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Long Term Dinoflagellate Bioluminescence, Chlorophyll, and Their Environmental Correlates in Southern California Coastal Waters

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1. Introduction

While many oceanographic studies have focused on the distribution of bioluminescence in the marine environment (Stukalin 1934, Tarasov 1956, Seliger et al. 1961, Clarke and Kelly 1965, Bityukov 1967, Lapota and Losee 1984, Swift et al. 1985, Lapota et al. 1988, Batchelder and Swift 1989, Lapota et al. 1989, Lapota and Rosenberger 1990, Neilson et al. 1995, Ondercin et al. 1995, Swift et al. 1995), little understanding of the seasonality and sources of planktonic bioluminescence in coastal waters and open ocean has emerged. Some previous studies with respect to annual cycles of bioluminescence were severely limited in duration as well as in the methods used to quantify bioluminescence (Bityukov 1967, Tett 1971). Only a few studies have measured bioluminescence on an extended basis, and these were short in duration, usually less than 2 years with long intervals between sets of measurements (Bityukov 1967, Yentsch and Laird 1968, Tett 1971). Others report data collected at different times of the year (Batchelder and Swift 1989, Batchelder et al. 1992, Buskey 1991) but do not address the seasonality of bioluminescence. Thus the detailed temporal variability of bioluminescence has never been characterized continuously over several years. Lack of such long-term studies leaves unanswered important questions regarding the role of bioluminescence in successional phenomena.

To adequately understand, model, and predict planktonic bioluminescence in any ocean, measurements must be conducted on a continual basis for at least several years in order to evaluate intra- and annual variability and long-term trends. In this study, bioluminescence was measured at two fixed stations on a daily long term basis: one in San Diego Bay (SDB) for 4 years (1992-1996) and the other for 2.5 years (1993-1996) at San Clemente Island (SCI), located 100 km off the California coast. Additional surface and at-depth bioluminescence data have been collected on a monthly and quarterly basis at both fixed stations and from a research vessel to provide a link between coastal and offshore waters. Additional factors such as seawater temperature, salinity, beam attenuation, and chlorophyll fluorescence were measured. Plankton collections were made weekly in SDB and monthly at SCI. This study provides unique correlated coastal and open ocean data collected on a long-term basis (Figure 1).
2. Methods and materials

2.1 Bioluminescence measurements

Two defined excitation moored bathyphotometers (MOORDEX, University of California, Santa Barbara) were used in San Diego Bay (SDB) and at San Clemente Island (SCI). Under control of on-board computers, these measured stimulated bioluminescence, flow rate, and seawater temperature hourly. Every hour, seawater was pumped for 120 sec at 7-8 L/sec for a total volume of approximately 840 - 960 L of seawater through a darkened cylindrical 5 l detection chamber approximately 406 mm long and 127 mm in diameter (Case et al. 1993, Neilson et al. 1995). Bioluminescence, excited by the chamber spanning input impeller, was measured by a PMT receiving light from 46 fiber optics tips lining the chamber wall and expressed as photons sec\(^{-1}\) ml\(^{-1}\) of seawater.

On monthly transits between SDB and SCI an "on-board" sensor system sampled seawater continuously from 3m below the sea surface from a 50m research vessel, the R/V Acoustic Explorer, measuring bioluminescence, seawater temperature, and salinity (Lapota and Losee 1984, Lapota et al. 1988, 1989). A vertically deployed bathyphotometer capable of measuring bioluminescence, temperature, salinity, beam attenuation, and chlorophyll fluorescence to a depth of 100m was used at 4 month intervals (summer, fall, winter, spring) at various stations in the Bight to examine the seasonal changes in the biological and physical structure of the water column (Lapota et al. 1989). Both systems were calibrated with the luminescent bacteria *Vibrio harveyii* in a Quantalum 2000 silicon-photodiode detector. The detector calibration is traceable to a luminal light standard (Matheson et al. 1984).

