural disasters. A trophic ecology component, using Stable isotope analysis (SIA), was added to identify feeding relationships among taxa, estimate trophic positions, and delineate energy flow (Richards et al., 2020). Understanding the flow of energy through this deep-sea
ecosystem is essential to be able to identify linkages which may be vulnerable to disasters such as an oil spill. A microbial ecology component was added to help characterize pelagic habitats (along with environmental and ocean modeling data) and to investigate the dynamics of diel vertical migration using acoustic backscatter data and eDNA sampling (Easson et al., 2020).

The time-series aspect of these two programs provides information on the patterns of abundance and distribution of the pelagic fauna, the concentration of PAHs therein, and the pattern of genetic diversity following a major marine disaster. Information such as this also provides a basis against which to compare hindcast-derived abundance estimates using proxies for data that did not previously exist (e.g., larval and adult deep-pelagic fish abundance relationships; the former data were collected prior to the DWHOS, while the latter were not). The multidisciplinary nature of these two programs facilitates an ecosystem-based approach to guide interpretations of assemblage-level data. For example, using ddRADseq in combination with physical oceanographic modeling provided evidence that the Loop Current could be facilitating genetic connectivity in pelagic shrimps, with its concomitant implications for the recovery and resilience of a species (Timm et al., 2020a). In another example, microbial assemblages were characterized using abiotic and biotic data collected via CTD sensing and their dynamics interpreted using MODIS satellite imagery (Easson and Lopez, 2019). In summary, results derived from each component were valuable in their own right, but each also added necessary information for other working groups.

**Transect Design**

During ONSAP field operations, a subset of the Southeast Area Monitoring and Assessment Program (SEAMAP; Eldridge, 1988) stations surrounding the DWHOS site was sampled (Figure 1), with original station nomenclature maintained. Sampling the entire 46-station grid took approximately 3 months, requiring that sampling be divided into several legs for resupply and personnel changes. This necessity dictated that sampling transects be arranged by logistics (time to station, weather, and personnel availability) in lieu of oceanographic and/or ecological considerations. Sampling, acoustic sensing, and water column profiling were conducted twice at each station (day and night). Sampling of the entire grid was conducted three times over a 9-month period, with each station being occupied either three (most stations) or two times over the course of ONSAP (Figure 1).

Due to time constraints, only a portion of the stations sampled during ONSAP were sampled during individual DEEPEND cruises, each of which lasted approximately 15 days. DEEPEND cruise tracks were designed to transect as many water masses (Common Water and Loop Current, sensu Johnston et al., 2019; Boswell et al., 2020) and mesoscale features (eddies, Mississippi River plumes) as possible during each cruise in order to model faunal assemblage structure, abundance, and distribution as a function of biophysical drivers. Because the location, intensity, and persistence of the GoM’s salient oceanographic features are constantly in flux, we considered both hindcasts and forecasts of hydrographic conditions from the United States Naval Research Laboratory’s Hybrid Coordinate Ocean Model (HYCOM; see section “Hybrid Coordinate Ocean Model”) along with satellite imagery (see section “Remote sensing/chlorophyll”) in selecting the location and timing of DEEPEND sampling stations from the original ONSAP sampling grid (Figure 2). This “directed sampling” approach allowed statistical analysis of population and assemblage variability as a function of environmental variability,
a methodology applied to both DEEPEND and the preceding ONSAP data.

Hybrid Coordinate Ocean Model

Hybrid Coordinate Ocean Model (HYCOM), implemented at 1/25° horizontal-resolution for the GoM (18 to 31° N., 77 to 98° W.), was run in “real-time” in the weeks before and during the DEEPEND cruises (DP01 through DP06, Supplementary Table 3), providing surface and sub-surface predictions through the pelagic ocean. In order to sample important features, pre-determined cruise tracks and stations were adjusted depending on proximity to these predicted mesoscale oceanographic features (e.g., eddies and fronts). Model predictions were delivered in the form of “first-look” visualizations via web portals. The model was configured with a 32-layer hybrid ($\sigma$/z/$\rho$) time-variant vertical structure, which was post-processed into a time-invariant, 50-level, z-vertical structure for end-user dissemination. In this configuration, the model assimilated daily observations using 3-D variational data-assimilation, received (initial) boundary information from the Global Ocean Forecasting System (Metzger et al., 2014), and was forced by 3-h momentum and heat fluxes from the Navy Global Environmental Model (NAVGEM). Tidal boundary conditions for water level and barotropic velocity were provided by the global Ocean Tide Inverse Solution (OTIS), and rivers were implemented as a “precipitation bogus,” specified by a monthly climatological database. Further information and detailed documentation about HYCOM can be found at hycom.org.

For the DEEPEND cruise campaigns, the model provided up to 120 h of forecasts, at 3-h frequency, of the 3-D oceanic physical environment (sea surface height, ocean currents, temperature, and salinity). The HYCOM model was initialized on January 1, 2015 and ran continuously through December 31, 2018. Its outputs for 2015 (Cruises DP01 and DP02), 2016 (DP03 and DP04), 2017 (DP05), and 2018 (DP06) were deposited in the Gulf of Mexico Research Initiative Information and Data Cooperative (GRIIDC; Supplementary Table 5).

