

1 **The Role of Rapid Changes in Weather on Phytoplankton Spring Bloom Dynamics**
2 **Captured by an Autonomous Uncrewed Surface Vehicle**
3

4 **G. M. Fragoso^{1,2†}, A. Dallolio³, S. Grant^{1,2}, J. L. Garrett³, I. Ellingsen⁴, G. Johnsen^{1,2,5} and**

5 T. A. Johansen^{2,3}

6 ¹Trondheim Biological Station, Department of Biology, Norwegian University of Science and
7 Technology (NTNU), Trondheim, Norway.

8 ²Centre of Autonomous Marine Operations and Systems (AMOS), NTNU, Trondheim, Norway.

9 ³Department of Engineering Cybernetics, NTNU, Trondheim, Norway.

10 ⁴SINTEF Ocean, Dept. Env. & New resources, Trondheim, Norway.

11 ⁵University Centre in Svalbard (UNIS), Longyearbyen, Norway.

13 Corresponding author: Glauzia M. Fragoso (glaucia.m.fragoso@ntnu.no)

14 † Bynesveien 46, 7018, Trondheim, Norway.

16 **Key Points:**

- 18 • The spring bloom in coastal high latitudinal regions consisted of multiple peaks
19 associated with gain and loss processes.
- 20 • Relaxation of strong winds and clear skies for 7-10 days in spring allowed phytoplankton
21 accumulation and bloom development.
- 22 • Episodic strong winds interluded spring bloom development after a period of calm, sunny
23 weather.

25 **Abstract**

26

27 The spring phytoplankton bloom plays a major role in pelagic ecosystems; however, its
28 dynamics is overlooked due to insufficient, highly-resolved observational data. Here we
29 investigate the start, peak and decline of a two-week phytoplankton spring bloom in Frohavet,
30 located at the coast of mid-Norway. We used observations from an uncrewed surface vehicle
31 (USV) combined with buoy measurements, satellite images, discrete water sampling and
32 modelling approaches. The spring bloom (March-June 2022) consisted of multiple peaks (up to 5
33 mg m⁻³), with a long peak in April, coincident with the period when the USV captured the
34 temporal and spatial dynamics of the bloom. Short-term (5 days) episode of calm weather in the
35 spring, such as clear skies and consistent low wind speed (< 7 m s⁻¹) shoaled the mixed layer
36 depth (< 15 m), after strong wind speed (average wind speed up to 20 m s⁻¹ in March) and
37 mixing events in winter. These rapid changes in the environment promoted the rapid
38 development of the spring bloom - from 1 to 5 mg m⁻³ in 5 days. Likewise, the collapse of the
39 bloom was rather quick, 1-2 days and coincides with low nitrate values and rapid increase in
40 wind speed (> 10 m s⁻¹), suggesting strong influence of the environment on phytoplankton
41 dynamics during early stages of the spring bloom. Understanding the dynamics of the spring
42 bloom is crucial for the management of marine resources. Integration of distinct observational
43 platforms has the potential to unveil the environmental factors underlying phytoplankton bloom
44 dynamics.

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46

47 **Plain Language Summary**

48

49 The phytoplankton spring bloom is an important recurrent phenomenon because it provides food
50 for the marine food web and regulates the climate. Although previous studies were focused on
51 the initiation of the spring bloom, its dynamics, meaning, rapid changes in formation and decline,
52 are usually not observed in detail. Here we used a combination of a technological (marine
53 sensors and robots) and traditional methods (water collection, laboratorial and microscopic
54 analyses) to observe the spatial and temporal variation of the spring bloom in a biological
55 hotspot of the coast of mid-Norway. Small windows of ‘good weather’, where few days of
56 sunny, clear skies and weak winds in the midst of ‘stormy spring’ promoted the rapid
57 development of the spring bloom dominated by the diatom *Skeletonema*. The bloom collapsed
58 after the wind speed got high again, suggesting the strong influence of environmental conditions
59 in the spring bloom. Here we demonstrated that the use of multiple ocean observation platforms
60 is crucial to understand, in detail, the processes controlling the spring phytoplankton bloom.

61

62 **Index terms**

63 4855 Phytoplankton 4894 Instruments, sensors, and techniques 4271 Physical and chemical
64 properties of seawater

65

66 **Keywords:** phytoplankton dynamics, environmental controls, phytoplankton spring bloom, non-
67 photochemical quenching, uncrewed surface vehicles.

68

69 **1 Introduction**

70

71 The spring phytoplankton bloom is a key event in the annual cycle of phytoplankton
72 abundance in high latitudinal seas (Chiswell et al., 2014; Rumyantseva et al., 2019). As a
73 recurrent seasonal phenomenon, the spring bloom plays a major role in pelagic ecosystems,
74 contributing to carbon export and sequestration, oxygen production and energy flow for higher
75 trophic levels (Alkire et al., 2014). Phytoplankton spring bloom are also sentinels of climate
76 change, where alterations in intensity and phenology have been observed (Edwards &
77 Richardson, 2004), with predictions of a continuous shift by the end of the century (Henson et
78 al., 2018; Yamaguchi et al., 2022)

79 In simple terms, phytoplankton spring blooms consist of positive biomass accumulation
80 rate (r) over a period of time, where growth (μ , e.g. cellular division) surpasses loss processes (l ,
81 e.g., grazing and sinking rates, viral lysis) ($r = \mu - l$, $r > 0$). While there is a scientific consensus
82 that spring blooms consist of accumulation of phytoplankton, the processes that determine their
83 start and the net balance between μ and l , are still on debate (Mojica et al., 2021). Moreover,
84 there is also a debate of what constitute a bloom, whether accumulation is explosive (rapidly
85 increase in r) or steadily increasing over time (Mignot et al., 2018). Historically, $r > 0$ in spring
86 was viewed as a bottom-up process, where l remained unchanged and μ increased rapidly as light
87 becomes a non-limiting factor with shoaling of the mixed layer (Sverdrup, 1953). Recently, this
88 hypothesis has been contested, where low l due to low grazing rates as a consequence of low
89 encounter rates of phytoplankton and zooplankton, rather than a fast change in μ in spring, would
90 account for slow $r > 0$ in winter (Behrenfeld, 2010; Behrenfeld et al., 2013; Behrenfeld & Boss,
91 2014). These authors also suggested that tight recoupling between grazers and phytoplankton
92 occur when the mixed layer start to become shallow, with a slight lag (1 day) in time, where l ,

93 the deterministic factor for $r > 0$, would eventually catch up with μ after an early stock of
94 phytoplankton biomass in winter (Behrenfeld & Boss, 2014).