2.2 Plankton and seawater analysis

Water and plankton samples were collected at 10 stations within the Bight (Figure 1). Monthly transits were made from March 1994 through June 1996 from SCI to SDB to measure surface (3m depth) bioluminescence and collect plankton and seawater samples to determine Chl \(a\) content. At SDB, weekly plankton and water samples were taken for 4 years while monthly plankton and water samples were collected at SCI for 2.5 years. Because plankton abundance within SDB is usually high, 10 L water samples were concentrated while 40 l samples were filtered for plankton at SCI. Fifteen-liter water samples were collected and filtered from select bathyphotometer depths on the quarterly stations (10, 20, 30, 40, 50, 70, and 90 m). This was accomplished by discharging the bathyphotometer's effluent from its submersible pump through a 130-m long, 2.54 cm (I.D.) hose into a 15 liter Imhoff settling cone. The bottom of the cone was modified with a valve that allowed water to be filtered into collection cups fitted with 20-µm porosity netting. One liter of seawater (unfiltered) was also collected at the each of these depths and frozen in precleaned polycarbonate bottles for later chlorophyll and nutrient analysis. Plankton samples were preserved in a 5% formalin seawater solution. Bioluminescent dinoflagellates were identified to the species level when possible. Chlorophyll \(a\) was extracted from the seawater samples using standard methods (APHA 1981) and measured by fluorescence as an estimate of biomass on a Turner Model 112 fluorometer (Sequoia-Turner Corp., Mountain View, CA, U.S.A.) and reported as µg L\(^{-1}\).
2.3 Upwelling, rainfall, and seawater nutrient data bases

Upwelling indices (North Pacific Ocean wind-driven transports) were collected from 1992 through 1996. The indices were computed for 33°N latitude (Schwing et al. 1996) and represent monthly average surface pressure data in cubic meters per second along each 100 m of coastline (Bakun 1973, Eppley 1986). Monthly rainfall data were acquired from the National Weather Service in San Diego. Nutrient and Chl a data were accessed from archived CALCOFI data (1992-1996) in the Bight and were averaged along CALCOFI lines 90 and 93 which run west from San Diego to the north and south of San Clemente Island (Hayward et al. 1996). Nitrates (µm L⁻¹) and Chl a (µg L⁻¹) along each of the CALCOFI transit lines (stations 93-26 to 93.45 and 90-28 to 90.53) were averaged from the surface to a depth of 50m for 12 cruises conducted from September 1992 through April 1995. These data were used to calculate correlations with bioluminescence, rainfall, and upwelling at SDB.

3. Results

3.1 Mean monthly bioluminescence

Hourly bioluminescence data were averaged for each month. Because minimal bioluminescence was measured during daylight hours, mean monthly values were based on data collected from 2100 h (9:00 P.M.) to 0300 h (3 A.M.) the following day.

Seasonal changes in bioluminescence were observed in SDB. Maximum bioluminescence (1 x 10⁸ photons s⁻¹ ml⁻¹ or greater as a threshold) was measured from March through September for 1993, May through June for 1994, December through May for 1995, and March through April 1996. Minimum bioluminescence (less than 1 x 10⁸ photons s⁻¹ ml⁻¹)
was measured in January through February for 1993, December through February for 1993-94, November for 1994-95, and January through February for 1996 (Figure 2).

Fig. 2. Mean monthly bioluminescence trends at San Diego Bay and San Clemente Island from 1992-1996.

A red tide of the bioluminescent photosynthetic dinoflagellate, *Gonyaulax polyedra*, developed in the winter of 1994 along the southern California coast and was correlated with an increase in bioluminescence in SDB in December and later at SCI in January through April 1995 (Figure 2). *Noctiluca miliaris* appeared in the plankton collections at SCI following this bloom and produced 57% of the bioluminescence in May 1995. Bioluminescence decreased in SDB during June-July 1995. Mean monthly minimum and maximum bioluminescence in SDB ranged from $1 \times 10^6$ (February 1993) - $4.5 \times 10^8$ (June 1993) photons sec$^{-1}$ ml$^{-1}$. Additionally, during the red tide, bioluminescence averaged $1-2 \times 10^8$ photons s$^{-1}$ ml$^{-1}$ from December through April 1995, a factor of 10 above the normal measured winter bioluminescence.

In contrast, mean monthly bioluminescence at SCI varied little from August 1993-February 1996 except during the red tide in January 1995 ($2 \times 10^8$ photons s$^{-1}$ ml$^{-1}$) and persisted through April (Figure 2). Mean monthly bioluminescence ranged from $8 \times 10^6$ - $3 \times 10^7$ photons s$^{-1}$ ml$^{-1}$ at SCI.

