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# Short-term variability in primary productivity during a wind-driven diatom bloom in the Gulf of Eilat (Aqaba)

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ABSTRACT: In the northern Gulf of Eilat (Agaba), sharp increases in the biomass of diatoms and rates of primary production occurred in April 2008. Within 24 h, diatom abundance rose from  $8 \times 10^3$ to  $228 \times 10^3$  cells l<sup>-1</sup>, and photosynthetic rates concomitantly doubled from 15 to 35  $\mu$ g C l<sup>-1</sup> d<sup>-1</sup>. Water transparency declined, as indicated by the vertical diffusion attenuation coefficient  $K_{\rm d}$  for photosynthetically active radiation (PAR), which increased from 0.076 to 0.090 m<sup>-1</sup> and decreased the euphotic depth from 60 to 45 m. During this time, a significant increase in silica deposition by the diatoms was also detected. We attribute the mentioned changes in environmental characteristics to wind-generated surface currents. Strong winds (up to 10 m s<sup>-1</sup>) during the measurements enriched the surface layers with unusually high nutrient concentrations within <1 d. Hence, primary production rates were observed at a relatively eutrophic nearshore station (MP) and pelagic station (A<sub>1</sub>, 10 km towards the center of the Gulf). They were compared with rates measured on the subsequent day. Values measured were twice as high as those at the pelagic station during the previous calm day. Routine monitoring programs with monthly or semi-weekly sampling are thus likely to miss brief but significant injections of nutrients, leading to the underestimation of seasonal and annual primary production. Our results demonstrate the impacts of transient events on the function and annual yield of aquatic ecosystems.

 $\label{eq:keywords} \mbox{KEY WORDS: Primary production} \cdot \mbox{Wind speed} \cdot \mbox{Upwelling} \cdot \mbox{Phytoplankton} \cdot \mbox{Diatom blooms} \cdot \mbox{Phytoplankton absorption}$ 

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#### INTRODUCTION

During the 8th International Workshop of the Group on Aquatic Primary productivity (GAP) that took place at the Interuniversity Institute in Eilat (northern Red Sea, Israel), concomitant measurements of primary productivity were conducted in accordance with the philosophy and practice of GAP (Bar Zeev et al. 2009, this Special Issue). Here we describe an unexpected event that occurred while measuring planktonic primary productivity in the Gulf of Eilat (Aquaba), and which exposes potential underestimation resulting

from the current routine practices of primary productivity monitoring.

Spatial and temporal distributions of primary production (PP) are consistent with limitations by light, nutrients, and temperature (Field et al. 1998). Nitrogen (N) limitation characterizes large areas of the world's oceanic, coastal, and estuarine waters (Dugdale 1967, Ryther & Dunstan 1971, Codispoti 1989, Mackey et al. 2006), and the supply rate of biologically available N to these waters is a key control of PP and the resultant trophic state (Ryther & Dunstan 1971, Eppley & Peterson 1979, D'Elia et al. 1986, Nixon 1986, Tyrrell 1999, Chen et al. 2008).

While nutrient limitation of PP in the ocean has traditionally been attributed to N, iron (Fe) availability has more recently been found to be important (Chase et al. 2006, Chen et al. 2008), and phosphorus (P) has been suggested as the ultimate limiting nutrient over geological timescales (Redfield 1958, Follmi 1996, Tyrrell 1999, Mackey et al. 2006, 2007). The major source of P in the ocean is the weathering of minerals on land and their subsequent introduction into the ocean by fluvial and aeolian transport (Filippelli & Delaney 1996, Benitez-Nelson 2000, Chen et al. 2008). Because there is no P input process analogous to N fixation, marine productivity over geological timescales is considered to be a function of the supply rate of P from continental weathering and the rate at which P is recycled in the ocean (Scanlan & Wilson 1999, Mackey et al. 2007).

There are a few events that can significantly increase oceanic PP. Hans (1997) found that atmospheric deposition (AD) contributes from 300 to >1000 mg N m<sup>-2</sup> yr<sup>-1</sup> to coastal waters in the biologically available forms NO<sub>x</sub>-, NH<sub>3</sub>/NH<sub>4</sub>+, and dissolved organic N; and groundwater (GW) can jointly account for 20 to >50% of total exogenous or 'new' N loading and may uniquely mediate coastal eutrophication by bypassing estuarine filters of terrigenous N inputs. Trace metal enrichment (e.g. Fe) in AD and GW may interact synergistically with N to stimulate coastal production. In deep (>200 m) oceanic regions away from land masses, much of the PP is dependent on regenerated ammonium derived from organic matter mineralization. Exceptions include production supported by N fixation or NO<sub>3</sub> from upwelled deep water, both of which are considered new N sources. Coastal and estuarine environments are heavily influenced by new N, which is supplied either naturally by weathering of minerals, decomposition, lightning, and geothermal emissions, or anthropogenically (Hans 1997).