95 The debate around spring bloom is mostly focused on their initial stage, although, in
96 reality, these such blooms are quite dynamic in high latitudinal and polar seas (Behrenfeld et al.,
97 2017). This means that weather changes (e.g. cloud cover, wind speed), particularly in spring, are
98 rather fast (< 1 day), where the volatile nature of these blooms consists of multiple peaks with
99 rapid formation and collapse over the course of few days or weeks. Thus, many studies ignore
100 the influence of ‘small windows of good weather’, where few days of sunny, clear skies and
101 weak winds during a ‘stormy spring’ might have a huge impact on phytoplankton dynamics.

102 Central to this problem is the lack of sufficient, highly-resolved observational data,
103 particularly for regions that are highly dynamic and complex. For many years, spring blooms
104 have been studied using satellites, which close progression of blooms are missed due to cloud
105 cover often occurring in high latitudinal seas (Behrenfeld, 2010). Discrete water and net
106 sampling over a long-term time series provides detailed information of plankton species
107 (González-Gil et al., 2022); however, microscopic analyses are time-consuming and, in many
108 times, not highly resolved in space and time. Fixed platforms, including buoys, can provide high
109 temporal resolution (< 1h), however, spatial patchiness is often be missed (Son et al., 2014).
110 Mobile platforms, including profiling floats (Boss & Behrenfeld, 2010; Mignot et al., 2018) and
111 gliders (Rumyantseva et al., 2019) are suitable, but only for deep, open waters (> 200 m) due to
112 the potential collision to a shallow seafloor. For coastal regions, with shallow and irregular
113 bathymetry, autonomous uncrewed surface vehicles (USV) can offer an affordable, flexible
114 solution for studying phytoplankton spring bloom progression, although observations comprise
115 only surface waters (Dallolio et al., 2021; Scott et al., 2020). Modelling approaches can offer

116 complementary information regarding areas not covered by satellites and USVs and can also
117 provide the vertical structure of the water column. An “observational pyramid” for ocean
118 systems, which combines the integration of in-situ autonomous platforms, fixed buoys, satellite
119 imagery and modelling approaches with discrete net and water sampling have the capacity to
120 unveil the dynamics of a phytoplankton spring bloom in a coastal, productive hot-spots (Fragoso
121 et al., 2022; Williamson et al., 2023).

122 In-situ chlorophyll *a* fluorescence (*FChla*) measurements derived from sensors attached
123 to autonomous or fixed platforms have been historically used as a proxy of phytoplankton
124 biomass in studies of bloom dynamics (Roesler et al., 2017). However, *FChla* measurements are
125 only useful when they reflect the true concentration of chlorophyll *a* [*Chla*] in the water.
126 Systematic errors, such as biofouling and non-photochemical quenching (*NPQ*) influence the
127 fluorescence signal to intracellular *Chla* pigment ratios, offering biased measurements (Carberry
128 et al., 2019; Johnsen et al., 2018; Scott et al., 2020). *NPQ* is a physiological response of live cells
129 to high light (usually observed in surface waters at daytime), where the excess of energy is
130 converted to heat, reducing the *FChla* signal (Huot & Babin, 2010). Thus, it is crucial that, for
131 surface *FChla* measurements derived from USVs, values are corrected for *NPQ*, particularly
132 during spring and summer.

133 Here, we use an USV equipped with environmental and weather sensors (fluorometer,
134 CTD, oxygen optode and weather station) to investigate the dynamics (start, peak and decline) of
135 a 2-week phytoplankton spring bloom in Frohavet, a coastal Norwegian biological hotspot. To
136 complement the USV data (resolved to 1 min binned), a combination of other observational
137 methods, including a fixed mooring buoy, satellite images, discrete water sampling and
138 modelling approaches were included. For *FChla* from the USV, we provide a solution for *NPQ*

139 for daily changes and investigate the biophysical controls of the bloom. Discrete water samples
140 for nutrient concentrations and phytoplankton abundances, in addition to *FChla* measurements
141 from a moored buoy, were collected from a fixed station ~25 km away from the trajectory of the
142 USV AutoNaut. Integration of distinct observational platforms, such as autonomous vehicles,
143 fixed buoys, discrete water sampling and modelling approaches has the potential to unveil the
144 environmental factors underlying phytoplankton bloom dynamics.

145 **2 Materials and Methods**

146

147 **2.1 Study area**

148

149 Frohavet is a wide, open stretch of sea surrounded by a large cluster of small islands on
150 the coast of Trøndelag, mid-Norway (Fig. 1). This region sustains high levels of primary
151 production and biological diversity and is a popular site for aquaculture activities. Frohavet is
152 highly productive because of the Norwegian Atlantic Water (NAW), which brings nutrient-rich
153 Atlantic Water (AW) along the shelf break. This water mass is located below the Norwegian
154 Coastal Current (NCC), which becomes thicker as freshwater input increases from spring to
155 summer (Fragoso et al., 2019). The NAW often reaches the surface through coastal upwelling
156 and internal waves (Fragoso et al., 2019). Diatoms are known to be the predominant
157 phytoplankton of the spring bloom (Fragoso et al 2021, Thu et al 2021), however, dinoflagellates
158 often co-occur. Diatoms blooms sustain high zooplankton (particularly copepods) abundance
159 predominantly found in this region (Fragoso et al., 2019).