### 3.2 Bioluminescent plankton - San Diego Bay

Most bioluminescence in SDB and SCI was emitted by the photosynthetic dinoflagellates *G. polyedra, Ceratium fusus, Pyrocystis noctiluca* as well as from the heterotrophic dinoflagellate, *Noctiluca miliaris*, and several species of *Protoperidinium*. Within SDB, maximum numbers of bioluminescent dinoflagellates ($2430 - 17,216$ cells L$^{-1}$) were collected during the spring-summer months while minimal numbers ($3 - 2,110$ cells L$^{-1}$) were usually collected in the winter months for 1992, 1993, and 1995 (Figure 3a).
In most months bioluminescent dinoflagellates represented a substantial percentage of total dinoflagellates (luminous and non-luminous species) (Figure 3b). Of these, *G. polyedra* and *Protoperidinium* spp. were most abundant; found in the winter, spring, and early summer months in SDB (Figure 4). *Gonyaulax polyedra* contributed more than 80% of all luminescent cells from early summer 1993 (Figure 3b). *Ceratium fusus* contributed minimally to the total number of dinoflagellates for the summers of 1993-1994 (Figure 4a). A shift was observed in the dinoflagellate species composition within SDB with *G. polyedra* becoming the dominant species. There were fewer *Protoperidinium* spp. and *C. fusus* at SDB in 1994 (Figure 4).
A light budget was calculated to estimate the contribution of light produced by the various species of bioluminescent dinoflagellates in SDB. Light output from each was measured with a laboratory photometer system by stirring individual cells for 30 sec (Lapota et al. 1989, Lapota et al. 1992). Mean light output for each species was calculated and then multiplied by the number of cells found in each of the monthly plankton samples. The mean light output values for single cells were: *Gonyaulax polyedra* $1 \times 10^8$ photons; *Ceratium fusus* $2 \times 10^8$ photons, *Protoperidinium* spp. $3 \times 10^9$ photons, *P. noctiluca* $1 \times 10^{10}$ photons, and *N. miliaris* $2 \times 10^{10}$ photons. Photons L$^{-1}$ for each group were plotted from monthly samples collected in SDB (Figure 5). Bioluminescence from each of the groups (% of total bioluminescence) was then estimated (Figure 5). *Protoperidinium* dinoflagellates contributed more than 80% of the bioluminescence in 41% of all months ($n = 51$ months) and more than 50% of the
bioluminescence in 73% of all months. In contrast, *G. polyedra* contributed more than 80% of bioluminescence in SDB in only 2% of all months and more than 50% of all bioluminescence in 18% of all months. *Gonyaulax polyedra* bioluminescence was most pronounced in the fall and winter months. Peaks in *C. fusus* bioluminescence were most pronounced in late summer and fall, however, the contribution to the light budget was minimal. In 1995 and 1996, *G. polyedra* dominance in the winter months was followed by an increase in bioluminescence from *N. miliaris* which attained concentrations of 95 cells L\(^{-1}\) in March 1995, 186 cells L\(^{-1}\) in April 1995, and 23 cells L\(^{-1}\) in May 1995. This same trend and similar cell numbers were encountered in the spring months of 1996.

Fig. 5. Bioluminescence produced by each species (photons L\(^{-1}\)) monthly in San Diego Bay from 1992-1996.

### 3.3 Bioluminescent plankton - San Clemente Island

Numbers of luminescent dinoflagellates were lower at SCI than at SDB, ranging from 3 - 211 cells L\(^{-1}\) of seawater from August 1993 through December 1994 (Figure 6). The principal species were *G. polyedra* and several species of *Protoperidinium*. The red tide was first observed in January 1995 and persisted through April 1995. Bioluminescence during this event increased approximately 10 times above former levels for both SDB and SCI, although this difference was measured at SDB one month earlier than SCI (Figure 2). Total dinoflagellates and bioluminescent dinoflagellates increased to 16,727 cells L\(^{-1}\) and 15,939 cells L\(^{-1}\), respectively at SCI in January 1995 (Figure 7a). Cell numbers remained high through April 1995. *Gonyaulax polyedra* was the predominant red tide bioluminescent dinoflagellate, however several species of *Protoperidinium* increased to numbers as high as 674 cells L\(^{-1}\) in February 1995 (Figure 7a). At SCI, bioluminescent dinoflagellates represented a major percentage of all dinoflagellates collected (Figure 7b). The light budget analysis indicated that the species of *Protoperidinium*, again, produced most of the bioluminescence, followed by *Gonyaulax* and *Ceratium* species (Figures 8a,8b). At SCI, *Protoperidinium* contributed more than 80% of all bioluminescence for 60% of all months (n = 30 months) and...
more than 50% of all bioluminescence for 77% of all months. In contrast, *Gonyaulax* contributed 80% of all bioluminescence for just 1 month (3.3% of all months) and 50% of all bioluminescence for only 10% of the months. During the red tide encountered in the winter and spring of 1995, *G. polyedra* contributed 59%, 42%, 58%, 48%, and 27% of all bioluminescence for the months of January through May 1995, respectively (Figure 8b). As in SDB, *N. miliaris* appeared (~2 cells L⁻¹) following the bloom of *G. polyedra* and produced 57% of the bioluminescence in May 1995 (Figure 8b). The open ocean bioluminescent dinoflagellate, *Pyrocystis noctiluca*, was also found in monthly collections. *Protoperidinium* spp. were present in greater numbers in the spring and summer months; while *G. polyedra* became more prevalent in the fall and winter months.