Increasing quantities of atmospheric anthropogenic fixed N entering the open ocean could account for up to % of the ocean's external (non-recycled) N supply and up to  $\sim\!3\,\%$  of the annual new marine biological production of  $\sim\!0.3$  Pg C yr $^{-1}$  (Duce et al. 2008).

In the Gulf of Eilat, aerosol-borne trace element deposition has an impact on the surface water chemistry (Chen et al. (2008) and may thus affect PP in the surface water by supplying important macro- and micronutrients or, possibly, high levels of toxic elements. The study by Chen et al. (2008) shows that, even in an area dominated by mineral dust deposition and surrounded by deserts, anthropogenic air emissions may make a significant contribution to the levels of various trace elements such as Cu, Cd, Ni, Zn, and P. The estimated residence time of a few years for most trace elements underscores the importance of atmospheric fluxes to the Gulf ecosystem and the episodic nature of such deposition events demonstrates that these may have a large impact.

Other phenomena that can change nutrient availability include the El Niño-Southern Oscillation (ENSO) and cyclones. Between September 1997 and August 2000, biospheric net PP varied by 6 Pg C yr<sup>-1</sup> (from 111 to 117 Pg C yr<sup>-1</sup>) (Behrenfeld et al. 2001). The passage of cyclone Wasa in December 1991 led to increases in biomass and PP that were related to nutrient enrichment either from terrestrial runoff or resuspended sediments. The predominance of diatoms during cyclones seems to be a general pattern that can be related to their higher growth rates compared to other phytoplankters (Delesalle et al. 1993).

Episodic eddy-driven upwelling may supply a significant fraction of the nutrients required to sustain PP of the subtropical ocean. Mode-water eddies can generate extraordinary diatom biomass and PP at depth relative to the time series near Bermuda. These blooms are sustained by eddy-wind interactions that amplify the eddy-induced upwelling (McGillicuddy et al. 2007).

The Gulf of Eilat, which is an arm of the Red Sea, is an oligotrophic region with predictable seasonal cycles in macronutrient concentrations and phytoplankton community structure that are similar to other oligotrophic areas of the world's oceans, such as the open ocean gyres. Changes in the dominance of different phytoplankton groups occur as the deep mixing conditions in winter relax, and stratification intensifies beginning in early spring and continuing through summer. Specifically, a bloom of eukaryotic nanophytoplankton characteristic of the winter–summer transition is replaced by a summer community dominated by picophytoplankton (<5  $\mu$ m), of which *Prochlorococcus* and *Synechococcus* are the most numerous (Lindell & Post 1995).

Atmospheric inputs of other nutrients gradually increase the likelihood of P limitation in the Gulf (Chen et al. 2007). Simultaneously, these atmospheric inputs provide an important source of seawater-soluble iron (Chase et al. 2006), which appears to remain in excess of that required for microbial growth throughout the

year (Chen et al. 2007). During summer and fall, stratified surface waters are depleted of nutrients and picophytoplankton populations comprise the majority of cells (80 and 88%, respectively). In winter, surface nutrient concentrations are higher and larger phytoplankton are more abundant (63%) (Mackey et al. 2007).

The aim of the present study was to describe a winddriven transient high primary productivity event in the Gulf of Eilat that was associated with a diatom bloom. The mechanism by which winds brought about the phytoplankton response is shown, and the implications of such dramatic spikes for estimates and models of PP are discussed.

#### MATERIALS AND METHODS

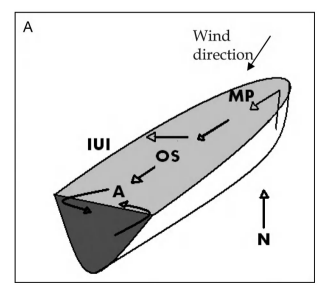
Observations were performed during the GAP Workshop held on 30 March to 6 April 2008 at the Interuniversity Institute of Marine Sciences in Eilat. *In situ* measurements and water sampling were carried out on 1 April 2008 at the following 3 stations: Stn A<sub>1</sub>, located ~10 km southeast of the northern shore of the Gulf of Eilat along its meridional axis at 29°28′N, 34°55′E, with a bottom depth of ~700 m; Stn OS (open sea) (29°29′N, 034°55′E), located in the middle of the Gulf opposite the IUI (Interuniversity Institute), with 400 m bottom depth; and Stn MP (Military Port) (29°32′N, 034°57′E), located at the northern shore 100 m from the coast, with 30 m bottom depth (Fig. 1). Depth, temperature, salinity, and fluorescence were measured with a CTD-19 (Sea-Bird Electronics) and an

in situ fluorometer (WetStar, WetLabs). Water samples were obtained with a rosette system of samplers from discrete depths of 1, 20, 40, 60, 80, and 100 m, and subsamples were taken from these depths for the measurement of PP. Laboratory analysis of the water samples comprised analyses of phytoplankton community structure, pigment absorption spectra, and heterotrophic bacterial count.