160

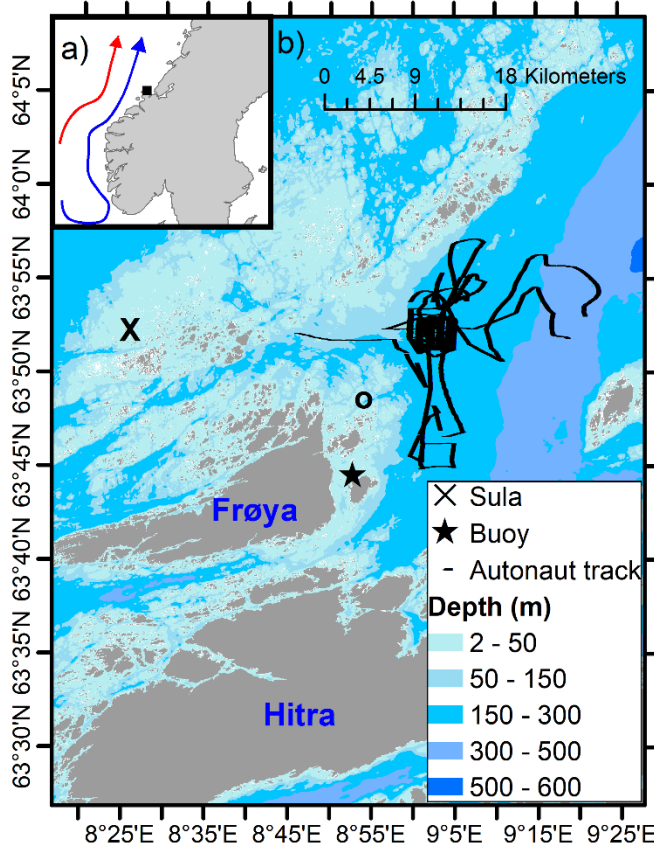


Figure 1- Study site in the coast of mid-Norway. a) Scheme showing the main currents flowing northwards – the Norwegian Coastal Current (blue) and the Norwegian Atlantic Current (red). b) Map of Frohavet region showing the islands of Frøya and Hitra, in addition to the weather station in Sula Island (cross symbol), the fixed buoy station near Frøya (star symbol) and the site where mixed layer depth was modelled by SINMOD (circle symbol).

2.2 Buoy and water sampling

To monitor the start and development of the bloom in the region of Frohavet, a C3 submersible fluorometer sensor (Turner Designs, USA) was attached into a buoy located about 2 km east of the coast of Frøya Island (Fig. 1) and placed at 4 m depth (Fig. 1). This sensor collected temperature ($^{\circ}\text{C}$), FCChla (calibrated later to concentration in mg m^{-3}) and turbidity (Relative Fluorescence Unit - calibrated later to Formazin Turbidity Unit (FTU)) every 10 min from mid-February to mid-June. Hourly wind speed (m s^{-1}) data (from February until mid-June) from Sula

177 meteorological station (located in the western part of Frohavet) were collected from the
178 Norwegian Weather Service Center (<https://seklima.met.no/>). A HOBO pendant temperature and
179 light logger (HOBO, USA) was placed at the top of the buoy to measure light intensity in air
180 (measurement in lux at every 30 min from February until June). Values were integrated daily and
181 converted to photosynthetic active radiance ($\mu\text{mol photons m}^{-2} \text{ s}^{-1}$) by using a conversion factor:
182 1 klux (kilolux)= 14-18 $\mu\text{mol photons m}^{-2} \text{ s}^{-1}$ (Sakshaug et al., 2009).

183 Discrete water sampling for nitrate and *in vitro* chlorophyll concentrations ($[\text{Chla}_{\text{in-vitro}}]$) was
184 collected at 3 m depth and every 2-3 weeks from mid-February to mid-June few meters away from
185 where the C3 sensor was attached. For nitrate analyses, triplicate water samples were filtered with
186 a 0.8 μm polycarbonate filter, where the filtrate water was immediately kept in a centrifuge tube
187 and stored frozen at -20°C. Nitrate analyses were determined in the laboratory using a continuous
188 flow automated analyzer (CFA, Auto-Analyzer 3, SEAL). For $[\text{Chla}_{\text{in-vitro}}]$, seawater was filtered
189 (0.25-0.5 L) onto a Whatman GF/F glass fiber filters, and immediately double-folded, wrapped in
190 aluminum foil and stored at -20°C for *a posteriori* analyses in the laboratory. For a fluorometric
191 determination of $[\text{Chla}_{\text{in-vitro}}]$, frozen filters were placed in glass vials with 100% methanol for few
192 hours on a dark fridge at 10°C. For $[\text{Chla}_{\text{in-vitro}}]$, the extracted solvent was determined using the
193 Turner Designs Trilogy fluorometer (model: 7200-000) and the non-acidification method (Fragoso
194 et al., 2019). For phytoplankton identification and quantification, water samples were fixed with
195 neutral Lugol's iodine solution to a final concentration of ~1% into dark amber bottles and stored
196 at room temperature and in the dark for later microscopic analyses in the laboratory.

197 2.3 USV AutoNaut sampling

198

199 The AutoNaut is a commercially available USV that relies on sea surface waves to
200 produce forward thrust, making it suitable for sustained operations at sea without human
201 assistance. Along with the instrumentation needed for navigating autonomously, the vehicle
202 carries several scientific payload integrated on the USVs hull and keel (between 25 and 50 cm
203 approximately below the waterline) that collects information about a wide range of
204 environmental variables. Among these, we had an Eco Triplet sensor (Wet Labs, Oregon, USA)
205 to measure *FChla* ($\lambda_{\text{ex}}=470 \text{ nm}$, $\lambda_{\text{em}}=695 \text{ nm}$), turbidity ($\lambda=700 \text{ nm}$) and fluorescence of
206 colored dissolved organic matter (*CDOM*, $\lambda_{\text{ex}}=370 \text{ nm}$, $\lambda_{\text{em}}=460 \text{ nm}$). A manufacturer
207 calibration factor converted the units of *FChla* to $[\text{Chla}] (\text{mg m}^{-3})$, *CDOM* to ppb and turbidity to
208 Nephelometric Turbidity Units (NTU). In addition, the USV was equipped with a Seabird CTD
209 SBE 49 (sampling rate of 16 Hz) for measurements of temperature and salinity and an oxygen
210 Optode 4835 (Aanderaa) for oxygen concentration (μM , later converted to mg L^{-1}). Finally, a
211 weather station (Airmar 220WX) for measurements of wind speed was located on the mast of the
212 USV. The USV was deployed from the Mausund field station (located north of Frøya island, in
213 the south-western part of Frohavet, Fig. 1) on 12th April 2022 and navigated in Frohavet until
214 28th April 2022. The log files of each sensor were converted to CSV format and merged in
215 Python. Data was binned for every minute.