The light budget analysis (photons L⁻¹) at SDB and SCI correlated with measured bioluminescence on a daily (photons ml⁻¹ day⁻¹) and monthly (photons ml⁻¹ month⁻¹) basis, although the light budgets were highly correlated with the later. The light budget analysis reinforces our understanding of which bioluminescent species contributed to measured bioluminescence. SCI had the higher correlations ($r = 0.876; p < 0.001$). This may reflect a more constant plankton assemblage over time in contrast to a more variable bay environment where tidal flow into and out of the bay may cause more variation in the populations sampled (Figure 8)

The change in dinoflagellate species composition between summer and fall is well shown in the monthly transits where surface (3-m depth) plankton samples were collected at 10 stations each month from July through October 1994 (Figure 9). We observed a gradual shift in the ratio of *Protoperidinium* spp. to *G. polyedra* between September and October (Figure 9). This trend was also observed in bathyphotometer profiles (0-90 m) for the July and November 1994 stations across the Bight (Figure 10), with *G. polyedra* in both instances dominating in the winter months.
Fig. 7. (a) Total and bioluminescent dinoflagellate cells collected monthly at San Clemente Island from 1993-1996. (b) Percent of total dinoflagellate cells that are bioluminescent monthly at San Clemente Island from 1993-1996.

Fig. 8. Bioluminescence produced by each species (photons L\(^{-1}\)) monthly at San Clemente Island from 1993-1996.
Fig. 9. Dinoflagellate species trends for San Clemente Island to San Diego Bay transits from July 1994 to October 1994.

Fig. 10. Dinoflagellate species trends for San Clemente Island to San Diego Bay transits from July 1994 to October 1994.
3.4 Bioluminescence, temperature, rainfall, and chlorophyll relationships

Seawater temperatures in SDB generally ranged from 14°C to 23°C (Figure 11) while mean monthly temperatures at SCI ranged from 14.7°C to 20°C (Figure 12). Correlation coefficients were calculated for both sites. No significant correlations were detected (SDB: n = 44 months; r = 0.131; p > 0.10) (SCI: n = 30 months; r = -0.241; p > 0.10).

Fig. 11. Mean monthly bioluminescence (photons sec⁻¹ ml⁻¹) and seawater temperature (°C) at San Diego Bay from 1992-1996.

Fig. 12. Mean monthly bioluminescence (photons sec⁻¹ ml⁻¹) and seawater temperature (°C) at San Clemente Island from 1993-1996.

3.5 Upwelling indices

Upwelling Indices for 33°N latitude were compared to mean monthly bioluminescence for SDB (Schwing et al. 1996). These indices are north Pacific wind-driven transports computed from monthly average surface pressure data in cubic meters of water per second along each 100 meters of coastline (Figure 13). Cold, nutrient rich water containing nitrates and trace metals are brought to the surface as waters are pushed away from the coast. Nitrates are limited in surface waters (Holm-Hansen et al. 1966, Armstrong et al. 1967, Strickland 1968), and are utilized by all phytoplankton for growth (Spencer 1954, Dugdale 1967, MacIsaac and Dugdale 1969). Inspection of the data shows there is a general trend for upwelling and bioluminescence to co-occur during the same months for 1993-1994 (February - November) but not for 1995 and 1996. Correlation coefficients were not significant for both years (1993:
n = 8 months; $r = 0.307; p > 0.10$), (1994: n = 8 months; $r = 0.617; p > 0.10$) and all 4 years (1992-1996: $r = 0.111; p > 0.10$; n = 47 paired monthly points). Bioluminescence actually began to increase in the winter of 1994 before the onset of upwelling (Figure 13) which suggests that some other factor than upwelling may be controlling the onset of maximum bioluminescence.

### 3.6 Rainfall effects

Total rainfall for San Diego County from 1992 through 1996 correlated with SDB bioluminescence. Four years of data (1992-1996) showed that years with the greatest precipitation also had measurably more bioluminescence at SDB (Figure 14). The year 1995 was different than prior years in that rainfall preceded upwelling (Figure 13) with a marked increase in bioluminescence. In addition, years with less monthly rainfall (3-4 inches per month for 1994 and 1996 vs. 8-9 inches per month for 1993 and 1995) exhibited less bioluminescence. Total bioluminescence (photons ml$^{-1}$ year$^{-1}$) at San Diego Bay was positively correlated with total rainfall (inches year$^{-1}$) for all 4 years ($r = 0.908; n = 4$;

![Fig. 13. Mean monthly bioluminescence (photons sec$^{-1}$ ml$^{-1}$) and upwelling index (m$^3$ sec$^{-1}$) for San Diego Bay from 1992-1996.](image1)