**Light measurements.** The underwater light field was measured using a standard high-resolution profiling reflectance radiometer (Biospherical PRR-800), and both downwelling ( $E_d$ ) and upwelling irradiance ( $L_u$ ) were recorded at 18 wavelengths (305, 313, 320, 340, 395, 443, 465, 490, 520, 560, 589, 665, 683, 694, 710, 765, 780, and 875 nm) as well as  $E_{\rm dz}$ PAR (photosynthetic available radiation) (400-700 nm) and  $L_{\rm uz}$ Chl (natural fluorescence). The PRR-800 also included internal sensors for temperature, detector-array temperature, inclination (roll/pitch), and depth. An onboard computer recorded the data during the casts. Light casts were done from the surface to 100 m depth. The diffusion attenuation coefficient  $(K_d)$  and euphotic zone depth  $(Z_{eu})$  were calculated from the  $E_{\rm d}$ PAR profiles.  $E_{\rm d}$  was determined at 1 m intervals, and  $K_{\rm d}$  was calculated over 5 m intervals, using PROFILER software as:

$$K_{\rm d}(\lambda) = \frac{\ln(E_{\rm d}(\lambda)_{(z_1)} / E_{\rm d}(\lambda)_{(z_2)})}{(Z_2 - Z_1)} \tag{1}$$

where  $E_{\rm d}$  is the downwelling irradiance (in  $\mu$ W cm<sup>-2</sup> nm<sup>-1</sup> for any specific wavelength  $\lambda$  or  $\mu$ mol photon cm<sup>-2</sup> s<sup>-1</sup> for PAR),  $Z_1$  and  $Z_2$  are 2 measured depths (m) and  $K_{\rm d}$  is the attenuation coefficient (m<sup>-1</sup>) calculated for the depth interval  $Z_2 - Z_1$ . The euphotic depth ( $Z_{\rm eu}$ )



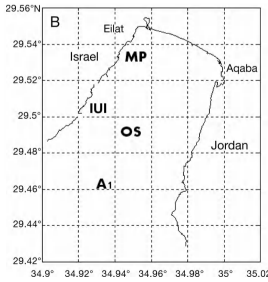


Fig. 1. (A) Schema of currents in the Gulf of Eilat in response to upwelling from northern coastal waters and downwelling from the west to the open sea deep water. (B) Map showing Stn  $A_1$ : 700 m bottom depth; Stn OS: 400 m; Stn MP: 30 m, and IUI: incubation station, 150 m

was calculated as the depth where 1 % of  $E_{\rm d}$ PAR at 1 m was measured. The  $E_{\rm d}$ PAR values at 1 m were used instead of subsurface values since the latter were unusable due to high surface noise.

Nutrients. Nitrites and nitrates were analyzed using the colorimetric method described by Grasshoff et al. (1999) using a flow injection autoanalyzer (FIA, Lachat Instruments Model QuikChem 8000). The analysis was fully automated and peak areas were calibrated using standards prepared in 'low nutrient' filtered seawater (surface seawater sampled during the oligotrophic season in the Gulf of Aqaba) over a range of 0 to 5  $\mu$ mol  $l^{-1}$ . The precision of this method is  $\pm 0.02~\mu$ mol  $l^{-1}$  for nitrites and  $\pm 0.05~\mu$ mol  $l^{-1}$  for nitrates.

**Chl** *a.* Chl *a* was estimated using natural fluorescence, i.e.  $L_{\rm u}$  from the PRR-800 at 683 nm. Chl *a* concentration (mg m<sup>-3</sup>) may be calculated from the natural fluorescence flux ( $F_{\rm f}$ , nmol photon m<sup>-3</sup> s<sup>-1</sup>), and the scalar irradiance ( $E_{\rm 0}$ , µmol photon m<sup>-2</sup> s<sup>-1</sup>), i.e. light flux from all directions:

$$Chl = \frac{F_{f}}{^{\circ}a_{c}PAR \times \phi_{f} \times E_{o}PAR}$$
 (2)

Two important optical assumptions are used in this equation:  ${}^{\circ}a_{\rm c}$ PAR is the chl a-specific absorption coefficient (absorption normalized to chl a concentration) and  $\phi_{\rm f}$  is the quantum yield of fluorescence. These parameters are treated as constants in the software for the PRR-800, with respective values of 0.04 m² mg² and 0.045 µmol photon fluoresced m² s²¹ (µmol photon absorbed)²¹ cm²² s⁻¹.