216 2.4 Non-photochemical quenching correction

217

218 *In vivo* (in situ) *FChla* is commonly used as a proxy for phytoplankton biomass.
219 However, when phytoplankton cells are usually exposed to high irradiance (particularly around

220 noon), excess energy is dissipated as heat, reducing the *FChla* signal in the water (Travers-Smith
221 et al 2021). This photophysiological process is called non-photochemical quenching (*NPQ*)
222 (reviewed by Brunet et al., 2011)). To visualize the overall trend of corrected *FChla*, the
223 beginning of each *NPQ*-induced trough was connected linearly to the end of the trough, resulting
224 in a straight line over the period affect by *NPQ*. This method excludes the *FChla* suppressed by
225 *NPQ* and ignores small variability of *FChla* during the day, however, it suits well to show
226 general trends of bloom dynamics (before, peak and post-bloom) in April 2022.

227

228 2.5 Satellite observations

229

230 Sentinel-3 images from OLCI (Ocean and Land Colour Instrument) sensor (multispectral
231 imager) are used to provide an overview of the sampling region, and to assess whether the USV
232 is observing spatial or temporal variations in the *FChla*. To determine the daily pattern of *Chla*,
233 all the Sentinel-3 *Chla* maps which contain the target region during the day were downloaded
234 and merged. The images are resampled to the same spatial grid and averaged. Pixels which
235 displayed land or clouds were excluded from the averaging process.

236

237 2.6 Mixed Layer Depth

238

239 Mixed layer depth was calculated from data from SINMOD, a 3D ocean model system. The
240 model has been established for the coastal region outside mid-Norway with a high spatial
241 resolution (800m). SINMOD's hydrodynamic component utilizes the primitive Navier-Stokes
242 equations to calculate ocean properties such as water current, velocity, water temperature, and

243 pressure (see Slagstad & McCliman, 2005 for more details on the hydrodynamic module). The
244 model is established on a z/z*-grid with fixed horizontal grid size and nesting for high spatial
245 resolution. The North Atlantic and Arctic region are modeled with a 20 km resolution, which
246 produces boundary conditions for a higher resolution configuration of 4 km horizontal grid size
247 for the Nordic Seas and further to 800 m grid size for the coastal region outside Mid-Norway.
248 The 20 km model uses specified boundary conditions, including 12 tidal components at open
249 boundaries, with data imported from the TPXO tidal model for global ocean tides. Interpolated
250 ERA5 atmospheric data from ECMWF (for more details see Hersbach et al. (2020) is used to
251 force the ocean model domain, including 3 hourly wind forcing, air pressure, and daily air
252 temperature, humidity, and cloud cover to calculate heat exchange. Norwegian freshwater
253 discharges from rivers and land are applied using data from simulations by the Norwegian Water
254 Resources and Energy Directorate with a version of the HBV-model (Beldring et al., 2003),
255 while historic data from SMHI Hype model data is used for other European rivers (<https://hypeweb.smhi.se/>). For more information, refer to Hersbach et al. (2020). Mixed layer
256 depth is calculated from the SINMOD output as the depth at which the density gradient exceeds
257 0.01 kg m⁻¹.
258

259 **3 Results**

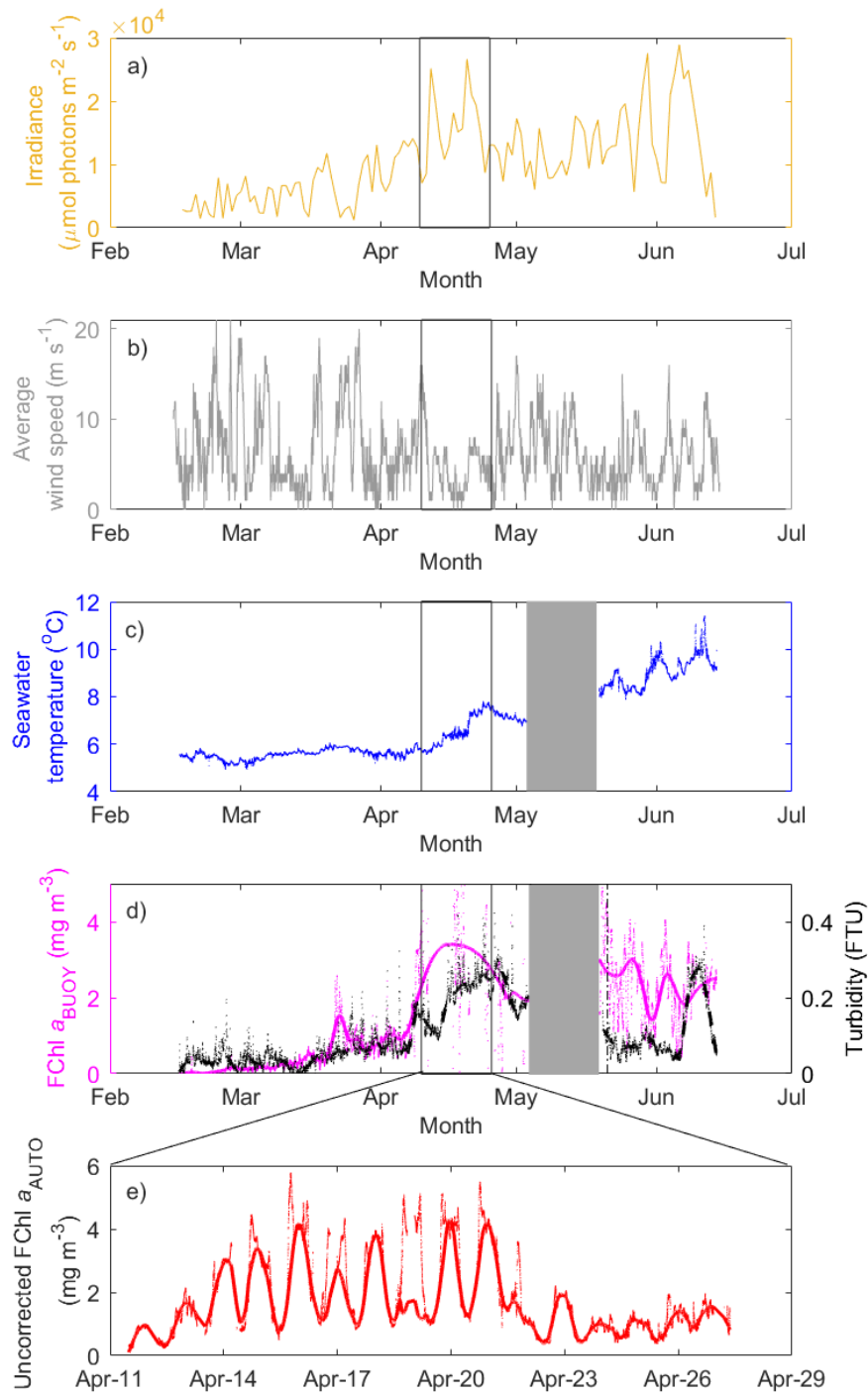
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261 3.1 Mooring buoy data