Remote Sensing/Chlorophyll

In the GoM, the location and intensity of mesoscale features can change dramatically in a few days, requiring that ocean color imagery be used to determine the precise location of surface features (e.g., Figure 3), especially the location of Mississippi River plumes and the Loop Current. While the location of surface fronts may not coincide with water mass boundaries at bathypelagic depths, the material and energetic relationships between euphotic and deeper waters were considerations when planning DEEPEND sampling transects.

Ocean color satellite images from the Moderate Resolution Imaging Spectroradiometer (MODIS) satellite were processed at the University of South Florida (USF) Optical Oceanography Laboratory through a Virtual Antenna System (VAS; Hu et al., 2013). Ocean color imagery is based on spectral reflectance of the surface ocean, which depends on the absorption and scattering of sunlight in near surface waters and therefore carries information on surface water constituents such as phytoplankton chlorophyll and colored dissolved organic matter (CDOM). The chlorophyll imagery was derived using NASA standard algorithms to remove atmospheric effects and convert surface reflectance to chlorophyll (Hu et al., 2012). Clouds, sunlight, and limited viewing angles can reduce the area of reliable ocean color satellite data. Thus, multi-day composites of MODIS ocean color imagery were created.
FIGURE 3 | Examples of MODIS ocean-color composite images created for the DEEPEND study region (26–30°N, 85–91°W) during DP06 (A–C for July 22, 25, and 30, 2018, respectively). Imagery from several days was combined to emphasize recent surface feature locations. In agreement with HYCOM model predictions, features in the left portion of these images tended to move toward the southwest at more than 20 km per day, while features in the lower central portion of the images were influenced by the Loop Current and moved to the east-southeast at about 50 km per day.

to decrease the fraction of the cruise area imagery that would otherwise have been masked or obscured. Due to the movement of surface fronts (sometimes more than 20 km over several days; Figure 3), satellite images from several days (up to a week) were combined such that the locations of ocean color features in the most recent images would be emphasized. The composites were sent to the Chief Scientist aboard the ship and the supervisor of glider operations so that transects could be adjusted to avoid or examine particular features. While sea surface temperature (SST) imagery was also examined, solar heating of the surface waters diminished the practical use of SST imagery to monitor the changing location of mesoscale features during the late-season (August) cruises.

FIELD SAMPLING AND WATER-COLUMN SENSING

Net Sampling
The vertical distribution of micronekton in the water column from the surface to 1,500 m was quantified by sampling discrete-depth intervals using a Multiple Opening Closing Net and Environmental Sensing System (Wiebe et al., 1985; Sutton et al., 2010) with an effective mouth area of 10 m² (referred to as MOC10 hereafter; Figure 4) when towed at a 45° angle. The MOC10 (3.41 × 4.69 m mouth opening) was equipped with six nets of 3-mm uniform mesh which were opened and closed at specific depth intervals on command from the ship through conducting trawl wire. This procedure yielded one oblique sample from the surface to the maximum depth sampled (net 0) and five discrete-depth samples (nets 1–5, Table 1). The rationale for these depth intervals, following Sutton (2013) and listed from deep to shallow, was: Net (1) sample the bathypelagic fauna living below the deep hydrocarbon/dispersant plume [i.e., below 1,200 m depth; Net (2)] sample the bathypelagic fauna within the stratum occupied by the deep plume (1,000 to 1,200 m depth); Net (3) sample the deep mesopelagic fauna (600 to 1,000 m, the daytime depths of occurrence of most vertically migrating taxa and persistent depth of occurrence of non-migrating mesopelagic taxa); Net (4) sample the fauna within the upper mesopelagic zone (200–600 m, daytime depths of shallow mesopelagic migrants and nighttime depth of weakly migrating taxa); and Net (5) sample the fauna of the epipelagic zone (0–200 m, the nighttime depths of most vertically migrating mesopelagic taxa and persistent depth of non-migrating surface fauna). Trawling was conducted twice at each station, centered at solar noon and midnight, to quantify diel vertical migration. Instruments were mounted to the trawl frame to measure depth, temperature, and salinity [conductivity], as well as the mouth angle of the net through the water. The volume of water filtered by each MOC10 net was measured by a Tsurumi-Sikie-Kosakusho Co., Ltd. flowmeter mounted on the MOC10 frame (adjusted for towing angle) facing directly into the flow of water. The trawl was towed at 1.5–2.5 knots and retrieved at a rate of 5 m min⁻¹. The total volume of water filtered varied by net depth stratum and ranged from 6,500 to 70,000 m³ (Supplementary Tables 1,3). During the ONSAP, MOC10 sampling on the M/V Meg Skanski occurred almost continuously from January to September 2011 (Figure 1). In total, 241 trawl deployments were conducted at 58 stations, yielding 936 quantitative, discrete-depth samples (Supplementary Table 1). During DEEPEND, sampling occurred in either May or August (the height of dry and wet seasons in the GoM, respectively) aboard the R/V Point Sur from 2015–2018 (Figure 2). In total, 122 trawl deployments were conducted at 24 stations, yielding 470 quantitative, discrete-depth samples (Supplementary Table 3). A quantitative sample was defined as having been collected within the depth bins detailed in Table 1 as well as having a valid measurement of the volume of seawater filtered by that net.

