Spatial and temporal distribution of sea turtles related to sea surface temperature and chlorophyll-\(a\) in Mexican Central Pacific waters

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\textbf{Resumen}.- En este estudio se describe la distribución espacio-temporal de tortugas marinas y su relación con parámetros oceanográficos en el Pacífico Central Mexicano (PCM) durante el 2010 (invierno, primavera y otoño). Los resultados muestran variaciones en la distribución de tortugas marinas. En invierno, la distribución de tortugas fue homogénea en áreas costeras y oceánicas; asociadas a la presencia de frentes térmicos debido a la interacción de una corriente de agua cálida del sur y la intrusión de agua fría proveniente del noroeste, así como con los límites de un giro ciclónico. En primavera, la distribución de tortugas se concentró hacia la zona costera y parte central de la zona oceánica, donde prevalecieron los efectos de una surgencia y los límites de un giro ciclónico. El mismo patrón de distribución ocurrió al inicio de otoño, mientras que las condiciones oceanográficas cambiaron para la segunda semana de muestreo, cuando ocurrió la formación de una surgencia costera. Los eventos de apareamiento solo se registraron durante otoño para la tortuga golfina (\textit{Lepidochelys olivacea}), la especie dominante de la región. Se concluye que en otoño el patrón de distribución de tortugas marinas fue hacia la zona costera del PCM, y se relacionó con actividades de reproducción; mientras que en invierno y primavera este patrón tendió a la región central/oceánica vinculado potencialmente con actividades de forrajeo.

\textbf{Palabras clave}: Tortugas marinas, distribución, parámetros oceanográficos, giro ciclónico, surgencia

\textbf{Abstract}.- In this study we describe the spatial and temporal distribution of sea turtles and their association with oceanographic parameters in waters of the Mexican Central Pacific (MCP) during 2010 (winter, spring and autumn). Our results showed variations in sea turtle distribution through the sampling year. Sea turtle distribution was homogeneous in coastal and oceanic areas during winter; there was an association with thermal fronts generated by a current of cold water flowing from the northwest and a warm current coming from the south, as well as with the boundaries of a cyclonic gyre. Sea turtles were distributed in the coastal zone and the central part of the oceanic zone in spring, where the effects of a cyclonic gyre and coastal upwelling prevailed. The same distribution trend was recorded at the beginning of autumn, whereas oceanographic characteristics changed during the second sampling week, when upwelling occurred. Mating events were only recorded in autumn for olive ridley sea turtles (\textit{Lepidochelys olivacea}), the dominant species in the region. It was concluded that sea turtles were distributed in the coastal zone of MCP waters in autumn due to reproductive activities, whereas in winter and spring sea turtles were distributed towards the central/oceanic region, potentially related to foraging activities.

\textbf{Key words}: Sea turtles, distribution, oceanographic parameters, cyclonic gyre, upwelling

\textbf{INTRODUCTION}

Seven sea turtle species are currently recognized worldwide within the Cheloniidae and Dermochelyidae families (Márquez-Millán 1990, 1996). They have complex life cycles that encompass oceanic and coastal ecosystems, and their distribution is affected by biological and ecological factors (Frazier 2001, Bolten 2003). Sea turtle species spend most of their life in oceanic waters (mainly the East Pacific green, leatherback, and olive ridley turtles), with only nesting females spending significant time on land; however, nesting corresponds to only ~1% of their life cycle. Soon after nesting, females return to the sea where they, along with juveniles, adult males and adult non-nesting females will remain some time in oceanic areas as their principal habitat (Spotila et al. 1997).

Sea turtle species are protected by national and international environmental laws, and they are listed at different levels ranging from vulnerable to critically endangered in the IUCN red list;
therefore, several ecological parameters need to be investigated
to conclude on their conservation status (National Marine
Fisheries Service & U.S. Fish and Wildlife Service 1998). The
distribution of species is an ecological parameter that contributes
to the identification of zones and seasons that could render
individuals vulnerable to certain threats, such as fishery bycatch
and marine debris ingestion. The latter is considered a potential
disturbance agent for sea turtles in the coastal area of the MCP
(Díaz-Torres 2014).

