Habitat use and activity patterns of female Deacon Rockfish (Sebastes diaconus) at seasonal scales and in response to episodic hypoxia

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Abstract We combined a high-resolution acoustic telemetry array with presence/absence receivers to conduct a preliminary study of the seasonal movements, activity patterns, and habitat associations of the newly described Deacon Rockfish (Sebastes diaconus). Eleven mature female Deacon Rockfish were tagged and monitored during an 11-month period, at a nearshore rocky reef off Seal Rock, Oregon, USA, an area of recurring seasonal hypoxia (defined as dissolved oxygen concentration [DO] < 2 mg l$^{-1}$). Two tags were detected leaving the study area by day 35, indicating predation or emigration. Three tags became inactive within the array, indicating tag loss or fish death. Six “resident” fish inhabited the array for 246–326 days. Resident fish exhibited high site fidelity, small home ranges (mean 95% KDE = 4907 m$^2$), and consistent activity patterns for the duration of the summertime high-resolution array (5 months), except during seasonal hypoxia. Resident fish were strongly diurnal in summer, with high levels of daytime activity above the bottom in relatively rugose habitat, followed by nighttime rest periods in deeper, less rugose habitat. During summertime hypoxia, resident fish exhibited less daytime activity during daytime hours with no rest periods at night, inhabited shallower water depths, and moved well away from their core activity areas on long, erratic forays. During the winter, diel patterns were less evident with higher activity levels at night (than in the summer) and lower activity levels in the day (than in the summer). We propose that some Deacon Rockfish continuously inhabit nearshore reefs throughout the year, but that daily/seasonal movement patterns, seasonally occurring hypoxia, and prey preferences for planktonic organisms influence relocation.

Keywords Sebastes diaconus • Acoustic telemetry • Movement • Behavior • Hypoxia • Seasonal • Planktivores

Introduction

The spatiotemporal scales of movement patterns for marine fishes are highly variable (Pittman and McAlpine 2003), with tunas and other large pelagics making migrations on the order of thousands of kilometers (Galuardi et al. 2010; Kraus et al. 2011) and other organisms remaining relatively sedentary in a singular spot (Bryars et al. 2012; Buston 2004). The temporal scales over which these movements occur vary widely from ontogenetic movements such as salmon and eels (Hansen et al. 1993; van Ginneken et al. 2005) to daily vertical migrations in sand lances and lanternfish (Engelhard et al. 2008; Dypvik et al. 2012). Fish movements vary within the same genera leading to niche partitioning (Sbragaglia et al. 2019). Movement can even differ within the same species resulting in distinct behavioral differences (e.g., for size, sex differences, reproductive behavior, etc.; Bell and Sih 2007; Barnett et al. 2011). Understanding intraspecies behavioral plasticity, especially within a habitat occupied by multiple
species, is ecologically necessary to understanding resource allocation (Leggett 1977; Hays et al. 2016). Further, in the event that the species are caught in fisheries, understanding the movement dynamics, especially in mixed stock fisheries, is essential for sustainable management (Peer and Miller 2014; Ogburn et al. 2017).

Oregon’s nearshore recreational fisheries primarily target schooling rockfish inhabiting nearshore rocky reefs. These schools are predominantly composed of Black Rockfish (Sebastes melanops), Blue Rockfish (Sebastes mystinus), and the newly described cryptic species Deacon Rockfish (Sebastes diaconus) (Fable et al. 2015). Although much is known about the movements of Black Rockfish, Blue and Deacon Rockfish have only recently been recognized as separate species, and little is known about their movements in Oregon (Parker et al. 2007; Parker et al. 2008).

In contrast to Black Rockfish, which are consistently captured throughout the year off the central Oregon coast, Deacon Rockfish are often nearly or completely absent from summer catches for weeks or months, causing fishers and managers alike to suspect seasonal and/or ontogenetic migration away from nearshore reefs (Dick et al. 2017; C Heath, pers. comm). An alternative to the seasonal and/or ontogenetic migration hypothesis is that catchability varies seasonally, and Deacon Rockfish remain at nearshore reefs throughout the year. Whether or not Deacon Rockfish migrate has distinct ecological implications, which in turn affect how to manage fisheries for this species. In addition to seasonal movements, some rockfish are hypothesized to exhibit diel shifts in their behavior. Black Rockfish have been shown to make large vertical movements associated with sunrise and sunset (Parker et al. 2008) whereas Copper Rockfish (Sebastes caurinus) and Quillback Rockfish (Sebastes maliger) exhibited home range movements associated with tidal flow in the Puget Sound (Tolimieri et al. 2009). A complete understanding of the inter- and intra-reef movement of Deacon Rockfish on daily and monthly timescales is important for fisheries management (Crossin et al. 2017). Specifically, managers increasingly seek non-lethal, spatially extensive surveys, making knowledge of seasonal fish locations essential to survey design (Berger et al. 2015; Berger et al. 2017).