262

263 Integrated daily irradiance above water from HOBO light loggers (converted from klux to
264 $\mu\text{mol photons m}^{-2} \text{s}^{-1}$, see methods) gradually increased from mid-February to mid-June, showing
265 peak in values (up to $2 \times 10^4 \mu\text{mol photons m}^{-2} \text{s}^{-1}$) from early April until early May (Fig. 2a).

266 This suggests a period of consistent clear and sunny skies in April for at least 2 weeks. Average
267 wind speed varied in the region of Frohavet from February until mid-June (Fig. 2b). Average
268 wind speed was particularly strong (up to 20 m s^{-1}) during March and early April (Fig. 2b). From
269 mid-April until late April (time when the USV AutoNaut was in Frohavet, Fig. 2e), average wind
270 speed was relatively weak for several days ($< 5 \text{ m s}^{-1}$) compared to March. Concomitantly,
271 seawater temperature from a buoy ($\sim 3 \text{ m}$ deep) located near the coast of Frøya island (Fig. 1)
272 gradually increased from early April until late June and shortly increased from ~ 6 to 8°C in mid-
273 April (Fig 2c). Chlorophyll a concentrations ($FChla_{BUOY}$) and turbidity from the buoy station
274 near Frøya island (Fig. 1) gradually increased from mid-March and peaked from mid to late
275 April. The spring bloom (March-June) consisted of multiple peaks (Fig. 2d) - a short peak in late
276 March where the haptophyte *Phaeocystis* sp. dominate the bloom and a second long peak during
277 April where diatom *Skeletonema* spp. is the dominant genus (Table S1. Supplementary material).
278 The USV AutoNaut was located in Frohavet before, during and after the *Skeletonema* bloom in
279 April (Fig. 2e), capturing the temporal and spatial dynamics of the bloom.
280



281

282 **Figure 2- Time series of environmental parameters and the spring phytoplankton bloom.** a)
283 integrated daily irradiance ($\mu\text{mol photons m}^{-2} \text{s}^{-1}$), b) average wind speed (m s^{-1}), c) temperature
284 ($^{\circ}\text{C}$), d) *in vivo* chlorophyll (mg m^{-3}) and turbidity (FTU) from the stationary buoy from mid-
285 February until mid-June 2022 and e) uncorrected *in vivo* chlorophyll from the AutoNaut from
286 April 11th – April 28th. Grey box in b) and c) represents lack of data due to malfunctioning of the
287 instrument. For d) and e) dot represent the median-calculated data from seven consecutive runs
288 and line represents the smoothing parameter (*rloess* method in Matlab).

289

290 3.2 Daily variation of $FChla_{AUTO}$

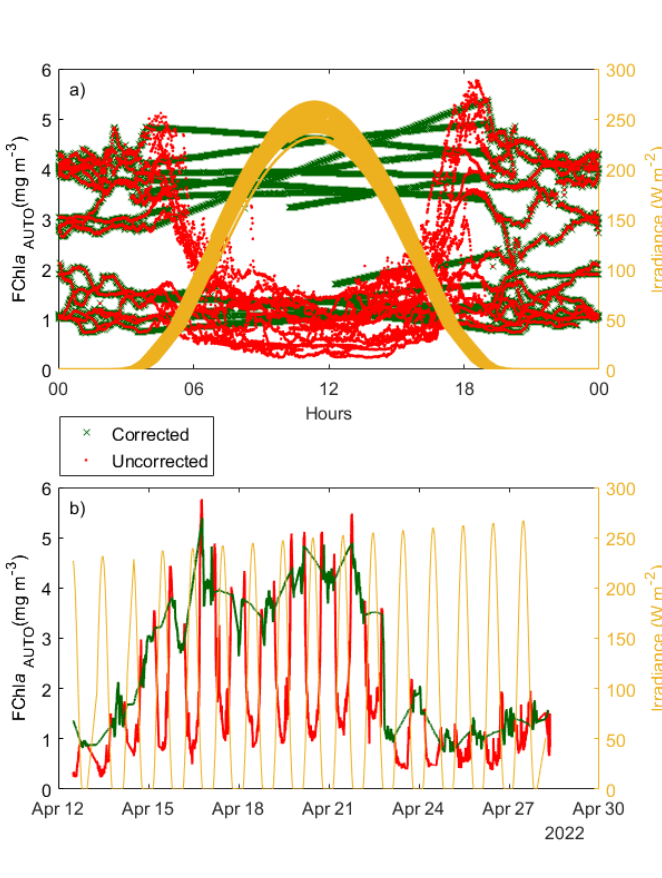
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292 $FChla_{AUTO}$ (mg m^{-3}) varied with time, with low values around 12th April (<1 mg m^{-3}), increasing
293 gradually until 16th April, reaching maximum average values $\sim 4 - 5 \text{ mg m}^{-3}$ and decreasing
294 again on the 23rd – 24th April (average $\sim 1 \text{ mg m}^{-3}$) (Fig. 2d). $FChla_{AUTO}$ (mg m^{-3}) varied also as a
295 function of irradiance during the day cycle, showing low values when irradiance is the highest
296 (around noon) (Fig. 3a). Such low $FChla_{AUTO}$ daylight values occurred due to NPQ . To correct
297 for this values, linear interpolation of night $FChla_{AUTO}$ was used, to show the robust trends in
298 [Chla] chlorophyll concentration before, during and after the bloom (Fig. 3b).