The olive ridley turtle *Lepidochelys olivacea* (Eschscholtz,
1829), is the most abundant sea turtle worldwide (Márquez-
Millán 1990). Distributed along the Pacific, Atlantic and Indian
Oceans in tropical and subtropical areas (Pritchard 1969), the
olive ridley principally inhabits the oceanic zone during all life
stages (Bolten 2003, National Marine Fisheries Service & U.S.
Fish and Wildlife Service 1998), where it exploits multiple
foraging areas (Plotkin et al. 1994) that are influenced by the
presence of coastal upwelling, oceanic fronts and cyclonic gyres

Olive ridley turtles have been detected in areas of the Eastern
Tropical Pacific where oceanographic parameters are variable.
For example, Polovina et al. (2004) reported the presence of
these turtles in two different kinds of environments in the central
part of the Pacific Ocean adjacent to the Hawaiian islands: a)
one with low uniform Chlorophyll-α (Chl-α) concentrations and
weak current dynamics; and b) a second characterized by high
Chl-α concentrations and the presence of strong currents. In
contrast, Swimmer et al. (2009) found olive ridleys distributed
in areas characterized by relatively high superficial Chl-α
concentrations (0.37 mg m⁻³) and the presence of a cyclonic
gyre (Costa Rica Dome). Both studies also reported the
aggregation of olive ridley turtles in areas with warm surface
temperatures (25-28°C) (Polovina et al. 2004, Swimmer et
al. 2009). Therefore, the spatial and temporal distribution pattern
of this species is unclear, and may vary among regions.

The coastal region of the Mexican Central Pacific (MCP)
has been recognized as a migratory destination for olive ridley
turtles to carry out breeding activities during certain seasons
(Márquez-Millán 1990, Quijano-Scheggia et al. 2006). Most
research about sea turtle ecology is based on data collected on
nesting beaches; knowledge has therefore been obtained on
only one population component, mature females (Björndal
2000). For this reason, any information that encompasses the
entire population should be considered valuable.

There are no studies that relate oceanographic parameters
to sea turtle distribution in the MCP, although several
oceanographic processes (at different scales ranging from 10
to 100 km) occur in this region due to the interaction of different
water masses that converge in the area (Torres-Orozco et al.

The present study is considered the first in providing spatial
and temporal distribution patterns for sea turtles (undifferentiated
adults and juveniles) in MCP waters, using direct observations
during at-sea surveys (e.g., Houghton et al. 2006), linking these
distribution patterns with oceanographic parameters to
demonstrate a potential influence on the ecological activities
of sea turtles, such as breeding and feeding.

**Materials and methods**

**Study area**

The Mexican Central Pacific (MCP) encompasses 87,291 km²
between 16°48.68'-18°07.54'N and 102°5.85'-104°22.17'W.
The coastal limits were located to the north at Cabo Corrientes,
Jalisco, and to the south at Maruata, Michoacán, with a 180
km extension offshore. The area was divided in two zones
parallel to the mainland: coastal (0-40 km) and oceanic (40-
180 km). The coastal zone was determined by the width of the
continental shelf and slope (see Figure 2 and 3 to identify the
study area).

**Field work**

Three research surveys were conducted in 2010 on board the
fisheries research vessel *BIP XII*: 1) winter (15-27 January),
2) spring (25 May-6 June), and 3) autumn (18-29 October).
We employed linear transect sampling during daylight hours
(7:00-19:00 h) at an average speed of 8-9 knots under favorable
sea state conditions (Beaufort scale (*B* = 0-3) to guarantee
sea turtle detection. Transects were systematically designed as
part of a study of sea turtle and marine mammal monitoring;
however, vessel routes varied depending on logistic and
environmental factors.