The MOC10 system was chosen for its discrete-depth sampling capability (versus non-closing nets), a key consideration for quantifying the abundance of vertically migrating taxa. The 10 m² MOCNESS was chosen over a 1 m² MOCNESS, as the former selects for micronekton (2–20 cm body
length) as opposed to plankton (Wiebe and Benfield, 2003). The fixed mouth area and the integrated flow meter allow for precise quantitative sampling, a prerequisite for time-series analysis. Lastly, the MOC10 can be deployed from an intermediate (regional-class) research size vessel with a single aft winch and conducting cable. Larger, dual-warp trawls require larger, fisheries-capable research vessels whose expense and availability are often prohibitive. That said, there are caveats with any sampling system, including the MOC10. The mouth and mesh size of rectangular midwater trawls limit the speed at which they can be towed, allowing for net avoidance by larger, more mobile taxa (Pearcy, 1983; Kaartvedt et al., 2012; Kwong et al., 2018). The multiple opening/closing nets may also be prone to “net contamination,” where animals from non-target strata can squeeze through the mouth bars of a closed net. We found the latter to be infrequent, and readily recognizable when it occurred. Taking all these factors into consideration, the MOC10 was determined to be the best gear to sample deep-pelagic micronekton/nekton in the GoM in a quantitative fashion in order to accomplish the goals of ONSAP and DEEPEND.

Midwater Trawl Sample Processing

**ONSAP**

After each MOC10 deployment and retrieval, individual nets were washed down with seawater and the contents of each codend were rinsed into separate numbered containers. Specimens from each codend sample were then concentrated with a sieve and placed into labeled collection jars and preserved. Larger specimens were curated separately in labeled jars. Nets 1–5 were preserved with buffered 10% formalin: seawater, while net 0’s were preserved in 95% non-denatured ethanol (EtOH) for genetic analyses. When the size or amount of gelatinous zooplankton exceeded storage capacity, individuals of each taxon were sorted into a graduated beaker, the volume and weight recorded, and the animals discarded at sea. The remaining gelatinous individuals were preserved with the rest of the catch. No sub-sampling occurred during at-sea processing. After each cruise, the samples were transported to Nova Southeastern University’s (NSU) Oceanic Ecology Laboratory, where they were sorted by major taxon, and distributed to the appropriate laboratory for species-level identifications by experts within each major taxonomic group. Specimens were then enumerated, weighed, and measured. Data were entered and stored as described in section “Biotic databases.”

**DEEPEND**

Midwater trawl sample processing at sea was more involved during DEEPEND than ONSAP (i.e., there was extensive subsampling for genetic and biogeochemical analyses), requiring

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**TABLE 1** | Discrete-depth ranges targeted for sampling via MOC10 during both ONSAP and DEEPEND cruises.

| Net number | Depth range (m) |
|------------|----------------|
| Net 0      | 0–1500         |
| Net 1      | 1500–1200      |
| Net 2      | 1200–1000      |
| Net 3      | 1000–600       |
| Net 4      | 600–200        |
| Net 5      | 200–0          |

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Written informed consent was obtained from the individual for the publication of any potentially identifiable images included in this article.
additional handling and data management procedures. Upon retrieval of the MOC10, catches from each net were rinsed into separate containers and kept in cold (4°C) seawater during shipboard processing. This step is extremely important; deep-pelagic animals tend to degrade quickly at room temperature due to their chemical composition. Samples were sorted separately and sequentially to avoid mixing specimens from different collection nets (i.e., depth strata). While each sample was being processed, all others were stored in a refrigerator at 4°C.

Fishes, crustaceans, and cephalopods were rough-sorted by higher taxon and then identified to lower taxonomic levels (usually species) by onboard taxonomic specialists. Identified animals were counted and weighed to the nearest gram on a motion-compensating scale in batches per lowest taxonomic unit. Up to 25 specimens of each taxonomic unit were measured to the nearest millimeter per sample. All data were entered directly into the DEEPEND Nekton Database at sea (see section “Biotic databases” for biotic database description). Animals that were not subsampled for other analyses (described below) were preserved and brought back to the lab. Animals that were not identified to species at sea were examined in the lab post-cruise for further identification.

**Genetics sub-sampling**

As part of DEEPEND’s initiative to catalog the species diversity of the deep-pelagic waters of the GoM, a ~650 bp segment of the mitochondrial Cytochrome c oxidase I (COI) gene and/or a ~550 bp segment of the large mitochondrial subunit 16s or 28s genes were sequenced from a subset of fishes, crustaceans, and cephalopod species. This method, “DNA barcoding,” allows researchers to use a partial DNA sequence to identify an organism to the species level. It is particularly helpful in cases where the specimen represents an undescribed, “cryptic” species, an undescribed early-life-history form, or when definitive morphological characters are not available (e.g., male anglerfishes, trawl-damaged specimens, etc.).