299

300

301



302

303 **Figure 3- Daily and temporal variations of *in situ* chlorophyll a fluorescence ($FChla_{AUTO}$
304 (mg m^{-3})) and modelled irradiance. a) Time series and b) daily cycle of $FChla_{AUTO}$ uncorrected
305 (red) and corrected for non-photochemical quenching (green) from the USV AutoNaut and
306 modelled daily irradiance (yellow) derived from 12th– 28th April 2022.**

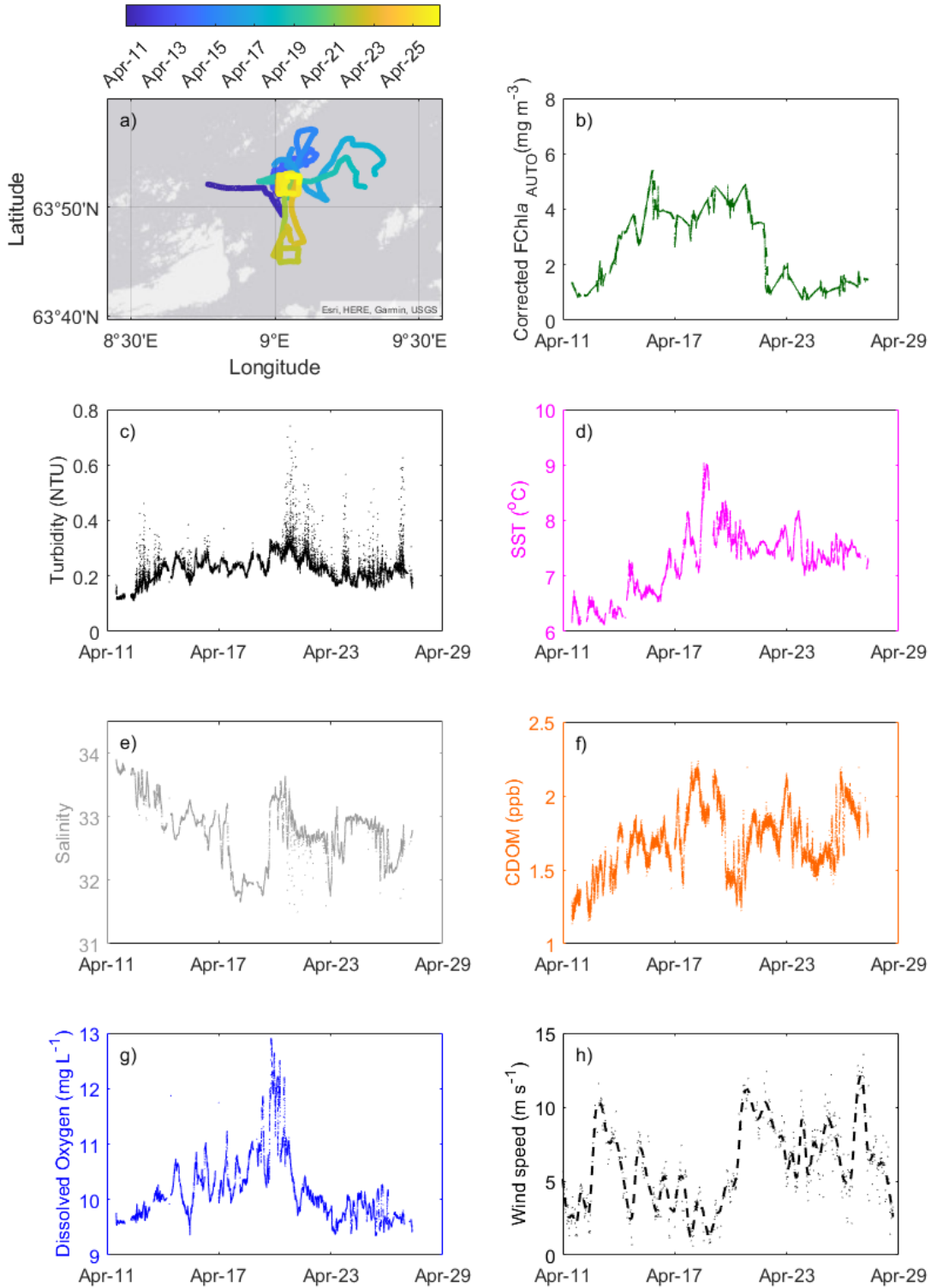
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308 3.3 USV AutoNaut

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310 The USV AutoNaut was present in surface waters of Frohavet from April 11th – April 27th (Fig.
311 4a). The USV departed from a small island, Mausund, north of Frøya island and moved towards
312 the deep waters of Frohavet. The USV initially moved out of Mausund Island and northwards
313 around 15th April, along Froan archipelago, then east around 19th April, then south (east of
314 Frøya) on April 23rd, completing its mission around the center, deep waters of Frohavet (Fig. 4a).
315 Corrected *FChla_{AUTO}* (mg m^{-3}) gradually increased from April 11th, reaching its peak (up to 5.5
316 mg m^{-3}) on ~ April 16th until ~ April 22nd when it abruptly declined, and reaching low values (<
317 2 mg m^{-3}) (Fig. 4b). Average turbidity values were lower than 0.2 (NTU) from April 11th – April
318 12th, slightly increasing during the period of the bloom (average > 0.2 NTU). Turbidity values
319 were less ‘noisy’ from 14th - 20th April and gradually declined until 23rd April, becoming again
320 noisier and around 0.2 NTU afterwards. Sea surface temperature (SST), colored dissolved
321 organic matter (CDOM) and dissolved oxygen (DO) gradually increases from April 11th until
322 April 18th (~ 6 - 9°C for SST, 1.1-2.2 ppb for CDOM and 9.7 - 13 mg L^{-1} for DO).
323 Concomitantly, salinity and average wind speed (m s^{-1}) decrease from 34 to 32 and from 10 m s^{-1}
324 until 3 ms^{-1} , respectively. Salinity sharply increases from 32 to 33 and CDOM and SST abruptly
325 decreases on April 19th, from 2.1 to 1.2 ppb and 8.9-7.5, respectively. At the same period
326 (particularly on April 20th, wind speed peaks up, reaching > 10 m s^{-1}). From April 18th onwards
327 (until April 26th), SST, CDOM, salinity and wind speed fluctuate. DO reach highest values on
328 April 19th and 20th and sharply goes down to < 10 mg L^{-1} after April 23rd.