Sea turtle sightings were recorded by 3 observers located
on the highest ship platform (6.7 m above sea level) using Fujinon
7x50 binoculars and facing in different transect directions. The
date, hour, sea state (*B* scale), ship course, number of sea turtles,
and reticle number and angle (data supplied by binoculars) were
recorded for each sighting. The geographic position of the ship
course and observed turtles were recorded using a global
positioning system Garmin GPSmap 76cs. If the sea turtle was
close to the ship’s transect line, its species and sex (mature
males have longer and thicker tails; Márquez-Millán 1990,
1996) were recorded, and a photograph was taken with a
Canon digital camera model EOS-5D to corroborate or
identify the species from its morphological features, using the
In addition, mating events (a pair of turtles joined with the male above the female) were recorded during these surveys.

**Sea turtle density calculations**

To analyze the spatial and temporal trends of sea turtle distribution, and to reduce the bias caused by differences in effort among the zones, the density \( D \) of organisms along each transect was calculated as:

\[
D_i = \frac{n_i}{2W_iL_i}
\]

where: \( n \) is the number of sea turtles recorded along transect \( i \), \( W \) is the width and \( L \) the length of transect \( i \).

The length of the transect \( L \) is the distance traveled (in km) from the point where the observation effort started to the point where it stopped, considering the following conditions to discriminate among transects: 1) decrease in speed or lack of navigation, 2) change of course, and 3) change of zone (transition between coastal and oceanic areas).

The width of each transect \( W \) was equivalent to the maximum perpendicular distance (with respect to the ship’s course; in km) at which a turtle was observed. Its estimation was obtained by incorporating the reticle number and angle in the equations proposed by Jaramillo-Legorreta et al. (1999). This variable \( W \) could be influenced by the sea state \( B \) scale observed during each transect, i.e., at a sea state \( B \leq 1 \), there was a higher probability of sighting a turtle at a greater distance than at transects with \( B \geq 3 \). We therefore obtained a standard \( W \) for each \( B \) scale \( (0, 1, 2, \text{ and } 3) \) for each zone (coastal and oceanic), established by the maximum \( W \) value during all transects with a similar \( B \) scale value. The corresponding \( W \) was used to calculate turtle density based on the \( B \) scale at which each transect occurred.

Only sightings corresponding to the olive ridley sea turtle \( (L. \text{ olivacea}) \) were taken into account for density analyses, as well as sightings of turtles that did not show clear morphological characteristics typical of other species, e.g., large size and longitudinal edges of the shell \( (Dermochelys \text{ coriacea}) \), darker coloration and flatter shell \( (Chelonia \text{ mydas}) \), lighter coloration and shell keels \( (Eretmochelys \text{ imbricata}) \) (Márquez-Millán, 1990). The assumption that these sightings could correspond to olive ridley turtles is supported by the great abundance of this species in adjacent waters of the Mexican Tropical Pacific (Eguchi et al. 2007), as well as along nesting beaches in the area (Quijano-Scheggia et al. 2006). Observations of sea turtles for which it was not possible to record reticle or angle value were omitted from density calculations and statistical analyses. To determine whether there were significant differences in densities among seasons and zones, we used non-parametric Chi-square tests \( (X^2) \) with \( \alpha \leq 0.05 \) significance, using the software *STATISTICA* ver. 7.0.

**Sea Surface Temperature and Chlorophyll-α analyses**

Sea Surface Temperature (SST) and Chlorophyll-α (Chl-α) data were obtained from the Aqua Modis (EOS PM) sensor from NASA (NASA’s Ocean Biology Processing Group, OBPG)$^1$, with 4 km spatial resolution. We used images of 8-day composite periods, corresponding to the weeks during which each survey was conducted. Data image processing was carried out using *R* ver. 2.14.1 software (Villalobos & González-Rodríguez 2010).

To describe the spatial-temporal distribution of sea turtles, we plotted weekly maps of each oceanographic parameter, of sea turtle sightings, and of sampling transects onto a map, using *MATLAB* ver. 7.0 software. We also included geostrophic current data from NOAA: Near Real-Time Altimeter page, calculated from climatic values and sea high anomalies from the Naval Research Laboratory (NRL) (Trinanes 2001).