Periodic variability in the abiotic environment of a fish can also impact its health and behavior in complex ways, altering normal activity and movements (Gray et al. 2002; Grantham et al. 2004). Oregon coastal waters are subject to seasonal hypoxia (dissolved oxygen [DO] < 2 mg L⁻¹) in the upwelling favorable summer months, the varied effects of which have been documented for other species of rockfish, but not for Deacon Rockfish (Diaz and Rosenberg 1995; Hopkins et al. 1995; Rankin et al. 2013). Responses to hypoxia can lead to fish dispersal or fish concentration, but have also been shown to affect home range, site fidelity, and refuge-seeking behavior in Rockfish (Pihl et al. 1991; Rankin et al. 2013). Changes in environmental conditions can result in fish remaining within an area, but cause fish to be behaviorally or physically “unavailable” to fishery capture or survey, or alternatively may make fish more susceptible to capture in a fishery (Hannah and Blume 2016; Stanley et al. 1999). As managers often use catch rates as a proxy for fish abundance, knowing the level of environmental influence on fish behavior is essential.

Investigating the movements of Deacon Rockfish over time, and throughout a range of environmental conditions, can provide much-needed data on their natural history, responses to changing environmental conditions, and can inform survey design. Determining the scale of movements for initial study design can be challenging for Rockfish as movements for reef inhabitants may range from high site fidelity over periods of months or years for some species (S. melanops, S. maliger) to the wide-ranging Canary Rockfish (Sebastes pinniger) (Parker et al. 2007; Hannah and Rankin 2011). Acoustic telemetry is an effective tool for establishing baseline movements and behavior of tagged fish, while providing scale of movements to inform the design of more comprehensive investigations. The goal of this study was to use high-resolution acoustic telemetry combined with simple presence/absence data to track Deacon Rockfish over a one-year period to: (1) provide preliminary scales of movement, (2) describe daily and seasonal patterns of behavior and use of habitat, and (3) examine the relative influence of seasonal hypoxia on movements, behavior patterns, and habitat usage.

Materials and methods
Study site and Array

The Seal Rocks study area, on the central Oregon coast, encompassed 11.84 km² (2.86 km × 4.14 km) of
nearshore rocky reef. The study area was comprised of bedrock benches, boulders, and rocky pinnacles for which 2 × 2 m multibeam resolution and side scan bathymetry data was available (Fig. 1). The primary habitat was variably rugose with swaths of smaller, complex secondary habitat in between. Specifically, our study area included an isolated rocky reef ranging in depth from 12 to 26 m that could be surrounded by a perimeter of receivers in low-relief habitat, to detect fish traveling beyond the reef. Results of a 15-day range testing study (conducted in the most rugose, high-relief area of the reef) provided worst-case detection statistics through a range of swell and wind conditions. The results suggested an optimal receiver spacing of 250 m (see online supplement for more information).

Based on these findings, and in consideration of habitat, ocean conditions, and potential range of travel for fish, we used two different acoustic telemetry array designs to maximize tag detections during both summer and winter periods (Andrews et al. 2011). The high-resolution summer array (May–September, 2016) included an inner grid of VPS (VEMCO Positioning System) receivers, surrounded by an outer perimeter fence of receivers used to detect any fish leaving the area (Fig. 1). The inner grid encompassed a 2.23-km² reef area to acquire fine-scale movements and habitat association, and was comprised of 21 VR2W receivers with co-located V16 coded synchronization tags (synctags, V16T-4L and V16-4L) moored at 250 m spacing to allow tag transmissions to be detected by presence/absence-only detection. Black circles indicate 250 m detection range for receivers used for presence/absence detection. Inset is the Washington, Oregon, and Northern California coastlines. Star in Oregon is the Seal Rocks study area. Triangle indicates the position of the moored CTD with oxygen sensor.
multiple receivers. Also moored were two permanent, centrally located V16 reference tags (VEMCO, Nova Scotia, Canada). Synctags are acoustic transmitters used for time synchronization that allow calculation of triangulated positions for each fish (VEMCO, Nova Scotia, Canada). This system can provide accurate (5–15 m) position information but only for a relatively small number of tagged fish, due to the high density of synctag transmissions (VEMCO, Nova Scotia, Canada). Acoustic receivers and synctags were positioned 3 and 4 m off bottom, respectively. The 18-receiver perimeter fence was 9.74 km long, positioned 1200 m outside of the VPS array, and surrounded the north, west, and south side of the array (excluding the shallow shoreline side). Perimeter fence moorings were placed 500 m apart for presence/absence detection by a single receiver, in depths ranging from 12 to 39 m.

A Sea-Bird SBE16plusV2 CTD, equipped with a SBE43 oxygen sensor, was moored in 27 m of water depth at the NW corner of the VPS array, to record conductivity, temperature, depth, and dissolved oxygen (Fig. 1; Sea-Bird Electronics, Bellevue, WA, USA). The CTD was positioned 1 m off bottom and sampled once every 60 min, during the summer months. The CTD was downloaded and serviced approximately bimonthly and was removed for winter.

Ocean conditions in Oregon during winter months (October–March) are typified by large, energetic storms that generate considerable acoustic noise. This noise greatly increases tag detection interference, particularly in shallow water. Thus, for the winter array, we removed all co-located synctags to increase the detection probability of fish tags. This low-resolution winter array had 9 receivers spaced 350 m apart covering an acoustic detection area of approximately 2.08 km² to acquire presence/absence-only data for fish within the reef area (Fig. 1). All acoustic receivers for both arrays (summer and winter) were downloaded and serviced approximately bimonthly on average.

