INTRODUCTION

Rhythms are present at all levels of biological organization, from the behaviors of populations and their constituent individual members down to the molecular functions of single cells (reviewed by Dunlap et al. 2004, Refinetti 2006). Studies of behavioral and physiological rhythms in marine species have generally been carried out in coastal areas in relation to day-night and tidal cycles (Naylor 2005). Only a few studies have dealt with these rhythms at deep-water continental margins and in the deep sea, which is one of the largest ecosystems on earth (Glover & Smith 2003). On shelf and slope areas where light is still present, rhythms have been characterized with respect to day-night and seasonal changes (reviewed by Aguzzi & Company 2010). Conversely, in the aphotic deep sea, behavioral rhythms, which are regulated by internal tides and inertial currents, are poorly understood (Wagner et al. 2007, Aguzzi et al. 2009d). For species at
these depths, hydrodynamic changes may replace fluctuations in light intensity as the zeitgeber (i.e. the environmental synchronizer of rhythms), hence modulating the behavior of organisms in relation to displacement for feeding and also affecting other important life traits such as colonization and dispersal (reviewed by Levin et al. 2001, Kritzer & Sale 2006).

The temporal functioning of an ecosystem depends on the sum of the biological rhythms of all of the species that constitute the community (Kornfeld-Schor & Dayan 2003). In deep-water zones, the rhythmic displacements of animals in and out of sampling windows produce changes in measurements of the biomass and biodiversity of a given region (Aguzzi et al. 2003, 2008, Aguzzi & Bahamon 2009, Coll et al. 2010). For example, the compositions of species in trawl hauls and the estimates of biodiversity that are derived from these species compositions can change within a given area depending on the time of the day or the season in which sampling occurs (Aguzzi et al. 2009a, Bahamon et al. 2009). Temporally scheduled trawling should be repeated in the same area so that variations in the species composition of the haul can be used as a proxy for behavioral modulation in the species in the sample (reviewed by Aguzzi & Sardà 2008). However, trawling repeatability is generally constrained for both technical and economic reasons (Raffaelli et al. 2003). In this context, the pace of technological development in sampling and direct observation methods represents one of the major bottlenecks for behavioral studies in relation to habitat cycles in deep oceanic zones (Aguzzi et al. 2009c, 2010).

In the past 2 decades, the number of cabled observation stations with video recording equipment has increased (Benoit-Bird & Au 2006). Unfortunately, the video footage that has been collected from these stations has generally been of short duration compared to what is required to produce a consistent behavioral time series analysis. The reason for this is the absence of proper automation of image analysis, which often obliges researchers to manually survey large quantities of images (Walther et al. 2004, Cline et al. 2007). In this study, we used an automated video-image analysis protocol that combined animal movement tracking and classification using multivariate morphometry (Aguzzi et al. 2009c) to search for possible tide-related behavioral rhythms in deep-sea species. We processed 2 mo of continuous footage showing the behavior of the demersal fauna of a geologically and ecologically important environment, the hydrocarbon seep of Sagami Bay (1100 m depth; central Japan). Time series based on visual counts of the most representative species in the footage were studied in relation to pressure variations at the seabed produced by changes in water depth at tidal pulls. Pressure variations were used as a marker of internal tide-associated bottom currents (Ohwaki et al. 1991), and this oceanographic parameter was also measured continuously during the same interval over which the video recording occurred.

**MATERIALS AND METHODS**

**Video footage and the video-image analysis.** Video footage (62 d from 17 April to 18 June 1999) was captured by a submarine infrared 3CCD video camera mounted on the cabled Real-Time Deep-Sea Floor Permanent Observatory (Iwase et al. 2003), which is a multisensory station located off of Hatsushima Island (Fig. 1; Sagami Bay, central Japan) in a hydrocarbon seep clam field (*Calyptraeogena soyae* Okutani, 1957) at a depth of 1100 m. The video camera acquired images continuously in time-lapse mode, capturing one frame every 5 s, under a constant source of white illumination. Four white 250 W halogen lamps were always on during video recording operations.