We performed a correlation analysis \( (Beta, \alpha \leq 0.05) \) between sea turtle densities and SST and Chl-α, by 1) identifying the values of these oceanographic parameters in correspondence to each sea turtle position using the program *WAM* (WIM Automation Module)$^2$; and 2) calculating the average of oceanographic parameters in each line transect to correlate with sea turtle density for the same line transect. Analyses were run using *STATISTICA* ver. 7.0 software.

**Results**

**Sea turtle density**

Sea turtle distribution was homogeneous throughout the study area (coastal: 0.07 turtles km\(^{-2}\); oceanic: 0.05 turtles km\(^{-2}\)) \( (X^2 = 2.57, \alpha = 0.11, n = 97) \) in winter (Fig. 1; Table 1). We identified more olive ridley sea turtles (21 confirmed sightings) than any other species: East Pacific green turtle \( (Chelonia \text{ mydas}) \) (4 sightings), loggerhead \( (Caretta caretta) \) (10 sightings), leatherback \( (Dermochelys \text{ coriacea}) \) (2 sightings), and hawksbill turtles \( (Eretmochelys \text{ imbricata}) \) (2 sightings). The total number of unidentified turtles was 358 (Table 2).

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$^1$[http://www.oceancolor.gsfc.nasa.gov/cgi/]

$^2$[http://www.wimsoft.com]
Sea turtle density in the coastal area (0.53 turtles km$^{-2}$) was significantly greater than density in the oceanic area (0.07 turtles km$^{-2}$) ($X^2 = 5.81, \alpha = 0.01, n = 119$) during spring (Fig. 1; Table 1). Only two species could be identified: olive ridley, which was the most observed (5 sightings) and East Pacific green (1 sighting) (Table 2).

Density in the coastal zone (0.73 turtles km$^{-2}$) was significantly greater than in the oceanic zone (0.06 turtles km$^{-2}$) ($X^2 = 5.95, \alpha = 0.01, n = 84$) during autumn (Fig. 1; Table 1). Two species could be identified: olive ridley turtle, which was the most frequent (12 sightings), and East Pacific green (7 sightings). Mating events were only recorded for olive ridley turtles, and occurred only during autumn ($n = 11$; Table 2).

![Table 1. Spatial and temporal sea turtle density in coastal and oceanic zones of the Mexican Central Pacific during three seasons in 2010 / Densidad espacial y temporal de tortugas marinas en zonas costeras y oceánicas del Pacífico Central Mexicano durante tres temporadas del 2010](image)

**Table 1. Spatial and temporal sea turtle density in coastal and oceanic zones of the Mexican Central Pacific during three seasons in 2010 / Densidad espacial y temporal de tortugas marinas en zonas costeras y oceánicas del Pacífico Central Mexicano durante tres temporadas del 2010**