The acoustic receivers for the VPS grid and perimeter fence were deployed on 5/10/16 and 5/11/16, and the Sea-Bird SBE16plus V2 was deployed on 5/17/16 (Fig. 1). These instruments were removed on 9/30/16 and replaced with the simple 9-receiver presence/absence winter array (Fig. 1). The winter array was downloaded on 11/16/16, 1/29/17 and removed on 4/18/17. Both winter and summer arrays performed well, providing good coverage throughout the 11-month period. One receiver in the summer VPS grid failed to download on 5/31/16 and was replaced and one receiver was found missing upon the final pull of the winter grid.

Acoustic tags and fish handling

Fish were tagged with V13AP coded acoustic transmitters (69 kHz, 153 dB, 13 × 42 mm, pressure sensing to 136 m) which transmit a unique tag ID to the receiver. The tags also transmit the tags depth (m) and acceleration rate (m s⁻²) which is used as a measure of activity (VEMCO, Nova Scotia, Canada). To maximize detections, we chose to use the larger V13 tag (weight = 6.5 g in water) which provided long battery life (365 days) at a high sampling rate (random delay 60–180 s); therefore, we elected to tag larger fish (fork length > 32 cm) to reduce the effect of tag weight.

Due to concerns with species identification of this cryptic species, Oregon Department of Fish and Wildlife Marine Fisheries Research staff were trained with known samples to visually differentiate between Deacon and Blue Rockfish. Further, species identification of tagged fish was visually confirmed by Dr. Wolfe Wagman, coauthor of the Deacon Rockfish Frable et al. (2015) species description.

Deacon Rockfish can suffer fatal injuries from barotrauma when captured from depths >28 m (Hannah et al. 2008; Hannah et al. 2012). Further, imprudent surface handling results in reduced submergence success of released fish (Hannah et al. 2008). To mitigate these effects, the following techniques were used to capture and handle fish chosen for tagging:

1. To compensate for the weight of the acoustic tag, we selected fish >32 cm total length which, for Deacon Rockfish, only include sexually mature females (Hannah et al. 2015).
2. Fish were captured using hook and line gear with small terminal tackle (Sabiki rigs), in water depths less than 26 m. Following Rankin et al. (2016), fish were immediately recompressed in drum-type cages and held at depth for 24 h to ensure survival and to resolve barotrauma before tagging. Studies have shown that many rockfish species, including Deacon Rockfish, do not experience barotrauma a second time upon returning to the surface within 24–48 h of initial recompression. Therefore, they can be handled at surface pressure during that time without further gas expansion injury (Hannah et al. 2012; Rankin et al. 2016).
Fish were tagged externally as per Hannah and Rankin 2011, in order to decrease surgical trauma and to increase tag detection in the array (Dance et al. 2016).

Fish were released into a floating bottomless sea pen after tagging, to descend under their own power (Hannah et al. 2008).

Data analysis

Oceanographic data

To examine how fish behavior changed during periods of hypoxia and normoxia, we used our CTD to identify periods of normoxia and periods hypoxia. Dissolved oxygen (DO) levels are strongly influenced by water temperature. Therefore, we minimized temperature as a confounding factor by identifying and selecting time periods of hypoxia and normoxia with similar temperature ranges and means. Following Diaz and Rosenberg (1995), the hypoxic period was defined by a mean dissolved oxygen \(<2 \text{ mg l}^{-1}\) and the normoxia period by a mean dissolved oxygen \(\geq 2 \text{ mg l}^{-1}\). Differences in oxygen and temperature during these two time periods were compared using a Welch’s two sample t test. Based on examination of the CTDs dissolved oxygen data, we designated two 11-day time periods: one normoxia period and one hypoxia period. These periods definitions were used to subset the acoustic telemetry data to be used in a focused analysis of how home range, site fidelity, activity, and depths differed between hypoxia and normoxia periods.

Residency and habitat usage

Acoustic tag transmissions from tagged semi-pelagic fishes (such as Deacon Rockfish) are highly detectable, generating hundreds of validated detections per day in most conditions. However, during high ambient noise events, such as high wind and swell generated by storms, detection rates are reduced; therefore, we used a minimum of six validated detections per day to establish residence. Six validated detections allowed us to determine whether a tag was on the sea floor or on a live fish, as indicated by variation in transmitted depth and/or activity level. Residence time was defined as the number of days live fish (active tags) were detected during the 11-month study period.

We first examined how the core area and home range differed between tagged fish. Spatial analyses were only conducted during summer months when we had high-resolution position data from the VPS array (May–September). Position data used for home range and space utilization analyses was filtered for \(<21 \text{ m HPE (horizontal position error)}\), defined as “a relative, unitless estimate of how sensitive a calculated position is to errors in its inputs”, and is the estimated precision of the system for this array and the conditions (VEMCO, Nova Scotia, Canada). Initial analyses filtering HPE data to values as low as 3 m demonstrated that the only effect on our data was to reduce our sample sizes and therefore we elected to only filter data to a value of 21 m. We estimated the home range and core areas for each tagged fish residing within the array, where home range was defined as the 95% kernel density estimator (KDE) and the core area was defined as the 50% KDE. KDEs were calculated using the program Geospatial Modeling Environment (GME, version 0.7.4.0) implemented in R (Beyer 2015; R Core Team 2018). Areas were estimated using the tool “isopleths” c(0.95,0.5) in GME.