Automated video-image analysis was performed according to the methods recently developed by Aguzzi et al. (2009c). A customized script was created in MATLAB 7.1 to recognize and then count eelpouts (*Zoarcid fishes*), red crabs *Paralomis multispina* (Benedict, 1894), and snails *Buccinum soyomaruae* Okutani, 1977. These were selected because they are the most representative fauna of the local hydrocarbon seep community (Fujikura et al. 2002).

Video originally stored on S-VHS tapes was pre-processed for digitization and consequent partitioning into frames at a rate equivalent to the frequency of video acquisition. Automated processing was based on motion detection of animals (i.e. tracking by means of image extraction) and recognition of animals (i.e. classification by profile identification). Motion detection identifies moving objects by following objects through consecutive frames (Lipton et al. 1998). The central area within the camera field was identified as a region of special interest, and the subtraction of consecutive frames took place within that area. Pixel differences indicated moving objects. Moving animals were discriminated from inanimate particulate matter (e.g. fouling), by filtering the extracted images based on size (pixel area) and color (pixel red/green/blue, RGB texture). The resulting animal profiles were classified as different species using the complex Fourier descriptors and supervised standard k-nearest neighbors morphometric analyses (Menesatti et al. 2009). The use of both of these techniques concurrently allowed the comparison of the profile of a newly tracked animal with images included in a reference library that was previously created by Aguzzi et al. (2009c).
**Biological and physical time series analysis.** The temporal variation in visual counts for the selected taxa was considered to be a proxy for their behavioral rhythms at a population level. For each species, time series of visual counts taken at 5-s intervals were binned using a 10-min bin size. Tidally related variations in depth in Sagami Bay are linked to up- and down-sloping deep-sea-bottom tidal currents (Ohwaki et al. 1991). Oceanographic data pertaining to water depth were also gathered at 5-s intervals throughout the period of video recording using a hydrostatic sensor that was mounted on the cabled observatory. The resulting time series were then binned for each 10-min interval to allow for comparisons with the biological data sets. Unfortunately, due to the malfunction of the mooring array, data on the bottom current speed and direction were not retrieved.

The time series analysis was carried out using El Temps software (A. Diez-Noguera; University of Barcelona). Serial Fourier analysis was used to quantify the amplitude of fluctuation on diurnal and semidiurnal temporal scales in the data sets of visual counts for the 3 species (Aguzzi et al. 2005). Time series were partitioned into 24-h segments, and 2 harmonics, 24-h and 12-h, were modeled as being representative of the diurnal and semidiurnal tidal periodicity, respectively. Both harmonics were fitted separately to each time series segment to obtain a Power Content (PC) value as a percentage of the contribution of each harmonic to the total data set variance (%Var$_{24}$ and %Var$_{12}$). PC

![Fig. 1. Geographic and depth location (triangle) of the cabled observatory of Sagami Bay (central Japan)](image-url)
values were considered to be a measure of the strength of rhythms (Hsu 1973). For each selected taxa, %Var time series containing 62 points were then obtained for both harmonics.

The occurrence of significant ($p < 0.05$) differences in the $\%\text{Var}_24$ and $\%\text{Var}_{12}$ time series values among species was evaluated using a 2-way ANOVA as a measure of the differential strength in the rhythms. To do this, species and harmonics were considered as factors. Then, a post-hoc Fisher’s test was conducted to detect significant ($p \leq 0.05$) differences between harmonics and species.

$\%\text{Var}_{24}$ and $\%\text{Var}_{12}$ time series for all selected taxa were screened with the Lomb-Scargle periodogram analysis (Ruf 1999) at the range of 1 to 30 d to assess the occurrence of any significant ($p \leq 0.05$) infradian periodicity (i.e. $\geq 1$ d) corresponding to spring-neap tidal alternations. The periodogram routine calculates the spectral power, $P$, for an increasing sequence of frequencies (in reciprocal units of the time array, $T$). The returned period values are associated with a spectral power.

A cross-correlation function was also calculated to measure the temporal agreement between the amplitude of the %Var time series and bathymetric fluctuations. For this purpose, the %Var time series were compared with water depth maxima and minima for each 24-h period throughout the testing period. The physical time series was lagged by 1 d against the %Var time series for the 24-h and the 12-h harmonics to obtain the correlation value and its significance for lag $= 0$ ($p_0$) (i.e. the contemporary series).