<table>
<thead>
<tr>
<th>Season</th>
<th>Stratum</th>
<th>E (km)</th>
<th>T</th>
<th>S.T.</th>
<th>Density (turtles km$^{-2}$)</th>
<th>Chi-square ($X^2_{df, n}$)</th>
<th>P</th>
<th>Density (turtles km$^{-2}$)</th>
<th>Chi-square ($X^2_{df, n}$)</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Winter</td>
<td>Coastal</td>
<td>434</td>
<td>30</td>
<td>26 (1)</td>
<td>0.07 ± 0.11</td>
<td>$X^2_{(1, 97)} = 2.57$</td>
<td>0.11</td>
<td>0.05 ± 0.09</td>
<td>$X^2_{(1, 97)} = 0.75$</td>
<td>0.39</td>
</tr>
<tr>
<td></td>
<td>Oceanic</td>
<td>955.9</td>
<td>67</td>
<td>348 (4)</td>
<td>0.07 ± 0.09</td>
<td></td>
<td></td>
<td>0.06 ± 0.10</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Spring</td>
<td>Coastal</td>
<td>255.2</td>
<td>31</td>
<td>88 (13)</td>
<td>0.53 ± 0.77</td>
<td>$X^2_{(1, 119)} = 5.82$</td>
<td>0.01</td>
<td>0.19 ± 0.46</td>
<td>$X^2_{(2, 309)} = 3.55$</td>
<td>0.17</td>
</tr>
<tr>
<td></td>
<td>Oceanic</td>
<td>1,143</td>
<td>88</td>
<td>108 (1)</td>
<td>0.07 ± 0.15</td>
<td></td>
<td></td>
<td>0.32 ± 1.47</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Autumn</td>
<td>Coastal</td>
<td>435.2</td>
<td>30</td>
<td>248</td>
<td>0.73 ± 2.37</td>
<td>$X^2_{(1, 84)} = 5.96$</td>
<td>0.17</td>
<td>0.32 ± 1.47</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Oceanic</td>
<td>956.6</td>
<td>54</td>
<td>56 (8)</td>
<td>0.06 ± 0.28</td>
<td></td>
<td></td>
<td>0.32 ± 1.47</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

E = effort; T = number of transects; S.T. = number of sighted turtles considered for density analysis, between parenthesis number of turtles excluded from the statistical analysis.
On a temporal scale, sea turtle density was highest during autumn (0.32 turtles km$^{-2}$), followed by spring (0.19 turtles km$^{-2}$), and winter (0.06 turtles km$^{-2}$); these differences were not significant, however ($X^2=3.56$, $\alpha=0.17$, $n=300$) (Table 1).

**DISTRIBUTION OF SEA TURTLES RELATED TO OCEANOGRAPHIC PARAMETERS**

Slightly colder water (25.5-26.4°C) was detected during winter, flowing from the northwest into the oceanic zone, promoting the formation of thermal fronts (26-27.3°C) where small groups of sea turtles were found (Fig. 2A1). In addition, a warm current (27.2-28.9°C) from the south was also identified, extending through the coastal zone and the southern part of the oceanic area (Fig. 2A1). Temperature values associated with sea turtles within this warm current were higher (27.4-28.9°C) in comparison with that for thermal fronts (Fig. 2A1), whereas Chl-a concentrations were low (0.16-0.25 mg m$^{-3}$), but similar to values reported for turtles within the cold intrusion water (0.17-0.24 mg m$^{-3}$) (Fig. 3A1). The correlation between sea turtle density and SST was not significant ($\alpha=0.03$, $\alpha=0.59$), nor was it significant for Chl-a ($\alpha=0.04$, $\alpha=0.84$).

A cyclonic gyre was detected (17.6-18.7°N and 104.6-105.9°W) during the second week of winter; it produced a transition zone in the oceanic area, where SST ranged from 26.2 to 28.5°C and Chl-a ranged from 0.2 to 1.4 mg m$^{-3}$ (Fig. 2A2). Sea turtles were distributed within the transition zone generated by the interaction of the cyclonic gyre and the remnant of cold water intrusion. The SST ranged from 26.5 to 27.6°C (Fig. 2A2), whereas Chl-a ranged from 0.18 to 0.63 mg m$^{-3}$ (Fig. 3A2). In contrast, turtles recorded in the coastal zone were located in warmer (27-28.5°C), and more productive (0.15-2.5 mg m$^{-3}$) waters (Figs. 2 and 3A2). The correlation between sea turtle density and the oceanographic parameters was not significant (SST: $\alpha=0.008$, $\alpha=0.71$; Chl-a: $\alpha=0.05$, $\alpha=0.36$).