Tissue samples for genetic barcoding were taken from up to 10 specimens of each fish species and up to five specimens of each crustacean and cephalopod species collected during DEEPEND cruises. Initial morphological identifications were conducted at sea, and subsequently checked by COI, 16s, and/or 28s barcoding (depending on taxonomic group). Tissues were preserved in either 95% non-denatured EtOH or RNALater. In addition to these samples, up to 50 tissue samples per cruise were collected for temporal population genomics studies (ddRADseq; Peterson et al., 2012) of eight fish species and six crustacean species (Timm et al., 2020b, Supplementary Table 6). Additionally, tissue samples from three species of cephalopods were used to compare the genetic connectivity of each species between the GoM (Supplementary Table 6) and the Bear Seamount region of the northern Atlantic Ocean (Timm et al., 2020a). In all cases, paired plastic identification tags were kept with each tissue sample and the corresponding individual voucher specimen to maintain data integrity before, during, and after barcoding procedures.

These methods have proven useful for the study of diversity, health and resilienc in the GoM (Judkins et al., 2016, 2020; Timm et al., 2019, 2020a,b). A major challenge of the DNA barcoding method was matching the genetic sequences with those already submitted by other researchers in the Barcode of Life Database. There were many instances where either one genetic sequence had more than one species name assigned to it, multiple sequences were attributed to the same species, or the species name listed in the database conflicted with the identification made by DEEPEND taxonomic experts.

**Stable isotope analysis sub-sampling**

A thorough understanding of deep-pelagic ecosystems requires detailed knowledge of food webs including descriptions of feeding relationships among taxa, estimations of trophic position, and delineations of energy flow. Food webs have traditionally been examined through gut content analysis (GCA) which can require thousands of samples, a high level of taxonomic expertise, and is best suited for organisms that ingest prey whole. SIA is a powerful complement to GCA, as it is not as dependent on taxonomic expertise, can be applied to a range of taxa regardless of feeding mode, and can be conducted with fewer samples. However, interpretation of SIA data can be difficult due to significant spatiotemporal variation in the isotopic signatures of primary producers (isotopic baseline) which can be conserved in higher-order consumers resulting in misinterpretation of feeding relationships and incorrect trophic position estimates. Amino acid compound-specific isotope analysis (AA-CSIA) is a more refined technique that can help distinguish between variation in consumer isotopic signatures caused by changes in the isotopic baseline and changes in the diets and feeding habits of consumers (Popp et al., 2007). The method uses “source” amino acids that accurately reflect the isotopic values of primary producers and “trophic” amino acids that can be used as indicators of change in consumer feeding and diet (McClelland and Montoya, 2002; Chikaraishi et al., 2009). Given the advantages of SIA, and because several high quality GCA datasets currently exist for deep-pelagic assemblages in the GoM (Flock and Hopkins, 1992; Hopkins et al., 1996), SIA and AA-CSIA were employed to provide a complementary description of the trophic structure of deep-pelagic assemblages in the GoM (Richards et al., 2019, 2020). To better inform the study design, catch data from MOC10 sampling and prior GCA investigations in the GoM were leveraged to identify numerically dominant species that represent important energy vectors connecting primary and secondary production with higher-order consumers. These species encompassed an array of migratory strategies (synchronous vertical migrants, asynchronou vertical migrants, and non-migrants) and feeding modes (Supplementary Table 6). Additionally, data from the HYCOM and MODIS were used to ensure specimens were collected from salient mesoscale features (e.g., cyclonic and anticyclonic eddies, Mississippi River plume), providing as complete a representation of deep-pelagic trophic structure as possible.

Following collection through MOC10 sampling, specimens for SIA and AA-CSIA were identified and enumerated at sea, with specimens selected for bulk SIA frozen whole at −20°C, while specimens selected for AA-CSIA were frozen whole in liquid nitrogen before transport to Texas A&M University at Galveston.
SIA and AA-CSIA specimens were kept in long-term storage at −20 and −80°C, respectively. Muscle tissue used for SIA and AA-CSIA was dissected from the lateral musculature of fishes, from the anterior portion of the mantle from cephalopods, and from the dorsal portion of the abdomen in decapod crustaceans. Samples were then rinsed with deionized water to remove trace carbonates, freeze dried, and homogenized using mortar and pestle. Information on remaining procedures during SIA and AA-CSIA can be found in Richards et al. (2019) and Richards et al. (2020).

Polycyclic aromatic hydrocarbon analysis sub-sampling
Polycyclic aromatic hydrocarbon (PAH) analyses were conducted on GoM deep-pelagic micronekton to determine the extent and persistence of DWHOS-derived oil contamination. Smaller fishes (<15 mm), cephalopods, and shrimp samples collected for PAH analysis were stored whole in pre-combusted (450°C for 4 h) glass vials and frozen in a −20°C freezer. The larger fishes (>15 mm) were dissected at sea to remove internal organs (liver, stomach, heart, and intestines), gills, muscle tissue, and eggs (if present). Each dissected tissue was stored separately and frozen. All samples were transported on dry ice to USF (Supplementary Table 6). Whole-body samples were dissected at USF to collect internal organs, muscle tissue, and eggs (if present) from fishes and shrimps, and mantle tissue and eggs (if present) from cephalopods. For a complete description of methods and findings for fishes see Romero et al. (2018) and for cephalopods see Romero et al. (2020).