A waveform analysis was carried out on the time series of biological and oceanographic data in order to estimate the mean daily profile of rhythms fluctuations. Each data set was subdivided into 24-h segments, and all of their constituting values were averaged through all of the subsets at corresponding timings (i.e. for each 10-min bin). The result was a waveform in the form of a consensus curve constructed using mean values that were represented along with their standard errors (SEM). A daily mean was estimated from each waveform profile in order to evaluate the temporal amplitude of the waveform peak (i.e. the phase). In order to do so, all waveform values were averaged, and the resulting mean was plotted as a line onto the plot. For a better visualization of the daily pattern of fluctuation, the waveform obtained from each time series was double-plotted by repeating its 24-h profile in a 48-h format.

**RESULTS**

A total of 10 808 visual counts were obtained for eelpouts (zoarcid fishes). The counts of red crabs *Paralimnias multispina* and snails *Buccinum soyomaruae* were 14 441 and 18 311, respectively. The range of fluctuation in the visual counts for these species during the 62-d study period was from a common minimum of 0, when no individual was observed, to a maximum of 138, 91, and 98 for eelpouts, red crabs, and snails, respectively. Behavioral observations showed alternating moments of swimming activity and rest on the seabed for eelpouts. Red crabs exhibited phases of active zig-zag locomotion interspersed with phases of arrested motion. Snails exhibited linear trajectories of crawling motion.

Time series of visual counts for eelpouts, red crabs, and snails are presented in Fig. 2, along with the concomitant sea-level variations. The time series of bathymetric data indicated a fluctuation of $\pm 1.5$ cm, showing alternate phases of large and small deviations that were related to predicted spring and neap tides respectively. Visual counts for eelpouts increased during spring phases, and also between the first and second half of the observation period. No clearly defined fluctuation patterns were discernible for red crabs or snails.

Serial Fourier analysis outputs (Fig. 3) indicated the occurrence of differential strengths in the coupling of depth fluctuations with fluctuations in the visual count time series for eelpouts, red crabs, and snails. The 24-h and 12-h %Var time series for eelpouts showed peaks during spring tidal phases. For crabs, the agreement between the $\%\text{Var}_{24}$ and $\%\text{Var}_{12}$ time series and the bathymetric fluctuations was less pronounced. %Var for both harmonics did not clearly peak during spring tidal phases. Likewise, no agreement between the time series and the bathymetric fluctuations was seen for snails. The $\%\text{Var}_{24}$ and $\%\text{Var}_{12}$ time series did not exhibit discernible fluctuations in relation to spring tidal phases.

The %Var time series exhibited different average ($\pm$ SD) values over the 62-d study; these average values are a measure of the differential strength of the tidal signal. For eelpouts, these percentages were 11.4% ($\pm$6.5) for the 24-h harmonic and 10.4% ($\pm$4.2) for the 12-h harmonic. In red crabs, these were 9.7% ($\pm$4.6) for the 24-h harmonic and 8.7% ($\pm$4.5) for the 12-h harmonic. In snails, the average variances were 11.3% ($\pm$4.0) for the 24-h harmonic and 8.4% ($\pm$3.8) for the 12-h harmonic. The %Var values for eelpouts were significantly higher than those values reported for the other 2 species. A 2-way ANOVA detected the occurrence of significant ($p \leq 0.05$) differences in the %Var between species ($p = 0.0171$) and harmonics ($p = 0.0008$). The post-hoc Fisher’s test (Table 1) revealed significant ($p \leq 0.05$) differences as follows: (i) eelpouts $\%\text{Var}_{24} >$ red crabs $\%\text{Var}_{24}$ and $\%\text{Var}_{12} >$ snails $\%\text{Var}_{12}$; (ii) eelpouts $\%\text{Var}_{12} >$ red crabs $\%\text{Var}_{12}$; and finally, (iii) snails $\%\text{Var}_{24} >$ red crabs $\%\text{Var}_{12}$.
Periodogram analysis outputs for PC time series of data are reported in Fig. 4 as an indication of differential coupling among the visual count time series for the selected taxa and bathymetric fluctuations driven by the spring and neap tidal regime. For eelpouts, a significant 13-d infradian periodicity was detected for the PC time series of both harmonics using Lomb-Scargle periodogram analysis. This periodicity corresponded to that of the