![Fig. 14. Mean monthly bioluminescence (photons sec$^{-1}$ ml$^{-1}$) and monthly rainfall (inches month$^{-1}$) for San Diego Bay from 1992-1996.](image2)
p < 0.05) (Figure 15). Total bioluminescence at SDB was approximately 41% more in 1992-1993 than in 1993-1994 while 1994-1995 was approximately 66% and 82% greater than in 1993-1994 and 1995-1996, respectively (Figure 16). Bioluminescence at SCI was also 287% greater in 1994-1995 than in 1993-1994. These data suggest that the development of bioluminescence were favored by the consequences of rainfall such as storm runoff nutrients from soil. During the period of 1992 through 1996, of the 77 rain events within San Diego County, 51 events or 66% of all rain events with > 0.1 inch of rainfall were associated with a 50% increase in bioluminescence within three days of the start of rainfall at SDB. Past monitoring programs in San Diego County have shown that storm runoff entering coastal and bay waters during the winter and spring months contains high levels of nitrates and phosphates as well as other nutrients and metals (City of San Diego Stormwater Monitoring Program 1994-1995). CALCOFI data sets also show elevated nitrate levels in surface waters off San Diego along transit lines 90 and 93 following peak rainfall periods.

Fig. 15. Correlation of total bioluminescence (photons ml⁻¹ year⁻¹) and total rainfall (inches year⁻¹) for San Diego Bay from 1992-1996. (r = correlation coefficient; p = significance level).

Fig. 16. Total bioluminescence (photons ml⁻¹ year⁻¹) measured at San Diego Bay and San Clemente Island from 1992-1996.
3.7 Seasonality of bioluminescence and chlorophyll \(a\) at San Clemente Island and San Diego Bay

Total bioluminescence (photons ml\(^{-1}\) year\(^{-1}\)) for each year at SCI and SDB was divided into seasons (Summer, Fall, Winter, and Spring). Total bioluminescence for each month was summed and that subtotal was divided by the entire year’s bioluminescence. Seasonal bioluminescence was calculated for SCI (Figure 17a) and SDB (Figure 17b). In 1993-1994, the percent of annual bioluminescence was fairly evenly distributed among all seasons, although the maximum percent of bioluminescence was measured in the winter months at SCI. A winter maximum was again measured the following winter (Figure 17a). For three of the four years at SDB, the maximum percent of annual bioluminescence was measured in the spring. Spring percentages ranged from approximately 30 - 50% of all bioluminescence measured for each of the years (Figure 17b).

Seasonal mean Chl \(a\) maxima were measured in the spring and winter for 1993-1994 and 1994-1995 at SCI, respectively (Figure 18a). Seasonal mean Chl \(a\) maxima were measured in the Spring from 1993-1996 for SDB (Figure 18b). Chl \(a\) was usually greater for all seasons at SDB than at SCI.

The seasonal percentages of bioluminescence for SCI and SDB for all years were averaged as were the seasonal means of Chl \(a\) for SCI and SDB for all years. Peak bioluminescence was measured in the winter at SCI and the spring at SDB. Forty-four percent of all bioluminescence measured at SCI was in the winter while only 16.5% of the year's total was measured in the summer (Figure 19a). Thirty seven percent of all bioluminescence measured at SDB was in the spring while in the fall, only 14% of the total bioluminescence was measured. Maximum mean Chl \(a\) was also measured in the winter (0.87 µg L\(^{-1}\)) at SCI as was bioluminescence while maximum mean Chl \(a\) in SDB was measured in the spring (2.39 µg L\(^{-1}\)) (Figure 19b). San Clemente Island can then be characterized as having winter maxima for bioluminescence and Chl \(a\) while San Diego Bay has spring maxima for both.

High levels of chlorophyll at SDB generally occurred either in the spring or summer, although an extended winter - spring peak was measured in January and April-May 1993. At SCI, measured Chl \(a\) ranged from a low of 0.04 µg L\(^{-1}\) in November 1993 to a high of 1.9 µg Chl L\(^{-1}\) measured in January 1995 which was probably attributable to the presence of high numbers of *Gonyaulax*, as well as chain diatoms and pico-plankton. Chlorophyll \(a\) levels were still high through June 1995 although decreasing through the remaining months. Simple correlations were calculated between monthly means of Chl \(a\) and bioluminescence at SDB and SCI. The correlation between monthly Chl \(a\) and bioluminescence at SDB was not significant \((r = 0.277; n = 43; \ p < 0.10)\). At SCI, the correlation was highly signifcant when red tide data were included \((r = 0.88; n = 26; \ p < 0.001)\). However, when the red tide data were deleted (January - March 1995), the correlation was similar to that at SDB and was not significant \((r = 0.237; n = 23; \ p > 0.10)\).