In situ Sensing
Abiotic Sensing
MOC10 sensors
The MOC10 was outfitted with pressure (depth), temperature, and conductivity (salinity) sensors, which were calibrated annually. The sensors recorded a reading once every four seconds during the entire tow.

CTD sensors
ONSAP. A Sea-Bird SBE 19 plus V2 CTD profiling package was deployed at each station to at least 1,500 m (when the bottom depth was greater than 1600 m). Stations with water depths less than 1,600 m were profiled to full water column depth within 100 m of the bottom (Supplementary Table 2). The CTD was mounted to a 12-Niskin bottle (12-L) rosette and equipped with a dissolved oxygen sensor (Sea-Bird SBE-43), two fluorometers (WET Labs CDOM and WET Labs ECO-AFL/FL), and a turbidity meter (WET Labs ECO-NTU). The CTD data were processed following the DWH-NRDA CTD processing protocol. Calibrated data from each sensor were averaged in 1-m bins within Sea-Bird’s SBE Processing software. For all deployments, only data from the downcasts were used in characterizing the water column structure.

DEEPEND. A 12-Niskin bottle (12-L) rosette with CTD was deployed from the R/V Point Sur at DEEPEND stations (Figure 2), usually to depths greater than 1,000 m. There were 106 CTD profiles collected during the DEEPEND cruises (Supplementary Table 4). The Sea-Bird 911plus CTD on the sampling rosette combined measurements of conductivity, temperature, and pressure, with additional sensors connected to the CTD on a per-cruise basis. These sensors included one or more dissolved oxygen sensors (Sea-Bird SBE-43), a transmissometer (WET Labs C-Star), and fluorometers (WET Labs ECO CDOM, ECO chlorophyll a, or Chelsea UV Aquatracka). The number and type of sensors varied between cruises, but information from the dissolved oxygen sensor and chlorophyll fluorometer was available at almost all of the DEEPEND stations. The CTD data were post-processed using Sea-Bird’s SBE Data Processing software, which converted the data to engineering units as well as computed salinity and dissolved oxygen concentrations. To increase the consistency of in situ chlorophyll a fluorometer results between the DEEPEND cruises, the measurements of water-sample chlorophyll a and CDOM absorbance were used to scale the CTD’s in situ fluorometer measurements. The measurements were binned (using median values) into 1-m depth intervals. Both the raw and binned CTD data for each DEEPEND cruise are available through the GRIIDC data repository (Supplementary Table 5).

Slocum glider sensing
During select DEEPEND cruises, a 1000-m depth-rated Slocum Electric Glider (Figure 5) was used to characterize the upper 400 m (on average) of the GoM water column. The glider was equipped with a Seabird SBE41CP CTD, two WET Labs fluorometers, two Satlantic radiometers, and an Aanderaa dissolved oxygen sensor. The fluorometers were equipped to sample for chlorophyll, CDOM, backscatter at 660 and 880 nm, and turbidity. All sensors sampled at 0.25 Hz. The radiance and irradiance sensors sampled at four wavelengths: ~412, 443, 556, and 683 nm. The glider transited vertically between 2 m and max depth (ranging from 400–800 m) at ~0.1 m/s which resulted in a vertical sample resolution of ~0.4 m. The measurements were taken at various depths and included conductivity, temperature, depth, chlorophyll fluorescence, gelbstoff fluorescence, dissolved oxygen, and light field measurements. While the glider was...
deployed at sea, it surfaced and communicated to an onshore control station at predetermined intervals, typically every 3 h. Launch, transit progress, and recovery of the glider position were planned and conducted, in part, by utilizing the HYCOM to provide context of the predicted current structure of the Loop Current and eddies. Model input for mission planning allowed glider adaptive sampling of features and assisted piloting to avoid unfavorable currents wherever possible. Once recovered, the complete measurement suite was downloaded from the vehicle, processed, and made available through USF and national data archives, including GRIIDC. The glider temperature and salinity data were assimilated into the HYCOM to assist in analyses of subsurface water characteristics and validation of ocean models, which was used to support the DEEPEND cruises.

**Biotic Sensing**

**Acoustic backscatter**

Two different vessels collected hydroacoustic data during the ONSAP and DEEPEND sampling programs. Simrad split-beam echosounder systems (EK60 and EK80) were used on both vessels; however, the transmitted frequencies varied according to vessel (Table 2). Transducers were mounted from a pole mount on both the M/V *Meg Skansi* and R/V *Point Sur*. While both vessels transmitted at 18 and 38 kHz, the higher frequencies were intermittently available. During each survey, the echosounder system was calibrated following the standard sphere method described by Demer et al. (2015). Measured gains and offsets derived from the Simrad lobe calibration program were recorded and input into the data analysis process. The echosounders were operated consistently among the surveys to ensure comparability over time. Multifrequency backscatter data were recorded simultaneously from each transceiver with a ping rate set to 0.2 Hz.