We next examined how home ranges and core areas differed between daytime and nighttime. Daytime and nighttime were defined as daily 5-h periods in the middle of the day or night. These time periods were chosen to remove potential crepuscular or transition behaviors from the analyses. For analysis of both the home range and core area data, we estimated a maximum distance from the calculated center of the core area using “genpointinpoly” in GME. We also examined the foray behavior of each fish in their core area and home range. Forays were defined as a series of movements, which were sequential in time and space, at least 100 m away from the calculated center of activity, and lasting for more than 1 h (Rankin et al. 2013). Following a similar study, in Rankin et al. 2013, foray distances were calculated from individual detections, and foray duration was estimated based on the time a tagged fish left and returned to a location less than 100 m from its center of activity. These analyses of home range and core areas were then repeated with telemetry data subset to only include the hypoxia and normoxia periods defined in the “Oceanographic data” section.

To determine if fish utilize habitats of different ruggedness during day and night, underlying bathymetric data from high-resolution Digital Elevation Model (DEM) layers were extracted from each tagged fish’s

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daytime and nighttime core area position data. Geomor-
phology analysis was conducted using the Vector Rug-
gedness Modeling (VRM) tool in the Benthic Terrain
Modeler (BTM) implemented in ArcGIS 10.3 (Hobson
1972; Sappington et al. 2007; ESRI 2015). Vector rug-
gedness provides a metric for how rugose a reef is;
specifically, higher values denote more rugose reef.
From the underlying DEM raster, a (2 m × 2 m) VRM
layer was created using BTM geomorphology tool, and
neighborhood size (1) was chosen at the smallest
resolution.

**Daily and seasonal behavior**

Analysis of within day and seasonal patterns of acceler-
ometer and depth data were conducted using general-
ized additive mixed effects models (GAMMs) using the
mgcv package in R version 3.5.1, Feather Spray (Wood
2004; Wood 2011). Spatial and temporal autocorrela-
tion were tested for using the DHARMa package
(Hartig 2020). Only data from resident fish were utilized
in these analyses. Data were first explored following the
protocols established by Zuur et al. (2010). Four models
were developed. Two models (one for activity, one for
depth) were developed to assess how fish behavior
changed over the course of our entire study, and two
models were developed to compare activity and depth
during the periods we defined as hypoxic and normoxic.
These data for the second set of models are a subset of
the larger seasonal dataset but were analyzed separately
so as to have an approximately equal number of obser-
vations for each category of oxygen level (see the
“Oceanographic data” section).

For both sets of models, fish activity was log trans-
formed to reduce spread and allow models to be devel-
oped using a normal distribution. Depth did not require a
transformation and was modeled using a normal distri-
bution. For all models, all observations recorded by the
receivers were included, in other words potentially up to
1440 observations per fish per day. However, the ran-
donized ping rates of the tags and the high potential for
every ping to not be received by a receiver reduced the
number of daily observations per tag. For the first set of
models (comparing activity and depth over the duration
of the study), the variables included in the model were
hour of the day (continuous), and month of the year
(categorical). An interaction between hour of the day and
oxygen level was also examined.

In both sets of models, continuous variables were
modeled using a cyclic cubic regression spline to account
for the cyclic nature of hour of the day. A cyclic cubic regression spline requires that the best fit line start
and stop at the same point on the y-axis. With hour of
the day, this is important because a large behavioral shift
from, for example, 23:59 to 00:01, is ecologically unre-
realistic. Regardless of model, we included the tag ID of
each individual fish as a random effect in the model to
account for intra-tag variability (Pinheiro and Bates
2000). We also included a first-order autoregressive
correlation structure (AR1) to account for temporal au-
tocorrelation between successive observations (Zuur
et al. 2009). No evidence of spatial autocorrelation
was found. Model selection was conducted by compar-
ing Akaike Information Criterion (AIC) values for all
possible model formulations (Burnham and Anderson
2004). The model structure with the lowest AIC value
was deemed the best-fit model.

**Results**

**Oceanography**

Periodic multiday hypoxic conditions were detected in
the Seal Rocks array throughout the summer season
(Fig. 2). The dates assigned for the 11-day comparison
periods were: (a) hypoxia; July 20–30 (DO range 0.63–
2.40 mg l\(^{-1}\), mean DO = 1.4 mg l\(^{-1}\), mean T = 7.91 °C),
(b) normoxia; August 16–26 (DO range 2.43–
5.60 mg l\(^{-1}\), mean DO = 3.19 mg l\(^{-1}\), mean T =
7.90 °C). (Fig. 2). Oxygen levels during hypoxia and
normoxia differed significantly from one another
\(t(478) = -40.14, p < 0.001\). Temperature was also sig-
ificantly different during the two periods \(t(478) =
3.58, p < 0.001\). However, the mean temperature differ-
ence between the two time periods was only 0.01 °C,
indicating conditions were highly static.

**Fish**

Eleven female Deacon Rockfish were tagged on 5/17/
2016. Fish ranged from 33 to 41 cm total length. Based
on length weight relationships, the tag weighed from 0.62–1.19% of fish body weight (unpublished data). Prior to release, tagged fish were alert, active, free of barotrauma signs, and were able to descend without assistance (Table 1).

Residency and habitat usage

After release, all 11 fish returned to the location of capture within 27 h (home foray, Table 1). One fish was detected in the grid for 112 days, before showing evidence of predation. Four fish had residence times between 13 and 37 days before leaving the array; two were located within the fence area, but were inactive, indicating tag loss, predation, or mortality. The other two fish were detected leaving the perimeter area via the north fence, and designated as “traveler” (Table 1, Fig. 1). Six fish “residents” remained in the grid and were consistently detected for 246–326 days (Table 1). 