Data from the bathyphotometer stations (\(n = 26\)) for all six cruises into the Bight (July and November 1994, February, June, November 1995 and March 1996) did not provide a significant correlation of water column bioluminescence with Chl \(a\) (0 - 90 m) \((r = 0.392; \ p > 0.10)\) (Figure 20). Further, integrated water column data for all stations from 1994-1996 or station averages also did not display an association between bioluminescence and Chl \(a\) (Figure 20).
Fig. 17. (a) Distribution of bioluminescence (% of annual bioluminescence) by season measured at San Clemente Island from 1983-1995. (b) Distribution of bioluminescence (% of annual bioluminescence) by season measured in San Diego Bay from 1992-1996.
Fig. 18. (a) Seasonal mean Chl $a$ (µg L$^{-1}$) distribution at San Clemente Island from 1993-1995. 
(b) Seasonal mean Chl $a$ (µg L$^{-1}$) distribution in San Diego Bay from 1992-1996.
Fig. 19. (a) Mean seasonal bioluminescence (% of total) for all years at San Clemente Island and San Diego Bay. Mean seasonal Chl a (µg/L) distribution for all years at San Clemente Island and San Diego Bay. (b) Mean seasonal Chl a (µg/L) distribution for all years at San Clemente Island and San Diego Bay.

Both the mean integrated water column bioluminescence and Chl a increased during the red tide in February 1995 (range: $6 \times 10^{16} - 4 \times 10^{17}$ photons s$^{-1}$ m$^{-2}$ and 16 - 112 mg Chl m$^{-2}$, respectively) (Figure 20) in comparison to previous measurements conducted in November 1994 (range: $7 \times 10^{15} - 2 \times 10^{16}$ photons s$^{-1}$ m$^{-2}$ and 20 - 31 mg Chl m$^{-2}$, respectively). Mean integrated water column bioluminescence in June 1995 returned to former levels measured in July and November 1994. Station averages of integrated bioluminescence increased as the Southern California mainland was approached. Minimum station averages of integrated bioluminescence were measured at stations 2 and 3, east of SCI in the outer Santa Barbara Passage. Maximum station averages were measured at stations 7, 8 and 9, west of the Coronado Escarpment. Water depths here are the shallowest of all stations measured. Station 7 is located in waters with a depth of 1050 meters while station 9 is located in waters with a depth of 155 meters. Stations 2 and 3 are found in waters with depths of 1660 meters and 1290 meters, respectively.
Fig. 20. Integrated bioluminescence (photons sec$^{-1}$ m$^{-2}$) and Chl $a$ (mg m$^{-2}$) averages of all stations from 1994-1996. Averages of individual stations are also shown.

The vertical structure within the water column with respect to other measured parameters (temperature, percent light transmission, and in vivo Chl fluorescence) changed seasonally (Figure 21). For example, at Station 3 in July (7/12/94), bioluminescence was significantly correlated with in vivo Chl fluorescence ($r = 0.673; p < 0.001$), beam attenuation ($r = 0.747; p < 0.001$), and temperature ($r = 0.892; p < 0.001$; Figure 21). Beam attenuation was positively correlated with in vivo Chl fluorescence ($r = 0.831; p < 0.001$). Maximum bioluminescence and in vivo Chl fluorescence were measured at the bottom of the mixed layer (20 m below the sea surface). The mixed layer deepened in November (11/10/94) as did maximum bioluminescence and Chl fluorescence. The correlation between bioluminescence and chlorophyll fluorescence ($r = 0.483; p < 0.001$) in November was significant as was bioluminescence and beam attenuation ($r = 0.954; p < 0.001$) and bioluminescence with...
temperature \( (r = 0.889; \ p < 0.001) \). Bioluminescence and \textit{in vivo} Chl fluorescence were not correlated February 12, 1995 at station 3 \( (r = 0.062; \ p > 0.10) \), however, bioluminescence was still significantly correlated with temperature \( (r = 0.765; \ p < 0.001) \). On June 13, 1995, bioluminescence and \textit{in vivo} Chl fluorescence were significantly correlated \( (r = 0.582; \ p < 0.001) \). Bioluminescence and beam attenuation \( (r = 0.788; \ p < 0.001) \) and bioluminescence and temperature \( (r = 0.703; \ p < 0.001) \) were significantly correlated. Beam attenuation was significantly correlated with \textit{in vivo} Chl fluorescence \( (r = 0.942; \ p < 0.001) \). These correlations improved as the mixed layer became shallower in June (Figure 21) in comparison to the deeper mixed layer observed in November 1994 and February 1995 (Figure 21).