![Graph](image_url)

**Fig. 2.** Eelpouts (Zoarcid fishes), red crabs *Paralomis multispina* and snails *Buccinum soyomaruae*. Fluctuations in water depth showing spring and neap tides and time series of visual counts obtained by automated video-image analysis. Vertical thin lines define consecutive days and have been added to visually correlate physical and biological fluctuations.
bathymetric fluctuations, which also showed a significant 13-d peak in the periodogram plot. The output of the periodogram analysis for red crabs only showed a significant periodicity for the %Var time series values for the 12-h harmonic. For snails, no infradian periodicity was detected by periodogram analysis.

The temporal relationship between fluctuations in the %Var_{24} and %Var_{12} time series and the effects of spring-neap tidal phases was statistically quantified using a cross-correlation test. In eelpouts, a significant crosscorrelation with the spring phase was found both for %Var_{24} ($\rho_0 = 0.30 > 0.25$ for $p = 0.05$) and for %Var_{12}

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**Fig. 3.** Eelpouts (Zoarcid fishes), red crabs *Paralomis multispina* and snails *Buccinum soyomaruae*. Daily variations in water depth and serial Fourier analysis outputs showing the time series of Power Content (PC) values for both the 12-h and 24-h harmonics (%Var_{24} and %Var_{12}), as calculated by fitting these onto 24-h visual count data sets.

**Table 1.** Results of a post-hoc Fisher’s test performed to detect significant differences among %Var time series for 24-h and 12-h harmonics for eelpouts, red crabs and snails. **Bold** indicates the occurrence of significant comparisons ($p \leq 0.05$)

<table>
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<tr>
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<tbody>
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Aguzzi et al.: Behavioral rhythms of hydrocarbon seep fauna

In this study, we analyzed the occurrence of internal-tide-related activity rhythms in different vertebrate and invertebrate species in a deep-sea hydrocarbon seep community. Using an automated video-image analysis protocol that was developed recently (Aguzzi et al. 2009c), we assessed the occurrence of these rhythms using visual counts of zoarcid fishes (eelouts), red crabs *Paralomis multispina* and snails *Buccinum soyomaruae*.

The time series analysis for eelpouts demonstrated the presence of a strong semilunar periodicity on the timescale of spring–neap tidal fluctuations (~13 d; see Fig. 3A). This was interpreted as a proxy for rhythmic increases and decreases in the swimming activity of the local zoarcid population in relation to alternately increasing and decreasing cycles of water flow. Previous studies have already demonstrated that eelpouts can adjust their behavior to match conditions of differing current strengths and directions (reviewed by Uiblein et al. 2002, Trenkel et al. 2004, Lorance & Trenkel 2006). Cummings and Morgan (2001) also described a strengthening of swimming rhythms during spring tidal phases for Zoarcids in the laboratory. The comparison of our results with those previously published (Kimura et al. 1990, Shinohara & Sakurai 2006) indicates the presence of a common tidal regulation of behavior in different species within the eelpout morphotype.

Data on the speed and direction of currents at the seabed were not available for comparison with biological time series due to the malfunctioning of the mooring array. These oceanographic data are necessary to determine the existence of direct cause-effect relationships between behavioral patterns and changes in water motion (Aguzzi et al. 2010). In Sagami Bay, bathymetric excursions have been associated with internal tides, which mix the water column from the surface down to the deep-sea bottom (Kitade et al. 2004), generating up and down currents along the slope of ~20 cm s\(^{-1}\) (Ohwaki et al. 1991). The along-slope motion of these currents was assessed by underwater photography, which identified stream-oriented sediment marks and characteristic megabenthos positioning (reviewed by Okada & Otha 1993).