The 19 weeks of high-resolution VPS data showed the six resident fish had high site fidelity and small home ranges ranging in area from 3511 to 6875 m² (mean 95% KDE = 4907 m²; Table 2, Fig. 3). These home ranges correspond to circles with a radius of approximately 33–47 m. Core areas ranged from 324 to 756 m² (mean 50% KDE = 477 m²; Table 2, Fig. 3). These core areas correspond to circles with a radius of approximately 10–16 m. Mean nighttime home range and core area size were 50 and 25% smaller than daytime home range and core area size, respectively (Table 2, Fig. 3). With the exception of fish TD03, which was undetected after day 246, five resident fish continued to be active and detected in the winter grid for 306–323 days (Table 1). Four resident fish (TD04, TD06, TD09, and TD10) consistently utilized an overlapping home range area (Fig. 3).

All six resident fish showed a substantial change in behavior during the hypoxic period, resulting in both larger and smaller core areas used. Foray behavior was exhibited by all six resident fish during the hypoxia period only. Forays ranged in duration from 23 to 88 h and from 923 to 2985 m traveled (Table 2, Fig. 4). The farthest distance from the calculated center of the core was 807 m for TD02 (Table 2, Fig. 4).

Resident fish used more rugose habitat during daytime period than at night, although the differences were much lower for fish TD03 than for other fish (Fig. 5). Further, fish utilized significantly more rugose habitat
during the day, night, or in total as compared to the average habitat rugosity for the area encompassed by the array (Fig. 5).

Daily and seasonal behavior

The best-fit models for both activity and depth included all variables and their interaction plus the addition of the random effect of fish tag (Table 3). For activity, the model explained 21.8% of the variability in the data, and for depth, the model explained 58% of the variability in the data (Table S1). During the summer months of May–September, resident fish showed a trend of higher diurnal activity for 8–12 h a day in shallower depths versus lower activity levels in deeper waters at night (Table 3, Figs. 6 and 7). This day/night trend became less evident in October, and was not evident from November through March, during which time there were no major differences in average activity level or depth readings between day and night (Figs. 6 and 7).

Table 1  Summary information and status of 11 tagged Deacon Rockfish. Fish status indicates area of final detection and classification of fish: fence (detected on fence), inactive (tag inactive), resident (in VPS array), and traveler (moved beyond fence). Residence time was defined as the number of days live fish (active tags) were detected during the 11-month study period. Home foray is the distance between the position of release of tagged fish to a location less than 100 m from the calculated center of activity. Home foray duration is the time elapsed between the time of release of a tagged fish to a location <100 m from the calculated center of activity.

| Fish ID | Total length (cm) | Residence time (day) | Home foray distance (m) | Home foray duration (hh:mm) | Status | Predation/mortality | Departure (date) |
|---------|------------------|----------------------|-------------------------|-----------------------------|--------|---------------------|------------------|
| TD01    | 33               | 13                   | 1896                    | 2:36                        | Fence/inactive | 05/31/2016*        |
| TD02    | 35               | 323                  | 581                     | 14:58                       | Resident   |                     |                  |
| TD03    | 33               | 246                  | 556                     | 18:35                       | Resident/traveler | 01/18/2017**    |
| TD04    | 34               | 312                  | 659                     | 10:22                       | Resident   |                     |                  |
| TD05    | 41               | 27                   | 1598                    | 16:11                       | Traveler   | 06/14/2016**       |
| TD06    | 35               | 306                  | 311                     | 26:46                       | Resident   |                     |                  |
| TD07    | 36               | 37                   | 341                     | 11:20                       | Fence/inactive | 07/06/2016*    |
| TD08    | 39               | 35                   | 382                     | 21:00                       | Traveler   | 06/20/2016**       |
| TD09    | 39               | 326                  | 311                     | 22:09                       | Resident   |                     |                  |
| TD10    | 40               | 322                  | 404                     | 8:00                        | Resident   |                     |                  |
| TD11    | 33               | 112                  | 589                     | 21:44                       | Resident/inactive | 07/08/2016*    |

Table 2  Home range, core activity area, and foray summaries for resident Deacon Rockfish. Home range is the 95% Kernel Density Estimator (KDE) and core area is the 50% KDE. Forays were defined as a series of movements, which were sequential in time and space, at least 100 m away from the calculated center of activity, and lasting for more than 1 h (Rankin et al. 2013).