Analyses of raw backscatter data were processed in Echoview (PTY. Ltd.). Data were manually scrutinized for interference, noise, and other artifacts, and data processing routines were applied to reduce the effects of these on the processed data following methods outlined in D’Elia et al. (2016). Specifically, data were corrected for the effects of attenuation due to propagation losses and absorption. Intermittent noise spikes and transient noise were removed with Echoview (Ryan et al., 2015). Following that, a background noise removal process was applied (De Robertis and Higginbottom, 2007). Data were re-sampled at 500 m × 5 m (horizontal by vertical) elementary distance sampling units (EDSU) to generate analysis cells in which the echo integral was derived for each transect. Multifrequency comparisons were drawn to examine water-column backscatter between 18 and 38 kHz (D’Elia et al., 2016; Boswell et al., 2020).

The main limitation or bias associated with this method is attributing the backscatter to specific taxa. Using “ground-truth” data from direct biological sampling (i.e., nets) to interpret the backscatter patterns is the ideal methodology and was employed in both ONSAP and DEEPEND. Another potential limitation when using acoustic data across wide depth ranges is the potential effect of resonance when vertically migrating animals with gas bladders change depth. This effect occurs because backscattering intensity changes as a function of the surface area of a gas bladder. Therefore, it is important to interpret acoustic data carefully to account for this possibility (Davison et al., 2015; Proud et al., 2019).

**Water Sampling**

During DEEPEND cruises, CTD profiles were used to identify the depths of four “features of interest” at each station where water samples were collected. The features of interest included the surface layer, the chlorophyll-maximum layer, the oxygen-minimum zone, and the maximum trawl depth at each station. The depths of the chlorophyll maximum and oxygen-minimum zone, which varied by station, were determined visually during the CTD downcast using real-time data collected by fluorometer and oxygen sensors. Once the station-specific water collection depths were determined, three Niskin bottles were fired at each of the four targeted depths during the CTD upcast, yielding 36 L per depth.

**Optical Absorption of Particulate and Dissolved Material and Determination of Chlorophyll a Concentration**

The absorption of light within the surface waters is a dominant factor in determining ocean color. Measurements of the optical absorption spectra for particulates and dissolved material in water samples were collected because they provide information for the validation of ocean color imagery (e.g., section “Remote sensing/chlorophyll”), information about the pigment composition of phytoplankton in a water sample, and a measurement of the concentration of chlorophyll and dissolved material in that water sample. Samples from waters near the sea surface (<5 m depth) and the chlorophyll maximum were used not only to estimate the chlorophyll a concentration, but also to separate the water’s optical absorption spectra into contributions from the sample’s particulate material, $a_p(λ)$, detrital material, $a_d(λ)$, and CDOM, $a_{CDOM}(λ)$. Shortly after collection, samples were filtered through a glass fiber filter to separate the particulate constituents from a water sample, with additional filtration to partition the dissolved material. Both the filter pad and filtrate was then stored for additional processing and analysis ashore.

The chlorophyll a and CDOM measurements from the water samples were used to standardize the *in situ* fluorometry values as mentioned in section “DEEPEND”. While the variability in the relationship between *in situ* fluorometric measurements and

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**Table 2** | Echosounder system properties used during multi-vessel studies in the GoM.

| Program and date range | Vessel                  | Frequency (kHz) | Pulse duration [kHz] | Pulse rate (pps) |
|------------------------|-------------------------|-----------------|----------------------|------------------|
| ONSAP, 2011            | M/V *Meg Skansi*        | 18, 38, 70, 120 | 4 ms [18, 38]; 1 ms [120] | 0.2             |
| DEEPEND, 2015–2018     | R/V *Point Sur*         | 18, 38, 70, 120 | 4 ms [18, 38]; 1 ms [70,120] | 0.2             |
chlorophyll \( a \) concentrations and the necessity of validation
measurements is often acknowledged. The normalization of the
fluorometric data is frequently omitted in presentations of the
in situ fluorometry (Roessler et al., 2017). Not only was the
in situ environment during the DEEPEND cruises different
than those used for factory fluorometer calibrations, different
fluorometers were used with the CTD on different cruises. The
optical absorption information from the filter pads was used to
improve the consistency of in situ fluorometric measurements
between different casts and cruises.

The determination of optical absorption using this filter pad
method requires several liters of sample water for the clear
waters found throughout much of the DEEPEND sampling
region. This relatively large volume of sample water, and the
time and effort needed to filter and process that water, limits
the number of water sampling depths that can be practically
collected from a CTD cast. Though the samples were intended
to capture representative waters from CTD profile features (e.g.,
chlorophyll maximum depth intervals), unsampled variations in
planktonic composition and optical properties may occur within
features. Chlorophyll \( a \) concentration and optical absorption
spectra data are available through the GRIIDC data repository
(Supplementary Table 5).