4. Discussion

The data show that bioluminescence changes seasonally in the Southern California Bight coastal waters with a maximum and minimum signal in the spring and fall in SDB (Figures 2,14,17b,19a). A winter maximum and summer minimum in bioluminescence was measured at SCI (Figure 2, 17a). In SDB and SCI, we observed a change in the dinoflagellate species composition over a year and its contribution to bioluminescence. We also observed a seasonal change in species composition (summer to winter) at SCI and within the bight (Figures 8a, 9, 10). Chlorophyll $a$ also showed similar seasonal trends with respect to location (Figures 18,19). However, measured monthly means of bioluminescence did not correlate with Chl $a$ either at SDB or SCI. Mean monthly surface seawater temperature did not correlate with mean monthly bioluminescence at either site; that is, maximum bioluminescence did not always correlate with either maximum or minimum seawater temperatures (Figures 11, 12), although minimum bioluminescence was measured during the coolest water temperatures (winter) at SDB in 1994 and 1996. The largest peak in bioluminescence measured at SCI (winter 1995) was associated with the coolest seawater temperatures (14-15°C) during winter (Figure 12). Coolest water temperatures did not correlate with the upwelling index as maximum indices for 33°N latitude, 119°W longitude generally occurred in June of each year.

Total bioluminescence (photons ml$^{-1}$ year$^{-1}$) was always greater at SDB than at SCI. Total bioluminescence at SDB ranged from $1.06 \times 10^{13}$ to $1.93 \times 10^{13}$ photons ml$^{-1}$ year$^{-1}$ (measured from 2100 to 0300 hrs each day) while total bioluminescence measured at SCI was from $3.45 \times 10^{12}$ to $9.91 \times 10^{12}$ photons ml$^{-1}$ year$^{-1}$. In 1993-1994, 3 times more bioluminescence was measured at SDB than at SCI. These differences lessened to a factor of 2 in 1994-1995 between both sites when a massive bioluminescence red tide was observed to extend south from Santa Barbara, California to Ensenada, Mexico and 100km offshore to SCI. At times, monthly differences in total bioluminescence were 8 times greater at SDB than at SCI in Spring 1994 and 1995.

Bioluminescent dinoflagellates, in most instances, comprised most of the dinoflagellates collected at SDB and SCI (Figures 3, 7). In SDB, bioluminescent dinoflagellates made up at least 80% of all dinoflagellates. Numbers of bioluminescent dinoflagellates dropped noticeably in the winter and spring at SDB (< 30% of total dinoflagellates) at SDB. Decreases in bioluminescent dinoflagellates were observed at SCI in late spring at SCI. The bioluminescent dinoflagellate assemblage at both SDB and SCI was composed of \textit{Ceratium}, \textit{Gonyaulax}, \textit{Protoperidinium}, and \textit{Noctiluca} species. \textit{Pyrocystis noctiluca} was a recurring species.
Fig. 21. Bathyphtometer profiles at station 3 for (a) July 12, 1994; (b) station 3 November 10, 1994; (c) station 3 February 12, 1995; (d) station 3 June 13, 1995.
found at SCI. *Protoperidinium* spp. and *Gonyaulax polyedra* contributed most of the bioluminescence at both sites. *Noctiluca miliaris* contributed substantial bioluminescence following increases in *G. polyedra* at SDB in 1995 and 1996 and at SCI in 1995.

Total rainfall was significantly correlated with measured bioluminescence at SDB ($r = 0.908$; $n = 4$; $p < 0.05$). Years with the greatest rainfall (1993, 1995) affected the total bioluminescence which implies that processes associated with rainfall, such as storm water runoff may be stimulating dinoflagellate and algal production in coastal waters (Anderson 1964; Eppley et al., 1978). We observed that the upwelling index did not directly correlate with SDB bioluminescence unless the index was shifted back 1 month ($r = 0.476$; $p < 0.001$). However, if mean monthly bioluminescence was shifted forward 2 months with mean rainfall, a significant correlation was observed ($r = 0.472$; $p < 0.01$). The upwelling index and nitrates ($\mu M L^{-1}$) measured in coastal waters, were significantly correlated when nitrate levels were shifted forward in time 1 month ($r = 0.679$; $p < 0.001$). We must then assume that some other factor besides upwelling is providing a stimulatory effect to dinoflagellate bioluminescence. Multiple regression analysis showed that rainfall, upwelling, and temperature were the most important conditions to predict bioluminescence and that when rainfall was moved ahead in time by 2 months, we could account for 24.7% of the observed variance to predict bioluminescence from 1992 - 1996 ($R^2 = 0.2468$; $F=2.469$; $p<0.05$). Increased nitrate levels were observed in coastal waters beyond SCI during the winter months and spring months; before maximum upwelling. The source of these nitrates may be in storm water runoff. Support for "new sources of nitrogen" versus "recycled nitrogen" and other nutrients entering the water column is not new. Some studies have shown that river inputs into the ocean can carry high levels of nutrients needed for algal growth (Harrison 1980, Fogg 1982, Mooers et al. 1978, Lalli and Parsons, 1993). Others have found that ferric iron is a limiting nutrient for phytoplankton growth (Menzel and Ryther 1961) and that high levels of iron are often associated with river runoff (Williams and Chan 1966). Iron is needed by phytoplankton to utilize nitrates for growth (Ryther and Kramer 1961). The availability of iron is enhanced by chelation with dissolved organic matter. That is, organically bound iron from storm runoff may stimulate the growth of phytoplankton (Kawaguchi and Lewitus 1996). Similarly, elevated phytoplankton levels off Del Mar, California following storm water runoff were attributed to increased nutrient inputs from land (Eppley et al. 1978).