329



330
331 **Figure 4- Track of the USV AutoNaut and the environmental parameters collected.** a)
332 Frohavet region and the track line of the USV AutoNaut from 11th – 28th April. Time series of b)
333 corrected chlorophyll a fluorescence ($F\text{Chla}_{\text{AUTO}}$ (mg m^{-3})), c) turbidity (NTU), d) sea surface
334 temperature (SST ($^{\circ}\text{C}$)), e) salinity, f) colored dissolved organic matter (CDOM (ppb)), g)
335 dissolved oxygen (mg L^{-1}), and h) wind speed (m s^{-1}) from the USV AutoNaut.

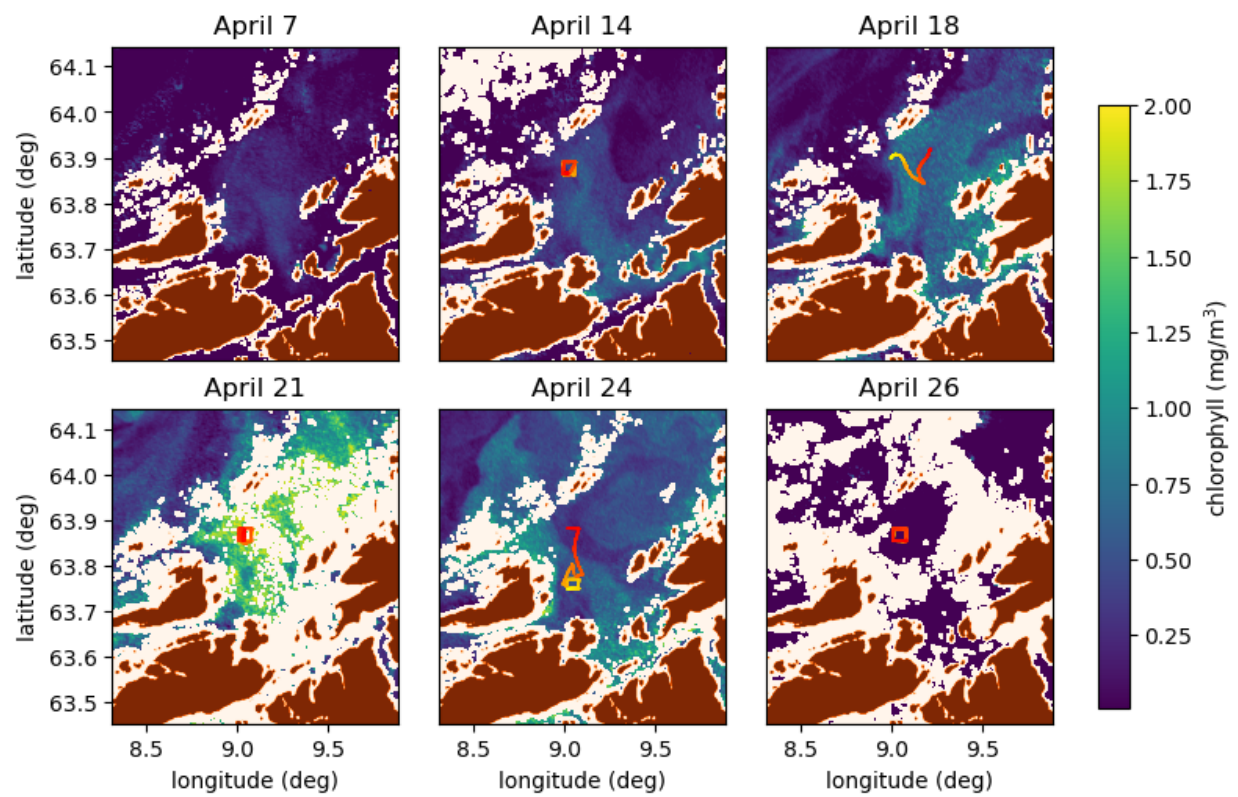
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337 3.4 Satellite images and USV AutoNaut track

338

339 To verify whether the bloom variability detected by the USV AutoNaut is temporal (e.g.
 340 start, peak and collapse) or spatial (patchiness), daily satellite images along with the USV tracks
 341 were analyzed together (Fig. 5). According to satellite images and, in alignment with the
 342 observations from the USV, the bloom started from ~ April 7th – 18th, peaked ~ April 21st,
 343 declined on April 24th, where the USV was confined in relatively low [Chla] regions (< 1 mg m⁻
 344 ³) and collapsed on April 26th (Fig. 5).

345



346

347 **Figure 5 - Satellite images (Sentinel 3) of chlorophyll α concentrations (mg m^{-3}).** Spatial and
 348 temporal variability of [Chla] in Frohavet region on April 7th, 14th, 18th, 21st, 24th and 26th. Note
 349 the tracking path of the USV AutoNaut for each day, during early (yellow) to late (red) hours.