**Microbial Community Characterization**
Seawater microbial sampling followed routine methods, as
described in Easson and Lopez (2019), to capture the dynamics
of microbial plankton communities in relation to a host of
biotic and abiotic factors. Briefly, seawater samples from all
four targeted depths (surface, chlorophyll maximum, oxygen
minimum, and maximum depth) were passed through 0.45-um
hydrophilic mixed cellulose ester filters, which were then frozen
and stored at \(-20^\circ C\) for subsequent DNA extractions post-
cruise. Subsequent next-generation sequencing and microbial
community analyses were conducted following the methods
of Easson and Lopez (2019). During seawater collection,
environmental metadata were simultaneously collected with
instruments on the CTD. These metadata provided context for
determining function and structure of the subsequently described
microbial communities.

The two main limitations of this method are that it does not
directly identify the function of community members or provide
an absolute abundance estimation (only relative abundance).
Assumptions are made based on substantial literature evidence
that these communities are responding directly to a particular
influence. Despite these limitations, these data remain useful in
capturing how microbial plankton dynamics are related to several
biological and oceanographic variables.

**Stable Isotope Analysis of Particulate Organic Matter**
Because a consumer’s isotopic signature is determined by both
its position in the food web and the isotope value of primary
producers, isotopic variation in primary producers can lead to
isotopic variation in consumers not reflective of a change in diet
or trophic status. Thus, when conducting SIA it is essential to
characterize the isotopic signatures of relevant primary producers
so that variation in consumer isotope values caused by shifting
isotope values in primary producers can be distinguished from
changes caused by differences in the feeding habits of consumers.
In order to establish an isotopic baseline in the pelagic GoM,
we conducted SIA on samples of particulate organic matter
(POM) to serve as a proxy for phytoplankton primary production
in the region. Water samples for POM were initially collected
from 12-L Niskin bottles deployed during CTD casts and then
transferred to clean 1-L Nalgene bottles, which were inverted into
500-ml Pall magnetic filter funnels. Samples of POM were then
obtained by filtering 5 – 20 L of water through pre-combusted
(2 h at 450°C) 47-mm (surface and chlorophyll maximum) and
25-mm (oxygen minimum and maximum trawl depth) glass
microfiber filters (GF/F) under low pressure. Once sufficient
material had been obtained, filters were stored frozen at \(-20^\circ C\)
until processing for SIA.

**COLLECTIONS AND DATABASES**

**Specimen Collections**
The majority of specimens (fishes, crustaceans, and gelatinous
zooplankton) collected during both ONSAP and DEEPEND
are housed at the Guy Harvey Oceanographic Center, Nova
Southeastern University, Dania Beach, FL and tracked through
the biotic databases described in section “Biotic databases.”
Molluscan specimens were deposited in the National Museum
of Natural History, Washington, DC, United States, or at
the USF St. Petersburg, St. Petersburg, FL, United States.
All crustacean specimens used for genetics, including tissue
and DNA extracts, were assigned catalog numbers (HBG#) in
curated, public-access databases. All voucher specimens
and tissues were archived in the Florida International Crustacean
Collection (FICC), which currently houses over \( \sim 10,000 \) curated
crustacean specimens.

The holotypes and paratypes for new species discovered
during these projects (e.g., Pietsch and Sutton, 2015; Judkins
et al., 2020) were deposited in museum collections appropriate
for each taxonomic group. Crustaceans and cephalopods
were deposited at the National Museum of Natural History,
Washington, DC, United States. Fishes will be deposited in one of
several museums based on the specific taxon: Lophiiformes will
be deposited at the Burke Museum, University of Washington,
Seattle, WA, United States; Stomiiformes will be deposited at
the Museum of Comparative Zoology, Harvard University,
Cambridge, MA, United States; and representative subsets of
the entire collection, or select specimens, will be deposited at
the Scripps Institution of Oceanography, the Louisiana
State University Ichthyology Collection, the Tulane Ichthyology
Collection, the Virginia Institute of Marine Science Ichthyology
Collection, the Yale Peabody Museum of Natural History, and
the Florida Museum of Natural History.

**Biotic Databases**
All biotic data are stored in Microsoft Access databases
at the Oceanic Ecology Laboratory at NSU (T. Sutton).
Data collected during the ONSAP are stored as “Nekton_Database DDMMYY.accdb” and data collected during DEEPEND are stored as “DEEPEND_Nekton_Database DDMMYY.accdb.” These are relational databases stored on NSU’s servers with replication. There are three main tables: (1) Field Sample/Trawl Field Data table containing the station and sampling depth information, (2) Nekton Database table containing the catch information (taxon, catch in numbers, weight, etc.), and (3) Taxon List table containing the hierarchical taxonomic information (class, order, family, etc.).

In addition, there are other tables to look up and combine data. A Primary Key connects these tables to one another.

Database Availability
DIVER
Biotic and abiotic data collected during the ONSAP are publicly available through NOAA’s Data Integration Visualization Exploration and Reporting (DIVER) tool found at https://www.diver.orr.noaa.gov/. DIVER is a data warehouse and query tool that allows public access to NOAA’s Damage Assessment, Remediation, and Restoration Program data. These data are collected in response to, and/or restoration of, environmental damage caused by oils spills, releases of hazardous waste, or vessel groundings. The DIVER Explorer query tool can be used to search, filter, and download these data using links to popular datasets, guided queries, or keyword searches. Data mentioned in this paper can be located by linking to the popular dataset “Deepwater Horizon NRDA data” and performing a keyword search for “Meg Skansi.”