| Resident fish ID | Home range/core area/day/night | Normoxia | Hypoxia | Hypoxia foray |
|------------------|--------------------------------|----------|---------|---------------|
|                  | Home range | Core area | Home range | Core area | Home range | Core area | Home range | Core area | Foray distance (m) | Distance from center (m) | Duration (hh:mm) |
| TD02             | 3715       | 324       | 3692      | 499       | 1949       | 162       | 3487       | 531       | 11,171           | 1339               | 27:37             |
| TD03             | 6530       | 480       | 7122      | 810       | 3087       | 112       | 5125       | 962       | 10,928           | 1328               | 87:48             |
| TD04             | 4546       | 352       | 4762      | 473       | 870        | 67        | 5392       | 610       | 6860             | 714                | 55:25             |
| TD06             | 3511       | 427       | 2922      | 373       | 1282       | 95        | 5016       | 868       | 3480             | 368                | 57:26             |
| TD09             | 4264       | 526       | 3550      | 374       | 5551       | 290       | 4830       | 974       | 6852             | 260                | 29:10             |
| TD10             | 6875       | 756       | 6945      | 875       | 1862       | 123       | 8597       | 1460      | 3682             | 367                | 23:04             |
| Mean             | 4907       | 477       | 4832      | 567       | 2434       | 141       | 5408       | 901       | 7162             | 729                | 46:45             |
The best-fit models describing how both activity and depth varied between hypoxic and normoxic events included all variables and their interactions (Table 3). For activity, the model explained 27.2% of the variability in the data (Table S2). For depth, the model explained 36.3% of the variability in the data (Table S2). During normoxia, fish increased activity and moved to shallower depths during sunrise hours and decreased activity and descended to deeper depths at sunset (Fig. 8). This was in contrast to hypoxia conditions during which activity levels remained relatively stable, and changes in occupied depth over the course of the day were drastically reduced (Fig. 8).

**Discussion**

Overall, in our study, there is strong evidence of site residency for female Deacon Rockfish. The resident Deacon Rockfish displayed high site fidelity having relatively small core areas during the day, and even smaller core areas during the night. Activity and location in the water column displayed a strong diel cycle that shifted seasonally with more static activity levels and depth usage during winter months. Hypoxic events altered daily behavior and depth occupancy as well as the way in which fish utilized the reef. Although we found strong evidence of residency, the three travelers detected leaving the array may indicate response to an
environmental cue, a shift in core area beyond the scope of the array, and/or emigration from the nearshore reef area.

High long-term survival and consistently high detection of resident fish during our 11-month study indicate that some Deacon Rockfish do not exhibit a seasonal migration away from nearshore reefs. This finding is supported by a concurrent study where researchers were able to collect Deacon Rockfish using hook and line gear, with small terminal tackle, throughout a 12-month period (Vaux et al. 2019). Further, underwater video observations in these areas routinely observed Deacon Rockfish during summer months (Rasmuson et al. 2020). In short, our work, in combination with other studies, indicates that at least a component of Deacon Rockfish do not likely undergo a seasonal migration away from nearshore reefs. However, since our study was restricted to large female Deacon Rockfish, we cannot exclude the hypothesis of an ontogenetic migration or a seasonal migration only exhibited by male rockfish or smaller females. Multiple behavior modes have been observed in the Australasian snapper *(Pagurus auratus)* and we cannot discount that possibility here (Egli and Babcock 2004). That said, in the nearshore, Vaux et al. (2019) observed both male and female Deacon Rockfish in all size classes, throughout the year, suggesting that at least a component of the smaller males and females remain in the nearshore. Thus, if there are different behavioral modes, it is more likely a behavioral syndrome as opposed to a sex or length based trait (Bell and Sih 2007).

Given that Deacon Rockfish appear to remain in the nearshore throughout the year, why does the recreational fishing fleet not capture them during summer months? We hypothesize the diet of Deacon Rockfish may be the answer to this question. Food items ingested by Deacon

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**Fig. 4** Combined resident fish positions in VPS array area for two comparison periods: Normoxia and Hypoxia. Data included both daytime and nighttime for the two 11-day periods. Forays away from the core center were only observed during the hypoxic period.
Rockfish in Vaux et al. 2019 included gelatinous zooplankton and small planktonic crustaceans; the colonial tunicate *Pyrosoma atlanticum*, hydrozoan *Velella vellela*, ctenophore *Pleurobrachia bachei*, brachyuran zoeae/megalopae, and pelagic amphipods. Feeding on gelatinous zooplankton requires specific visual abilities which allow some species of Rockfish (e.g., Blue Rockfish) to see and feed upon very small and/or transparent prey items (Hobson and Chess 1988; Hobson et al. 1996). In the closely related planktivorous Blue Rockfish, electroretinograms demonstrated that Blue Rockfish had low sensitivity to lower light levels than more nocturnally active rockfish (Reilly and Thompson 2007). Thus, the cryptic Blue and Deacon Rockfish may have similar unique visual capabilities making them well adapted to daytime light levels and to feeding on small, clear planktonic prey (Green et al. 2014). Accordingly, we hypothesize that Deacon Rockfish may be resistant to standard fishing techniques because recreational terminal gear typically does not mimic the

Fig. 5 Vector Ruggedness Measure (VRM) comparing resident Deacon Rockfish area usage to the VPS array area. Resident Deacon Rockfish daytime VRM (yellow), nighttime VRM (gray), and the entire VPS acoustic array area (red). Text denotes results of Kolmogorov-Smirnov tests comparing Deacon Rockfish usage between time periods and to the grid VRM values. Day-Daytime usage, Night- Nighttime usage, VRM metric for the VPS array, Total-Usage of the tag during day and night combined. Most resident fish used more rugose habitat during the daytime period than during the nighttime period. All resident fish used more rugose habitat during the day, night, or in total compared to the average habitat rugosity for the area encompassed by the array.

![Graphs showing VRM comparisons](image-url)
preferred summer prey of Deacon Rockfish. These results suggest that periods of relatively low catch of Deacon Rockfish in the recreational fishery may not be indicative of a small population size, but rather a seasonal shift in diet and/or behavior. Furthermore, this dietary shift may provide some resistance to overfishing (under current effort levels), for which the high site fidelity and small home ranges of Deacon Rockfish might otherwise make them vulnerable (Patrick et al. 2010).