Reported differential strengths in tidally oriented rhythms for the 3 studied species may be explained by their different mode of active displacement, which in turn accounts for a markedly different substrate interaction (reviewed by Aguzzi & Company 2010). Eelpouts are benthic boundary layer swimmers that are potentially more vulnerable to changes in water flow than walking crabs and crawling snails (Eguchi et al. 1997). A similarity in the mode of locomotion in phylogenetically different marine species can drive the morphological and metabolic convergence of taxa (re-

The behavioral rhythms that we report here may be the product of an exogenous reactive response, or alternatively, these may be the result of anticipation; therefore, they may be fully endogenously generated via biological clock control (reviewed by Naylor 2005). Small changes in hydrostatic pressure reinforce the tidal rhythm in coastal fishes (Gibson 1984) and crustaceans (Akiyama 2004). Additionally, the modulation of behavior in response to tidal hydrodynamics was previously observed in fishes performing diel vertical migrations (Michalsen et al. 1996) and in coastal crabs (Thurman 2004; Brown et al. 2010). The activity cycles reported herein were not observed under laboratory constant conditions, and therefore, the presence of a biological clock sustaining an endogenous rhythm cannot be proven.

Any discussion on the exogenous or endogenous nature of tidally oriented rhythms should be carried out by considering 2 important aspects of the deep sea: the highly variable character of current flow regimes (reviewed by Garrett 2003, Simmons 2008) and the food-depletion (Childress 1995). Most of the effect of currents on the behavior of deep-sea animals has been related to feeding (Lampitt et al. 1983, Priedie et al. 1994, Jones et al. 1998, Levin et al. 2001). An exogenous response to food odor from one current direction may evoke a swimming response. In the deep sea, scavengers can locate corpses over great distances by olfaction of organic chemicals dissolved in water plumes (Tamburri & Barry 1999, Bailey et al. 2007). Food can also be involved in the endogenous response because its temporally scheduled administration reinforces endogenous behavioral rhythms in fishes and decapods. Food pulses set the phase of functioning of light-independent peripheral oscillators such as the liver and the hepatopancreas (in fishes and decapods, respectively; Fernández de Miguel & Aréchiga 1994, Reebs 2002). In this scenario, directional food odors within tidal wave fronts may reinforce the hydrodynamic entrainment of behaviors in deep-sea fauna. Food perception, localization (by behavioral activation), and consequent food retrieval and ingestion may all represent important aspects of the behavioral repertoire that interact in tidal entrainment (Aguzzi et al. 2009d). However, to our knowledge, no study to date has demonstrated that food can entrain a tidal rhythm.

Our deep-sea observations were carried out under bright lights, creating an unnatural environmental that may affect behavioral tests by damaging the optic apparatus of local fauna (Widder et al. 2005) and by favoring an escaping response (Gaten et al. 1999). Additionally, the circadian rhythmicity of individuals can be altered by constant light-on conditions in the laboratory (Dunlap et al. 2004), but that phenomenon has not yet been tested in deep-sea animals. However, our recording conditions did not impair the detection of clear swimming rhythms for fishes, as have been reported in other behavioral studies where light attraction responses have also been observed (e.g. Trenkel et al. 2004; Lorance & Trenkel 2006). In this work, we detected a tidal-related rhythm in fishes, which usually had trajectories avoiding the central lighted seabed area (Aguzzi et al. 2009c). Conversely, the other 2 invertebrate species had trajectories that fully encompassed the light field, but their tidally oriented rhythms are comparatively less clear. Crabs and snails may be
less affected by direct illumination exposure than fishes (Watanabe 2002). The possibility that these animals may have rhythms that might be disturbed by the artificial lighting should be further investigated.

Our study indicates that video-imaging techniques can be successfully used for long-term and remote monitoring of deep-sea rhythms when a suitable level of automation in digital footage processing is achieved. The population approach can be successful for the determination of species responses to cyclically fluctuating habitat parameters, when the observed rhythmic pattern is considered to be an average property of the animals within the group (Simonetta et al. 2009). Video-imaging technology will contribute to our understanding of the temporal functioning of these ecosystems on diel and seasonal timescales based on the integrated analysis of activity rhythms of species in relation to recurrent habitat events such as internal tidal fluctuations.

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