GRIIDC
Biotic (ONSAP and DEEPEND) and abiotic (DEEPEND only) data are also publicly available through the GRIIDC, housed at the Harte Research Institute for GoM Studies at Texas A&M University, Corpus Christi. GRIIDC is a team of researchers, data and topic specialists, and information technology professionals who have developed a data management system to organize, store, and disseminate data collected by GoM researchers as part of the Master Research Agreement between British Petroleum (BP) and the GoM Alliance. GRIIDC has secured a funding agreement with the GoMRI, the funding body of the GoM Alliance, to continue providing data management and the dissemination of datasets to the scientific community (both GoMRI funded and non-GoMRI funded research) for a minimum of 10 years beyond the conclusion of formal GoMRI funding in 2020 (i.e., through the year 2030, at a minimum).

All data produced by GoMRI-funded individuals and research consortia (such as DEEPEND) are required to be submitted to the GRIIDC repository in a timely fashion, typically within one year of data collection and/or processing. Upon submission, all datasets undergo a rigorous vetting process led by GRIIDC subject matter experts who work with researchers and the Data Manager to ensure data integrity, organization, and discovery, including descriptive, ISO-19115-2 compliant metadata. All datasets housed by GRIIDC are assigned Digital Object Identifiers (DOIs) in the same manner as publications to allow future researchers to use and cite the data. See Supplementary Table 5 for a list of datasets and their corresponding DOIs. All datasets are available at https://data.gulfresearchinitiative.org/.

NCEI
Most environmental data, such as CTD and ship along-track measurements, were submitted on behalf of DEEPEND to the National Centers for Environmental Information (NCEI) through a proprietary process developed between NCEI and GRIIDC. Calibrated water column acoustic backscatter data, including associated metadata, collected during the DEEPEND program are also archived at NCEI. The archive includes raw acoustic backscatter data for each station that has corresponding net tow data (Supplementary Table 5).

NCBI
DNA sequences obtained from barcoding were submitted on behalf of DEEPEND to the National Center for Biotechnology Information (NCBI) database. A compendium of specimen information that includes ID Number, Cruise Number, Collection Date, Collection Location, Taxonomic Species Identity (Order, Family, Genus, and Species), and NCBI GenBank Accession Numbers have been deposited into GRIIDC (Supplementary Table 5). Additionally, the small subunit rRNA gene was sequenced from samples collected in the water column to identify the microbial community. These sequences were deposited in the NCBI Short Read Archive where bioproject accession numbers were assigned.

DATA AVAILABILITY STATEMENT
Data are publicly available through the Gulf of Mexico Research Initiative Information & Data Cooperative (GRIIDC) at https://data.gulfresearchinitiative.org (doi: 10.7266/N7R49P43; 10.7266/N7MC8XDC; 10.7266/n7-ceq1-5g82; 10.7266/N7GM85P1; 10.7266/N73R0QXS; 10.7266/N7PV6HS1; 10.7266/N7Q52N08; 10.7266/N7KD1W8J; 10.7266/N7VM49MP; 10.7266/N7Q4VK0; 10.7266/N7X065FZ; 10.7266/N7K2W65; 10.7266/n7-77gs-w736; 10.7266/N7FF3QFK; 10.7266/N7610XD5; 10.7266/N72805Q8; 10.7266/N7X9P97F; 10.7266/N7DV45V; 10.7266/n7-j3c9-4s47; 10.7266/N7TM78J6; 10.7266/N7ZC81X8; 10.7266/N7NC5ZK6; 10.7266/N7ZK5F24; 10.7266/n7-bhjkh-nh73; 10.7266/N7CN29W; 10.7266/N7H4IP6; 10.7266/N74871; 10.7266/N76T0K11; 10.7266/N7BK19RB; 10.7266/n7-ac8e-0240; 10.7266/N7POX3T; 10.7266/N7VXDK2; 10.7266/N7XP7385; 10.7266/N7902234; 10.7266/n7-dd3p-t155; 10.7266/n7-05f6-th15; 10.7266/N7ZG6QQ9; 10.7266/N7XP73B2; 10.7266/N7HM56TD; 10.7266/N71CITZC; 10.7266/n7-1x7-4n30; 10.7266/n7-3py-g470; 10.7266/n7-hhnpq-kh83; 10.7266/N75M63Q3; 10.7266/n7-c56k-dp86; 10.7266/N7VX0F19;
ETHICS STATEMENT

The animal study was reviewed and approved by the Florida Atlantic University Institutional Animal Care and Use Committee.

AUTHOR CONTRIBUTIONS

All authors conducted the research, analyzed the data, and contributed to the manuscript. AC and TS oversaw all aspects of this research, including specimen, sample, and data collection, analysis and data management. All authors have agreed to being listed as such and approve of the submitted version of this manuscript.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmars.2020.548880/full#supplementary-material

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