Table 3 Summary of model selection for fish acceleration (proxy for activity level) and depth occupied. Model selection for activity level (left) and depth occupied (right) includes the entire year’s dataset (upper) and the shorter dataset comparing oxygen level (lower)

A) Seasonal model

| Model structure                                      | Acceleration: $n=190,595$ | Depth: $n=380,889$ |
|------------------------------------------------------|-----------------------------|---------------------|
| $s(Hr \times \text{Month})+\text{Month}+\text{re(Fish)}$ | 0                           | 0                   |
| $s(Hr \times \text{Month})+\text{Month}             | 1156                        | 274,935             |
| $s(Hr)+\text{Month}+\text{re(Fish)}$                | 11,894                      | 19,545              |
| $s(Hr)+\text{Month}$                                | 13,085                      | 287,202             |
| $s(Hr \times \text{Month})+\text{re(Fish)}$         | 17,320                      | 64,143              |
| $s(Hr \times \text{Month})$                         | 19,524                      | 313,897             |
| $s(Hr)+\text{re(Fish)}$                              | 31,763                      | 82,271              |
| $s(Hr)$                                              | 33,844                      | 326,280             |
| $\text{Month}+\text{re(Fish)}$                       | 23,783                      | 33,659              |
| $\text{Month}$                                       | 25,103                      | 291,683             |
| $\text{re(Fish)}$                                    | 44,461                      | 94,205              |
| 1                                                    | 46,721                      | 330,311             |

B) Hypoxia and normoxia model

| Model structure                                      | Acceleration: $n=11,097$ | Depth: $n=22,299$ |
|------------------------------------------------------|-----------------------------|---------------------|
| $s(Hr \times \text{Oxygen})+\text{Oxygen}+\text{re(Fish)}$ | 0                           | 0                   |
| $s(Hr \times \text{Oxygen})+\text{Oxygen}$            | 980                         | 4588                |
| $s(Hr)+\text{Oxygen}+\text{re(Fish)}$                 | 519                         | 1671                |
| $s(Hr)+\text{Oxygen}$                                 | 1688                        | 6135                |
| $s(Hr \times \text{Oxygen})+\text{re(Fish)}$          | 2303                        | 1297                |
| $s(Hr \times \text{Oxygen})$                          | 570                         | 6318                |
| $s(Hr)+\text{re(Fish)}$                               | 1688                        | 2560                |
| $s(Hr)$                                               | 1057                        | 7424                |
| $\text{Oxygen}+\text{re(Fish)}$                       | 2303                        | 5915                |
| $\text{Oxygen}$                                       | 1774                        | 9348                |
| $\text{re(Fish)}$                                     | 2495                        | 5296                |
| 1                                                    | 3473                        | 10,006              |

Bold denotes the most parsimonious model
$s$ variable was modeled as a cyclic spline, $re$ variable modeled as a random effect, $Hr$ hour of the day, $Month$ month of the year, $Oxygen$ Categorical definition of oxygen concentration, $Fish$ fish tag ID

Although methods for calculating home range vary between studies, mean home ranges were considerably smaller for resident Deacon Rockfish (0.0049 km$^2$) than other semi-pelagic species like Blue Rockfish in kelp habitat off Central California (0.23 km$^2$) (Green et al. 2014), and Black Rockfish inhabiting the same reef complex in Oregon (0.55 km$^2$) (Parker et al. 2007). Deacon Rockfish utilized small, consistent core areas with mean nighttime core areas being approximately 75% smaller than the daytime core area. Additionally,
all fish displayed a distinct behavior trend of switching away from utilizing more rugose habitat during the day, to utilizing less rugose habitat at night. Pairing this observation with the nighttime video surveys where Deacon Rockfish have been observed laying directly on flat bedrock (Rasmuson unpublished data) indicates a shift away from more rugose habitat during the day to a flat bedrock sleeping environment. Why sleep out in

![Graph showing the relative effect of month on daily activity level of resident Deacon Rockfish.](image)

**Fig. 6** Relative effect of month of the year on daily activity level of resident Deacon Rockfish. The black line within the gray shading (95% CI) is the smoothed response. Vertical gray bars denote the hours after sunset and before sunrise. Fish exhibited a strong diurnal pattern of activity in the summer months, with relatively high levels of activity in the daytime and lower levels during the nighttime.
the open directly on the bottom? In short, we are unsure. At night, Deacon Rockfish adopt much darker coloration so it is possible that laying near or on the bottom provides camouflage from visual predators. However, one would then wonder why lay in the open rather than in cracks like other rockfishes. Alternatively, they may stop their activity at night because as a visual predator of clear planktonic organisms, they may be unable to see

Fig. 7 Relative effect of month of the year on daily depth level occupied by resident Deacon Rockfish. The black line within the gray shading (95% CI) is the smoothed response. Vertical gray bars denote the hours after sunset and before sunrise. Fish occupied relatively deeper water depths during the nighttime in the summer months, than in the winter months.
Off Oregon, coastal circulation has two primary periods: the period from the fall transition to the spring transition, hereafter winter, and the period from the spring transition to the fall transition, hereafter summer (Huyer 1979; Strub and James 1988). Characteristically during the summer, conditions alternate between periods of upwelling and downwelling on the continental shelf, though in the nearshore, where this study was conducted, these effects are strongly muted (Austin and Lentz 2002). In the wintertime, the conditions are characteristically downwelling favorable and large storms systematically influence circulation. In 2016, the year our study started, the spring transition occurred on March 27 and the fall transition occurred on September 29, whereas in 2017 the spring transition occurred on April 26 (http://damp.coas.oregonstate.edu/windstress/index.html). It seems likely that the change from heterogeneous acceleration and depth during summer relatively to the more homogenous acceleration and depths of winter is associated with the spring and fall transitions. This likely indicates that the seasonal behaviors we observed are due to prevailing changes in circulation. Unfortunately, the rough winter storms off Oregon during precluded us from leaving our CTD mooring in the water and associating our observed behaviors with abiotic factors.