Oceanographic characteristics recorded during two weeks in spring suggested the presence of coastal upwelling off the Jalisco coast, characterized by SST between 24.1 and 25.6°C (Fig. 2B1-B2) and Chl-a concentrations between 0.57 and 10 mg m$^{-3}$ (Fig. 3B1-B2). Sea turtles within this area were associated with the boundaries of the upwelling, where high Chl-a gradients were found (0.14-1.8 mg m$^{-3}$) (Fig. 3B1); however, these values were lower (0.12-0.48 mg m$^{-3}$) during the second week (Fig. 3B2). In addition, a cyclonic gyre off Colima state (17.8-18.9°N and 104.3-105.7°W), was detected during the two sampling weeks. This gyre resulted in Chl-a concentrations between 0.15 and 0.63 mg m$^{-3}$ (Figs. 3B1-B2) and SST ranging from 25.9 to 28°C (Figs. 2B1-B2). Although the distribution pattern of sea turtles was associated with the boundaries of this gyre (Figs. 2 and 3B1-B2), the correlation of sea turtle density with SST and Chl-a during the first week (SST: $\alpha=0.03$, $\alpha=0.37$; Chl-a: $\alpha=0.08$, $\alpha=0.10$) or second week (SST: $\alpha=0.03$, $\alpha=0.47$; Chl-a: $\alpha=0.03$, $\alpha=0.45$) was not significant.

Warmer SST was recorded throughout the entire study area during the first sampling week of autumn (27.3-30°C). Sea turtles density was highest near the coast, and sea turtles aggregated in two large groups: one near the northern part of Jalisco state (19.5-20°N) with a relative density of 1.44 turtles km$^{-2}$, and the other near the southern part of Colima and northern Michoacán (18.2-19°N), with a relative density of 0.46 turtles km$^{-2}$. The SST associated with sea turtle positions ranged from 27.4 to 30°C (Fig. 2C1), whereas the Chl-a observed over sea turtle aggregations reflected more productive waters (0.41-10 mg m$^{-3}$), compared to turtles in the oceanic zone (0.15-0.3 mg m$^{-3}$) (Fig. 3C1). However, the correlation analysis between sea turtle density and SST and Chl-a was not significant (SST: $\alpha=0.02$, $\alpha=0.55$; Chl-a: $\alpha=0.0004$, $\alpha=0.92$).

We observed coastal upwelling in the second week (SST: 24.1-25.5°C; Chl-a: 0.75-10 mg m$^{-3}$), extending from Cabo
Corrientes, Jalisco, to the southern part of the study area. This upwelling was also confirmed by the dynamics of the geostrophic currents (Figs. 2 and 3C2). The sea turtles were located over the warm part of the oceanic zone, with an associated SST of 27.4-28.3°C (Fig. 2-C2) and low Chl-a concentration (0.18-0.37 mg m⁻³), except for a small turtle aggregation (17.4-17.6°N and 103.8-104.6°W), located in a Chl-a front area (0.32-1.17 mg m⁻³) (Fig. 3-C2). Sea turtles in the Colima coastal area were located in slightly cooler (26.8-28°C) (Fig. 2-C2) but more productive waters (0.72-2.3 mg m⁻³) (Fig. 3-C2). The correlation between sea turtle density and both variables was not significant (SST: Beta= 0.10, α= 0.24; Chl-a: Beta= 0.05, α= 0.38).


DISCUSSION
This study presents calculations of sea turtle density in waters of the MCP, as well as oceanographic features of the region that could potentially explain the observed spatial-temporal distribution pattern.

Most of the sea turtles observed during this study (~93%) were labeled as unidentified species, because we were no able to detect morphological characteristics particular to each species, e.g., large size and longitudinal shell edges for D. coriacea, darker coloration and flatter shell for C. mydas, lighter coloration and shell keels for E. imbricata (Márquez-Millán 1990). It is however possible that most of them corresponded to the olive ridley sea turtle (L. olivacea), which was the most sighted species in the MCP (38 confirmed identifications) compared with the other four identified species (26 identifications, Table 2). In addition, this species is the most abundant in waters of the Mexican Tropical Pacific and Central America (X= 1.38 million, CV= 19.7%, CI: 1.15-1.62 million) (Eguchi et al. 2007), and the nest and hatchlings reports released by sea turtle camps on the Jalisco, Colima and Michoacán coasts have also demonstrated the dominance of this species (Quijano-Scheppia et al. 2006, Trejo-Robles et al. 2006, Comisión Nacional de Áreas Naturales Protegidas & Secretaría de Medio Ambiente y Recursos Naturales 2009). It is important to keep in mind that most sightings recorded during this study in 2010 probably corresponded to the olive ridley sea turtle, but this is only an assumption and for this reason we will refer to them below as ‘sea turtles’.