High-resolution measurements (4 to 6 times per day) resulted in a map of chl a values during the period before and after the study period. Data were plotted and interpolated using Ocean Data View (ODV) software (Schlitzer 2008). Since the chl a estimation algorithm is based on empirical constants (the chl a-specific absorption coefficient and the quantum yield of fluorescence), which were not derived specifically for our phytoplankton assemblages, we calibrated the estimated chl a against extracted chl a concentrations throughout 2008 (including profiles acquired during the present study), using 51 samples over a range of 0.01 to 1.14 mg chl a m $^{-3}$ . The calibration factor was calculated as chl a (mg m $^{-3}$ ) = 0.04  $F_{\rm f}$  chl - 0.0192 (R $^2$  = 0.804).

**Absorption spectra.** *In vivo* phytoplankton absorption was measured using the filter-pad method (Mitchell et al. 2000) from the discrete samples collected at different depths. Amounts of 600 to 2000 ml water were filtered onto 25 mm GF/F (Whatman). The filters were placed in histoprep tissue capsules (Fisher Scientific) and stored at  $-20^{\circ}$ C until analysis 1 or 2 d later. The filters were scanned at 350 to 750 nm using a UV-visible spectrophotometer (Cary 50) to obtain total

particulate optical density ( $OD_{part}$ ). After pigment extraction with methanol, the filters were rescanned to obtain the optical density of the detrital material ( $OD_{det}$ ). OD spectra were then corrected for pathlength amplification (Cleveland & Weidemann 1993) and converted to absorption coefficient ( $a_{part/det}$ ) using:

$$a_{x}(\lambda) = \frac{2.303 \times \text{OD}_{x}(\lambda)}{X}$$
 (3)

where the subscript x denotes either particulate or detrital OD ( $\lambda$ ) and X is the ratio of the filtered volume to the filter clearance area.

The absorption spectra were then corrected for residual scattering by subtracting the average over 748–752 nm from all measured spectra.

Phytoplankton absorption  $(a_{\rm ph})$  was calculated as the difference between total particulate  $(a_{\rm part})$  and detrital absorption  $(a_{\rm det})$ :

$$a_{\rm ph} = a_{\rm part} - a_{\rm det} \tag{4}$$

**Phytoplankton enumeration.** Samples (100 ml) for phytoplankton enumeration were collected at 5 m depth and fixed with 100  $\mu$ l of a 25% glutaraldehyde solution. Samples were kept in low light and counted within 2 d of collection. For each sample, a 50 ml subsample was counted using a phase contrast inverted microscope (Utermöhl 1958). Only phytoplankton cells with >5  $\mu$ m diameter were identified (Tomas 1996). Phytoplankton biomass was estimated according to Edler (1979) and Menden-Deuer & Lessard (2000).

Flow cytometry. Samples (1.8 ml) for ultraphytoplankton were preserved with 1% glutaraldehyde (final concentration) and immediately frozen in liquid N and stored at  $-80^{\circ}$ C. Before analysis, samples were quickly thawed at 37°C and counted in a flow cytometer (FACScalibur, Becton & Dickinson). Phytoplankton samples were run at approximately 60  $\mu$ l min<sup>-1</sup> and 10000 events were acquired in log mode. We added yellow-green latex beads (0.93  $\mu$ m, Polysciences) to each sample as an internal standard.

Silica deposition. Biogenic silica deposition was measured in natural phytoplankton assemblages using **PDMPO** (2-(4-pyridyl)-5-((4-(2-dimethylaminoethylaminocarbamoyl) methoxy)phenyl)oxazole) labeling following the modified protocol of Leblanc & Hutchins (2005). On 1 to 3 April, fresh surface and deepwater samples were incubated with PDMPO (final concentration of  $0.125 \mu M$ ) using 250 ml flasks. The samples were incubated in water tables with flowing seawater, and the tables were covered with window screening to reduce surface irradiance to ~25%. After a 24 h incubation, samples were brought to the laboratory and filtered through 2 µm pore size polycarbonate filters (Osmonics, Poretics). Filters were rinsed with distilled water to eliminate any unbound PDMPO, placed on a drop of immersion oil on a glass slide and sealed under a glass cover with another drop of oil above the filter. A portion of cells actively depositing silica was counted immediately after filter preparation; PDMPO fluorescence was measured later. Epifluorescence microscopy coupled with digital image analysis was used for quantitative evaluation of PDMPO fluorescence ( $F_{\rm PDMPO}$ ), which was expressed in relative fluorescence units (RU) cell<sup>-1</sup> and calculated as described in Znachor & Nedoma (2008). A paired 2-tailed Student's t-test was performed to examine differences in Si incorporation between surface populations and those at the deep chlorophyll maximum (DCM), with differences accepted as significant when p < 0.05.

Productivity (carbon assimilation). PP measurements were made using the <sup>14</sup>C uptake method described by Steeman-Nielsen (1952) and modified by Strickland & Parsons (1968). Bottles containing 50 ml samples of seawater were incubated *in situ* for 24 h at their original sampling depth after addition of ~5 μCi of carrier-free NaH<sup>14</sup>CO<sub>3</sub> (New England Nuclear). The mooring line with the incubation bottles was deployed ~1 km offshore of the IUI (Fig. 1) at a bottom depth of 140 to 150 m before first light (~04:00 h), and retrieved at the same time the following morning.