The bathyphotometer stations showed that bioluminescence and Chl fluorescence were positively correlated, during the summer months, when the water column stabilized with a shallow thermocline. These significant positive correlations broke down with water column mixing during the fall and winter months but were reestablished with the development of the thermocline during the spring and summer months. Several species of *Protoperidinium* were the predominant dinoflagellate in the spring and summer months while *G. polyedra* was important during the fall and winter months; not only in surface waters, but at depth. The increased bioluminescence and chlorophyll levels associated with the red tide at SCI are remarkable for their duration since they persisted from January through April 1995. This strengthens the inference that the physical environment in the bight is fairly stable with respect to seasonality, and that bioluminescence is strongly influenced by seasonal rainfall and runoff.

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Southern California Bight bioluminescence is similar to that found in coastal waters of Vestfjord, Norway (Lapota 1990, unpublished), and the Arabian Sea (Lapota & Rosenberger 1990), but higher than that found in the Sargasso Sea (Batchelder & Swift 1989), the North Atlantic (Neilson et al. 1995) and the Beaufort Sea (Lapota et al. 1992). The vertical structure of bioluminescence was correlated with Chl fluorescence for some of the stations in the Bight. However, integrations between bioluminescence and chlorophyll were positively correlated, but weak. Strong positive correlations between bioluminescence and chlorophyll fluorescence were observed during the red tide in February 1995. At depth, seawater temperature correlated strongly with the vertical distribution of bioluminescence, as did transmission. In contrast, weaker correlations were observed between bioluminescence and Chl fluorescence. Other studies have infrequently observed correlations which may be dependent on the season the study was conducted (Lapota et al. 1989 Young et al. 1992, Neilson et al. 1995, Ondercin et al. 1995). An obvious conclusion is that the primary dinoflagellates which are contributing much of the bioluminescence do not contain Chl a. These would include the heterotrophic Protoperidinium dinoflagellates. These dinoflagellates produce as much as 30 times more light per cell than does G. polyedra (Biggley 1969, Lapota et al. 1992). This could explain the poor correlations between bioluminescence and chlorophyll. Consequently, these results impact models predicting bioluminescence from global ocean primary production and ocean color (Young et al. 1992, Ondercin 1995) since these are based on the assumption that much of the oceanic bioluminescence is derived from photosynthetic bioluminescent dinoflagellates (Ondercin 1995). It is clear that from this and other studies (Lapota et al. 1989, 1992, 1993 a, b, Swift et al. 1995, Neilson et al. 1995) that Protoperidinium dinoflagellates dominate surface water bioluminescence in the world's oceans for a significant portion of the year.

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6. References

Long Term Dinoflagellate Bioluminescence, Chlorophyll, and Their Environmental Correlates in Southern California Coastal Waters


Tarasov, N.I. (1956). Marine luminescence. USSR Academy of Sciences, Moscow [in Russ.] [Engl transl by Naval Oceanographic Office (No NOOT-21); National Space Technology Laboratories Station, Bay St. Louis, MS].


We now find ourselves utilizing luciferase - luciferin proteins, ATP, genes and the whole complex of these interactions to observe and follow the progress or inhibition of tumors in animal models by measuring bioluminescence intensity, spatially and temporally using highly sophisticated camera systems. This book describes applications in preclinical oncology research by bioluminescence imaging (BLI) with a variety of applications. Chapters describe current methodologies for rapid detection of contaminants using the Milliflex system, and the use of bioluminescence resonance energy transfer (BRET) technology for monitoring physical interactions between proteins in living cells. Others are using bioluminescent proteins for high sensitive optical reporters imaging in living animals, developing pH-tolerant luciferase for brighter in vivo imaging, and oscillation characteristics in bacterial bioluminescence. The book also contains descriptions of the long-term seasonal characteristics of oceanic bioluminescence and the responsible planktionic species producing bioluminescence. Such studies are few and rare.

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