The distribution trends of sea turtles in MCP waters showed clear spatial variations, perhaps related to particular biological and ecological aspects, such as feeding (in winter-spring) and breeding (in autumn) (Plotkin et al. 1994, National Marine Fisheries Service & U.S. Fish and Wildlife Service 1998, Frazier & Secretaria de Medio Ambiente y Recursos Naturales 2009). This study presents calculations of sea turtle density in waters of the Mexican Coastal Current (Kessler 2006). Oceanic fronts are highly productive systems due to the interaction of several water masses with different densities. They promote the formation of convergence and vertical divergence zones that favor the accumulation of organic/inorganic material, enriching biological productivity at different trophic levels (Font et al. 1987, Lalli & Parsons 1997, Miller 2011). This process could have occurred in the oceanic zone during the second week, where more productive waters were located (0.2-1.4 mg m⁻³), and where turtle aggregations were observed (Fig. 3-A2).

Pelayo-Martínez (2013) analyzed the in situ response of nutrients, Chlorophyll-α, and zooplankton biomass versus the oceanographic variability in the oceanic/central region of MCP waters during winter, revealing the presence of Euphausiacea, Decapoda and Amphipoda crustacean groups, as well as gelatinous species of the Doliolida group (pyrosomes and salps) (volume: 14-22 ml 1000⁻³). Foraging activity may have been occurring in the region, as these organisms are important components of the olive ridley diet (Montenegro-Silva et al. 1986, National Marine Fisheries Service & U.S. Fish and Wildlife Service 1998, Polovina et al. 2004).

Sea turtle distribution was observed in the coastal zone and the central part of the oceanic area in spring, where a cyclonic gyre again prevailed (Figs. 2 and 3B1-B2), confirming its semi-permanence in the region as previously had been reported (Salas-Pérez et al. 2006). The tendency of sea turtles to aggregate in cyclonic gyre areas (Luschi et al. 2003, Swimmer et al. 2009) has been explained by the oceanographic characteristics of cyclonic gyres (shallow thermocline, wide gradients of Chl-α and nutrients), which stimulate enrichment of the trophic chain and the presence of potential prey (Lalli & Parsons 1997). Coastal upwelling was also detected off the Jalisco coast (Figs. 2B1-B2). It resulted in high Chl-α concentration (0.57-10 mg m⁻³) distributed over the area due to the presence of the cyclonic gyre, and favoring the formation of frontal zones (Figs. 3B1-B2). Previous studies have indicated that coastal upwelling off the Jalisco coast is a cyclic phenomenon that is due mainly to the wind influence (Torres-Orozco et al. 2005, López-Sandoval et al. 2009); a period of upwelling intensification during spring has also been noted. According to the wind pattern from the CoastWatch software of NOAA¹, it was noted that winds blew parallel to the continent mainly in the west-northwest direction during the spring sampling period, favoring the formation of upwelling due to Ekman transport (Torres-Orozco et al. 2005, Stewart 2008, López-Sandoval et al. 2009).