Upon retrieval, the incubation bottles were kept in a black box until filtration. The contents of the bottles were GF/F (Gelman) filtered with a Millipore glass filtration system under light vacuum (15 cm Hg). These filters were then treated with HCl vapors to remove traces of inorganic carbon. The filters were placed in glass vials with 10 ml Insta Gel (Packard) and counted in a liquid scintillation counter (Packard 3255) for 10 min each. Added activity was measured by sampling 50 µl from each bottle and placing it on a GF/F filter soaked with NaOH to trap the CO2. The addedactivity samples were counted in the manner described above. The total inorganic carbon was determined from alkalinity (Strickland & Parsons 1968). Productivity rates were then estimated using the JGOFS (Joint Global Ocean Flux Study) protocol (UNESCO 1994).

Estimated productivity ( $F_c$ -683). Productivity estimated was estimated based on chlorophyll natural fluorescence at 683 nm wavelength, measured with an underwater spectro-radiometer (Biospherical PRR-800). Natural fluorescence per volume ( $F_f$ ) was derived from  $L_u$  in the chlorophyll emission spectrum.  $F_{c-683}$  productivity estimation uses the volume fluorescence ( $F_f$ , nmol photon m<sup>-3</sup> s<sup>-1</sup>) and the scalar irradiance ( $E_o$ PAR, µmol photon m<sup>-2</sup> s<sup>-1</sup>) as well as photosynthetic parameters such as the maximal ratio of the quantum yields of photosynthesis to fluorescence ( $\phi_{rmax}$ , C atom photon<sup>-1</sup>) and the irradiance at which this ratio is ½ of the maximum ( $k_{cf}$ , µmol photon m<sup>-2</sup> s<sup>-1</sup>]). These parameters were determined empirically and assigned as  $\phi_{rmax} = 4$  C atom photon<sup>-1</sup> and  $k_{cf} = 133$  µmol photon

 $m^{-2}$  (Kiefer et al. 1989, Chamberlin et al. 1990, Chamberlin & Marra 1992).

$$F_c = F_f \frac{k_{cf} \phi_{r \max}}{k_{cf} + E_o PAR}$$
 (5)

 $F_{\rm c}$ -683 estimations were made using PROFILER software (Biospherical Instruments). An onboard computer recorded the data online. Measurements were done from the surface down to  $100\,\mathrm{m}$  at 13:00 to  $14:00~\mathrm{h}$ .

Regression of PP (estimated using *in situ* <sup>14</sup>C uptake) against estimated productivity based on fluorescence ( $F_c$ -683, measured in 3 profiles on different dates) showed a significant correlation: Y (mg C m<sup>-3</sup> d<sup>-1</sup>) = 1.469 + 0.592 X ( $F_c$ -683 units);  $r^2$  = 0.823; p < 0.0001; n = 24. Each profile was taken from 5 to 100 m; no fluorescence readings were recorded in the upper 5 m because of the high ambient irradiance interference with readings from the PRR-800 reflectance radiometer.

Wind speed and direction data were obtained from the meteorological database maintained at the IUI.

#### RESULTS

Wind velocity changed dramatically throughout the study period and showed a diurnal variation of mainly high values during the day and low values at night. Wind speed was  $\sim 3 \text{ m s}^{-1}$  at midnight prior to sampling and increased to almost 10 m s<sup>-1</sup> the next day (2 April 2008) (Fig. 2B). The changes in wind speed also affected the temperature profiles, which displayed a sharp decrease (from 21.7 to 21.4°C in 15 h) at the surface (Fig. 2A). The thermocline, which was at 5 m on 1 April at 13:40 h. moved to a depth of ~35 m the following day (Fig. 2B). Temperature profiles during 20 March to 7 May 2008 showed significant changes in temperature during the storm event (Fig. 3). Following the strong wind event on 1 April, water transparency declined;  $K_{\text{d(PAR)}}$  increased from 0.076 to  $0.090~\mathrm{m}^{-1}$  and  $Z_{\mathrm{eu}}$  increased from 60 m to 45 m, which is shallower than typically observed at this time of the year (Iluz 1997).  $Z_{\rm eu}$  continued to decrease for 4 d showing high diel variability, and increased once again by 6 April (Fig. 4).

Chl a concentration was low and showed the typical vertical variation for the study area. The deep chlorophyll maximum (DCM) was at 30 m on 31 March, with a maximum value of ~1.5  $\mu$ g l<sup>-1</sup>, but deepened during the remainder of the observation period, with declining chl a concentrations that reached a minimum on 8 April (Fig. 5).