Tagged fish showed greatly reduced activity in their nighttime core areas, but this pattern was less evident during the wintertime. Some potential hypotheses for this wintertime behavior are that the fish receive less rest during storms because high wave action creates turbulence in nearshore reef areas, or that the fish change to nightly foraging activity during these months. Undoubtedly, other hypotheses exist. The seasonal pattern of high levels of daytime, off-bottom activity, followed by a definitive period of rest observed in the summer months aligns with Green et al. (2014), indicating nocturnal sheltering for Blue Rockfish off Central California. Although Deacon Rockfish had not been established as a separate species at the time of Green’s study, the authors confirmed they were already visually differentiating between the two cryptic species and that tagged specimens were all Blue Rockfish (R. Starr, pers comm). The diurnal behavior of Deacon Rockfish is in contrast to Black Rockfish from the Seal Rocks Reef area, which demonstrate mixed diurnal/nocturnal movement patterns (Parker et al. 2008).
All resident Deacon Rockfish exhibited a response to hypoxic conditions, which interrupted 5 months of otherwise consistent movement patterns. The long-duration forays, well away from core activity areas, were atypical as were the depth and activity patterns for that time of year; fish were moderately active at shallower depths, with no high-level diurnal activity and no nighttime on-bottom rest. Our analyses suggest oxygen and temperature differed significantly between the hypoxic and normoxic time periods. However, on average, temperature only differed by 0.01 °C between the two time periods. Accordingly, although temperature is statistically different between the time periods, we hypothesize that the difference is not ecologically relevant, but rather, the larger difference in oxygen levels between the time periods is a better explanation for the observed behavioral shifts. Deacon Rockfish response to seasonal hypoxia differed considerably from the responses of more benthic rockfish in Oregon. For example, Copper Rockfish reduce their home ranges and activity in response to hypoxia, while Quillback Rockfish showed no change (Rankin et al. 2013).

Hypoxia may also result in increased respiration rate in fish, which is a physiological response to decreased oxygen availability. Movement patterns exhibited by resident fish suggest fish increased their baseline activity to compensate for respiratory stress, and moved beyond their core activity areas towards the surface, to seek areas with higher oxygen levels (Nakanishi and Itazawa 1974; Furse et al. 1996; Kim et al. 1995; Palsson et al. 2008). The independent and irregular forays were likely not prey-seeking behavior, as resident fish maintained a high level of site fidelity, as well as habitat and school fidelity through the rest of the summer season, during which variable upwelling/downwelling/relaxation patterns intermittently interject pelagic planktonic prey into the system (Hobson and Chess 1988; Checkley Jr and Barth 2009). Additionally unlikely is reproductive activity, which takes place in late summer/fall with parturition in the wintertime (Hannah et al. 2015).

Although our sample size was necessarily small to accommodate the high numbers of synctag detections needed for high-resolution position data, detection numbers and position data for tagged fish was excellent, as mid-water schooling behavior of this semi-pelagic species benefits acoustic transmission. Detection rates can be problematic for more benthic rockfish in high-relief habitat, as habitat can block or distort the acoustic transmission. The high-resolution inner VPS array, combined with the perimeter fence, and fish tags equipped with accelerometer/depth sensors, provided additional certainty about the fate of fish that remained inside or left the array. A larger study in southern Oregon, using similar methods, but tagging both Deacon and Blue Rockfish inhabiting the same area, could shed light on differences in the cryptic pairs’ movements in various habitats including un-fished offshore reefs, which may act as refuges for older, more fecund fish found in rockfish conservation areas in Oregon.

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**Availability of data and material and code** Data are available on the Oregon Department of Fish and Wildlife’s Natural Resources Information Management Program.

**Authors’ contributions** This manuscript was designed, executed, and written by Leif K. Rasmuson, Matthew T.O. Blume, and Polly S. Rankin. The field investigation was designed and conducted by Blume and Rankin; analysis and writing was by Rasmuson, Blume, and Rankin.

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**Declarations**

**Ethics approval** The Oregon Department of Fish and Wildlife did not have an IACUC (Institutional Animal Care and Use Committee) process in place at the time of the study, one is currently being developed. The collection, handling, and tagging of *S. diaconus* was conducted using the best available science and veterinary practices for minimizing stress and ensuring longevity in captured marine fish, with special emphasis on mitigating the effects of barotrauma. We followed the protocols developed by the American Fisheries Society in Guidelines for the Use of Fishes in Research, which were reviewed and approved by veterinary staff.
of the Oregon Department of Fish and Wildlife, and were found to be consistent with acceptable standards.

Consent to participate and publication All authors participated in this study and consent to its publication.

Conflict of interest The authors declare that they have no conflict of interests.

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