¹<http://coastwatch.pfeg.noaa.gov/erddap/griddap/erdQAstress1day.graph>
Sea turtles occurred in frontal zones where SST varied between 25.9 and 28°C (Figs. 2B1-B2), in slightly productive waters (0.12–1.8 mg m⁻³) (Figs. 3B1-B2), where the principal organisms in the olive ridley turtle diet have also been reported (Pelayo-Martínez 2013). Therefore, we infer that the presence of sea turtles near these oceanographic structures was related to foraging activities, as has been reported by other authors for the Tropical Pacific Ocean (Frazier 2001, Luschi et al. 2003, Polovina et al. 2004, Swimmer et al. 2009). This behavior may be influenced by the narrow continental shelf and by the 4,000 m deep canyon that lies near the coast in the study area (De la Lanza 1991), which leads sea turtles (such as olive ridley turtles) to forage in the mid-water column, and not in benthic areas, as has been proposed based on satellite tracking (McMahon et al. 2007).

Sea turtles were distributed throughout the coastal zone in autumn. This could be influenced by breeding activities such as mating and nesting. Olive ridley turtles reproduce from July to January (Márquez-Millán 1990) with a peak in August and September (Aguilar-Olguín et al. 2006, Trejo-Robles et al. 2006). This coincides with the sightings of mating turtles that were recorded only during the autumn survey (Table 2). Most mating events occur before nesting peaks (Márquez-Millán 1990, Godley et al. 2002). As we were not able to conduct a survey before August-September, when we would have expected a higher count of mating events, we cannot provide an exhaustive discussion.

Nesting cycles of olive ridley turtles occur every 14 days during solitary events, and every 28 days during synchronized nesting events, termed ‘arribadas’; ridley turtles can therefore nest approximately three times per season (Plotkin 2003, Abreu-Grobois & Plotkin 2008). Thus, the highest density recorded during autumn (0.32 turtles km⁻²) (Table 1) can be associated with the arrival of olive ridley turtles to mate and nest in the area, which is supported by the previously mentioned numerous beaching arrivals of olive ridley sea turtles to conservation camps.

The presence of upwelling off Cabo Corrientes, Jalisco, was also detected in autumn, favoring the enrichment of biological productivity in the coastal zone and part of the oceanic zone (0.75–10 mg m⁻³) (Figs. 3C1-C2). This could have favored the increase of potential prey for sea turtles, allowing them to feed during their stay near the coast (Lalli & Parsons 1997, Stewart 2008). López-Sandoval et al. (2009) pointed out that Cabo Corrientes is characterized by coastal upwellings all year long, which tend to occur during the relaxation phase that occurs from July to December. The wind recorded during autumn had a north-northwest direction, also blowing parallel to the coast (NOAA)⁴. This favored the development of coastal upwelling due to Ekman transport (Stewart 2008), also confirmed by the geostrophic current dynamics (Figs. 2 and 3C2).

Integrating the information, we can conclude that sea turtle aggregations, for the most part of olive ridley turtles, were recorded in the MCP region during winter-spring, when we would not have expected records of sea turtles due to their migratory behavior. Sea turtle aggregations were found in oceanic areas where potential conditions appropriate for feeding could occur. Autumn aggregations occurred in the coastal zone and were associated chiefly with breeding activities.

Given this result, a potential inference could be established: that these sea turtles could not travel to distant locations, and their distribution pattern could have been influenced over a small spatial scale in the MCP, where oceanographic conditions favored their main ecological activities, such as feeding and breeding (Kernohan et al. 2001). Nevertheless, it should be taken into account that 2010 was an atypical year, because during the first months of the year a warmer temperature anomaly prevailed (‘El Niño’), and during last months of the year the anomaly temperature was colder (NOAA)⁵, and changes in turtle migration patterns in response to ‘El Niño’ were reported (Plotkin 2010). Therefore, we recommend that a similar study be conducted during a typical year to elucidate the distribution pattern of sea turtles as a function of oceanographic parameters. This ecological aspect could contribute substantial information to the knowledge of the ecology of these species, and contribute useful tools for adequate species management, because at present, all sea turtle species are considered in the IUCN (2013) red list.

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⁴<http://coastwatch.pfeg.noaa.gov/erddap/griddap/erdQAstress1day.graph>

⁵<http://www.cpc.ncep.noaa.gov/products/precip/CWlink/MJO/enso.shtml>
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