### Phytoplankton community structure

Throughout the experiment, picophytoplankton ( $<5 \mu m$ ) increased from 25 to 50  $\mu g$  C l<sup>-1</sup> (see Bar Zeev

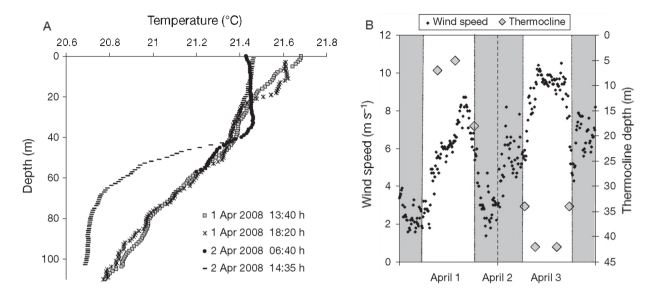


Fig. 2. (A) Temperature profiles and (B) wind speed during 1 to 2 April 2008. Grey shading in (B) indicates dark periods

et al. 2009). The >5 µm fraction of the phytoplankton population was dominated by diatoms, which represented up to 99% of the abundance and 90% of the total biomass of the nanoplankton. Diatoms were at their lowest abundance and biomass on 1 April ( $6 \times 10^3$  cells  $1^{-1}$  and 0.5 µg C  $1^{-1}$ , respectively). The values increased again sharply on 3 April to  $228 \times 10^3$  cells  $1^{-1}$  (abundance) and 20 µg C  $1^{-1}$  (biomass) (Fig. 4A). From 4 April, diatom abundances decreased to values

recorded prior to the strong wind event of 1 April. A centric diatom, *Chaetoceros* cfr. *compressus*, dominated the phytoplankton population during the sampling period, representing up to 84% of the cells identified in the samples and up to 64% of the total biomass. Other chain-forming centric diatoms (e.g. *Leptocylindrus danicus*, *Eucampia cornuta*, and *Chaetoceros curvisetus*) accounted for a maximum of 28% of the remaining phytoplankton biomass.

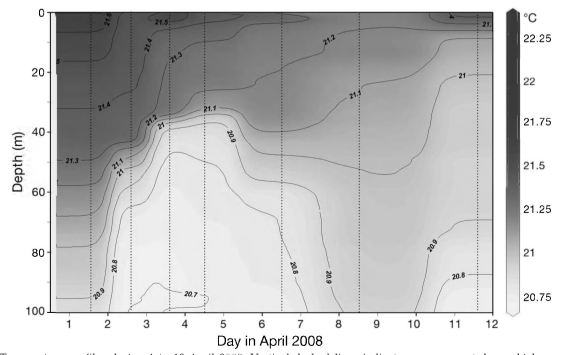


Fig. 3. Temperature profiles during 1 to 12 April 2008. Vertical dashed lines indicate measurement days which occurred in parallel to the storm event, showing significant changes in temperature

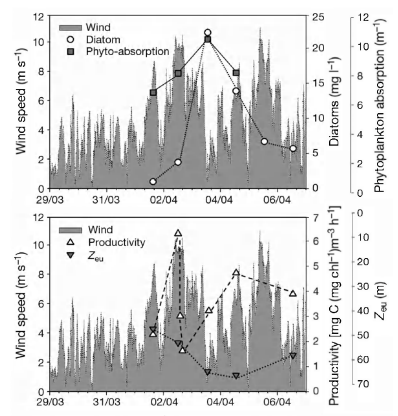


Fig. 4. (A) Wind speed (m s<sup>-1</sup>) during 28 March to 9 April 2008, diatom biomass (µg C l<sup>-1</sup>) at 5 m depth from 1 to 6 April 2008, and phytoplankton absorption of photosynthetically active radiation (PAR) (m<sup>-1</sup>). (B) Wind speed (m s <sup>1</sup>) during 28 March to 9 April 2008, estimated productivity ( $F_c$ -683; mg C [mg chl a]<sup>-1</sup> m<sup>-3</sup> h<sup>-1</sup>), and euphotic depth  $Z_{cu}$  (average of 3 to 5 measurements d<sup>-1</sup>, y-axis reversed)

## Phytoplankton absorption

Phytoplankton pigment absorption changed temporally. Phytoplankton absorption at 435 nm,  $a_{\rm ph}(435)$ , increased from 0.045 m<sup>-1</sup> on 1 April to 0.060 m<sup>-1</sup> on 2 April and 0.062 m<sup>-1</sup> on 3 April, and then decreased to 0.043 m<sup>-1</sup> on 4 April. The  $a_{\rm ph}(675)$  also varied accordingly. Phytoplankton pigment absorption at PAR (400–700 nm) showed a similar trend as  $a_{\rm ph}(675)$ ,  $a_{\rm ph}(435)$ , and the diatom biomass (Fig. 4A).

# **Primary productivity**

Within 1 d, productivity at 20 m depth increased from 15 mg C m<sup>-3</sup> d<sup>-1</sup> at Stn OS (1 April 2008) to 35 mg C m<sup>-3</sup> d<sup>-1</sup> at Stn A<sub>1</sub> (2 April 2008) (Fig. 6). The daily integrated PP in the coastal Stn MP on 1 April was 1141  $\mu$ g C m<sup>-2</sup>. We obtained similar and even higher values (1330  $\mu$ g C m<sup>-2</sup>) at Stn A<sub>1</sub> (located 10 km from Stn MP) on 2 April.

# Silica deposition

Biogenic silica deposition was measured using PDMPO fluorescence to compare the growth rates of the dominant diatom

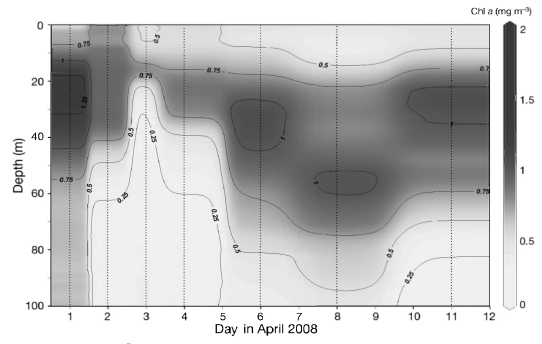


Fig. 5. Chl a concentration (mg m $^{-3}$ ) at various depths, showing changes in the deep chlorophyll maximum (DCM) during 1 to 12 April 2008. Vertical dashed lines indicate measurement days that occurred concurrently with the storm event

Chaetoceros spp. during the study period. Due to low abundance of the other diatom species, their contribution to Si deposition could not be measured. In general, Si deposition of Chaetoceros was significantly higher at the DCM depth than at the surface (t = 2.9, p < 0.01). On 1 April, low abundance of Chaetoceros spp. concurred with low silica deposition rates ( $F_{\rm PDMPO} = 0.07$  RU cell<sup>-1</sup>). During a peak in the Chaetoceros bloom on 2 April, Si deposition rates markedly increased ( $F_{\rm PDMPO} = 0.38$  RU cell<sup>-1</sup>) and most of the population was involved in Si deposition at both the surface and the

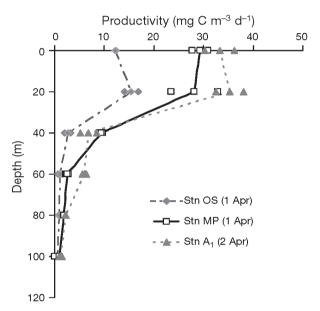


Fig. 6. In situ primary production profiles at Stns MP (Military Port, 1 April), OS (open sea, 1 April), and A<sub>1</sub> (2 April). The incubation at Stn A<sub>1</sub> was on 2 April 2008

DCM, indicating high growth rates. However, the day after the bloom, Chaetoceros growth rates dropped down to previous values, most likely due to nutrient depletion. On 1 April, concentrations of dissolved inorganic nitrogen (DIN) (nitrite and nitrate) ranged between 0.01 and 0.02  $\mu M$  N, while it increased abruptly in near-surface waters on 2 April, reaching concentrations of 0.08 to 0.1  $\mu M$  N. The day after the bloom, DIN levels decreased to 0.01 to 0.03  $\mu M$  N.

#### DISCUSSION

Daily PP and chl a concentration during the study period were the highest ever recorded in the Gulf of Eilat: 1330 mg C m<sup>-2</sup> d<sup>-1</sup> and up to 1.5  $\mu$ g chl a l<sup>-1</sup>, respectively, which is almost 3× higher than average chl a values at this time of year in this area. Our results suggest an abrupt increase in mixing. The data show a sharp increase in the biomass of diatoms and in the rates of PP over a short time. Wind direction was from the coast towards the open Gulf, and upwelling was induced. PP is usually higher in the northern and coastal stations than in the open Gulf (Iluz 1997), but in the experiment reported here, we observed that on 1 April 2008 and the following day, the rates at Stn  $A_1$ (10 km from MP) increased to a level similar to that of the coastal Stn MP. The changes were limited to the upper 40 m (Fig. 6), with no significant increase in either biomass or productivity below this depth. We assume that the increased wind speed induced the transfer of a coastal water mass from the northern shore towards the open sea. This water is more nutrient rich than that of the open sea, and thus supports

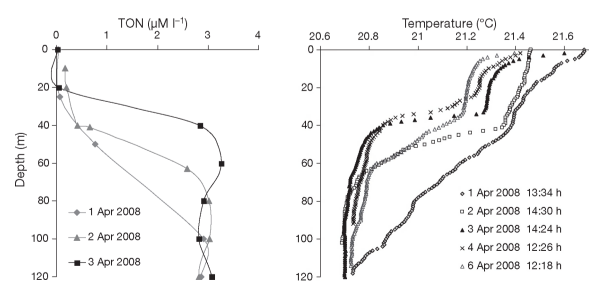


Fig. 7. (A) Profiles of total organic nitrogen (TON) on 1 to 3 April 2008, and (B) temperature on 1 to 6 April 2008, showing mixing from the surface and mixing of upwelled water at depth

the higher productivity near the northshore. This movement of the water mass is confirmed by temperature profiles (Figs. 2, 3, 7). In the period from 1 to 3 April, total organic nitrogen (TON) and temperature profiles revealed 2 current components: upwelling of colder and nutrient rich water from deep layers in the north towards the surface waters of the open sea, coupled with lateral downwelling of warm, surface waters that sink along the western shore of the Gulf and flow as a deep current towards the center of the Gulf (Fig. 1A) (Labiosa et al. 2003). Diatom abundance and biomass increased sharply over 2 d from 1 to 3 April, which accounts for the increased Si deposition by these cells and the concomitant depletion of Si in the water. The growth rates of the dominant diatom *Chaetoceros* doubled within 2 d and dropped the following day to pre-bloom values, probably due to nutrient exhaustion. As expected, the increase in PP preceded that of the biomass, which was phase-shifted by 1 d in relation to PP. Picophytoplankton are usually the main producers in the Gulf of Eilat and contribute >90 % to PP (Lindell & Post 1995); these phytoplankters probably responded rapidly to nutrient enrichment. Diatoms have a slower response than picophytoplankton, which explains the observed lag between the PP peak and maximum diatom cell density (Fig. 4).

During the reported period, the sky was cloud-free; therefore, insulation may be excluded as a factor that might have induced the changes in phytoplankton density and vertical distribution, which were attested by changes in chl *a* concentration (Fig. 5). On the other hand, wind speed changed and correlated well with water column optical properties (Fig. 4B).

In the Gulf of Eilat, the main seasonal ecological patterns are driven by mixing that usually takes place between February and April and which is characterized by deep convective mixing of the northern Gulf, which may extend down to hundreds of meters and even to the very bottom (Wolf-Vecht et al. 1992, Genin et al. 1995). Nutrients upwelled from depth and from coastal waters enrich the euphotic zone and increase PP. There are exceptional events that can raise PP, such as a high degree of mixing resulting from a particularly cold winter, as happened at the end of the winter in 1992, when the highest total daily PP rates ever recorded in the region were measured (738 mg C  $m^{-2} d^{-1}$ ). (During 1989–1997, the multi-annual average productivity rate was  $\sim 200$  mg C m<sup>-2</sup> d<sup>-1</sup>; Iluz 1991, 1997). Winters in the Gulf of Eilat are exceptionally cold, resulting in very intense deep mixing (>750 m) and increased nutrient injection from deep waters of the Gulf. As a result, a strong spring bloom develops at the onset of stratification (Gordon et al. 1994, Genin et al. 1995). Additional exceptional but local events are floods of the Shlomo River, as occurred in 1993 and which increased productivity to 604 mg C  $\rm m^{-2}~\rm d^{-1}$  (Iluz 1997).

The northern edge of the Gulf of Eilat borders the relatively populated areas of the cities of Agaba and Eilat, where anthropogenic pollutant sources such as fish farms and sewage may produce impacts on the marine environment. However, the influence of these pollution sources seems, so far, to be of a slow and gradual nature. In contrast, the event that we report here was abrupt and prominent in its effects on the biology of the water. As the situation observed was apparently wind-driven, it is expected that it may occur repeatedly in the Gulf. Northern wind events with speeds exceeding 9 m s<sup>-1</sup> occurred ~10× between 1 April 2007 and 1 May 2008, each lasting for at least 2 to 3 d. We do not know if these events had the same effects as those recorded in this report, and it is obvious that such observations should be repeated over several seasons to allow reliable estimation of their impacts on annual PP.

It is noteworthy that routine ocean monitoring programs in which PP is measured once or twice a month are likely to miss their brief but significant spikes, leading to underestimation of seasonal and annual PP values. If these values are significant for total biomass, we should consider them in our calculations, as well as study their impact on the storage and flow of energy through marine food webs and ecosystems.

Our recommendation is that, whenever possible, the sampling frequency for PP measurements should be increased using PP fluorescence calculations based on the  $F_c$ -683 method described here. Otherwise, where different automated systems tracking chlorophyll are installed, closely spaced PP measurements should be conducted when any sudden increase in pigment concentration is detected, in order to include otherwise undetected peaks. Such measurements would significantly increase and improve our annual estimates, and provide better databases for ecosystem models. Furthermore, such peaks are likely to trigger microbial and zooplankton peaks that may be missed or remain inexplicable. The advantage of the  $F_c$ -683 method is its high resolution and ease of operation that allow several measurements of depth profiles throughout the day. Such work would be nearly impossible with <sup>14</sup>C incubations.

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