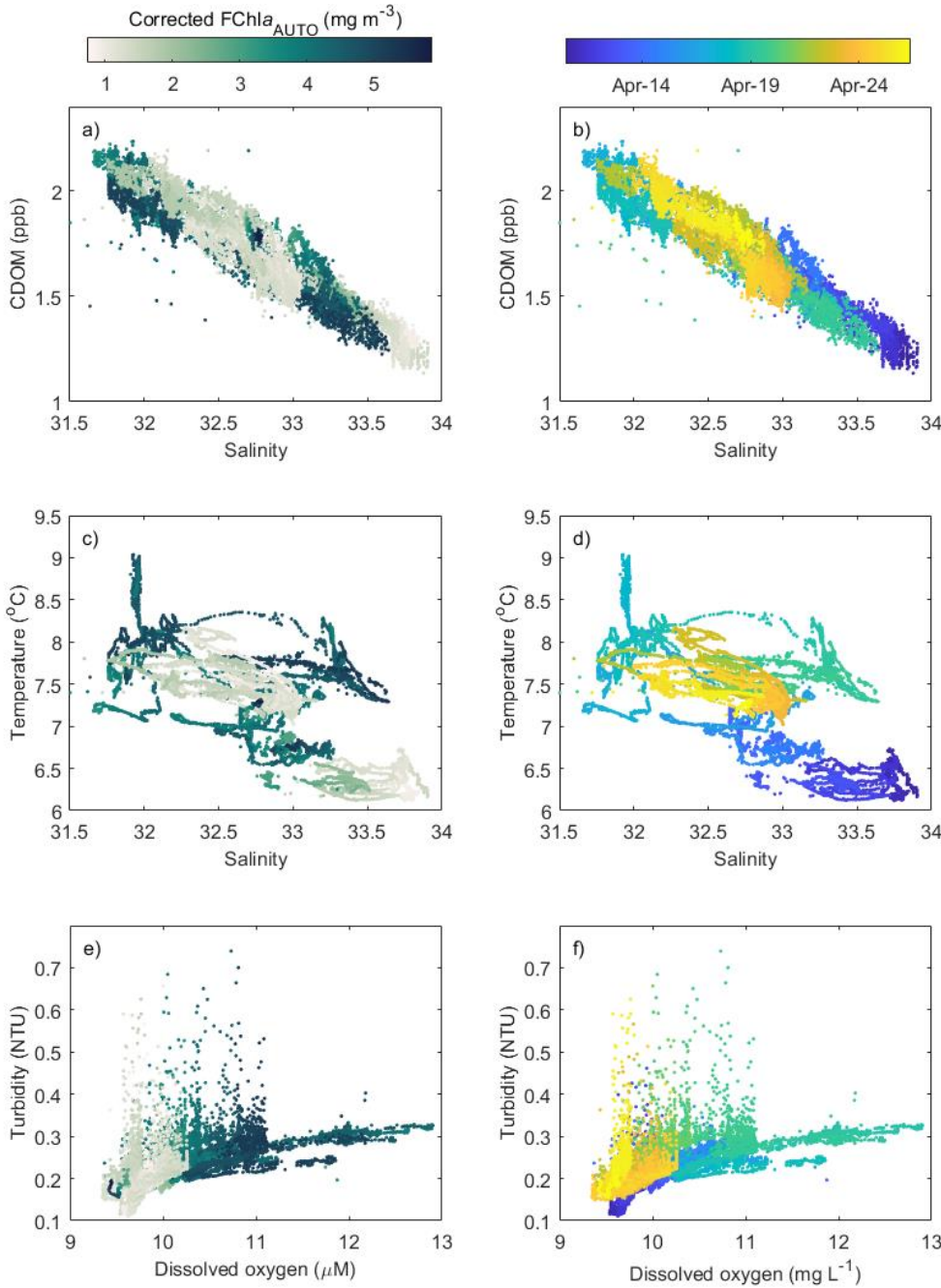
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351 3.5 Correlation of USV AutoNaut parameters

352

353 The phytoplankton bloom observed as corrected $FChla_{AUTO}$ seemed, in general, not to be
354 concentrated within a certain water mass, being widely present in waters with low and high
355 salinity and temperature values (Fig. 6a and 6c). The phytoplankton bloom seemed to, rather,
356 have a temporal trend, starting from 12-13th April and ending on 21st April. Salinity and CDOM
357 presented a negative relationship, changing progressively in values in the early phase of the
358 bloom (Fig. 6a and 6b). Temporally, the USV AutoNaut traveled waters with a wide range of
359 salinity and temperature in the beginning of the mission, concentrating later in the center of
360 Frohavet (after April 19th – salinities are between 32 and 33) (Fig. 6d). Lowest daily turbidity
361 values had a positive relationship with DO and chlorophyll (Fig 6e and 6f). Waters with high
362 dissolved oxygen levels ($> 11 \text{ mg L}^{-1}$) had less variability (which coincided with a period of
363 weak winds and high chlorophyll concentrations) (Fig. 4c and 4h).

364



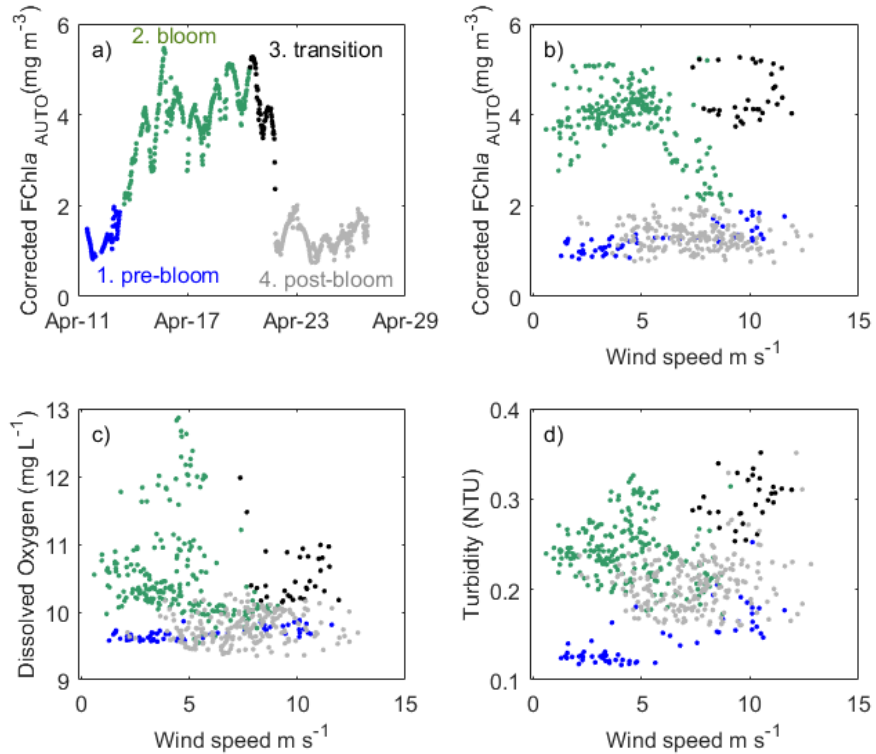
365

366 **Figure 6- Correlation plots between parameters from the USV AutoNaut.** Correlations
 367 between a,b) salinity and coloured dissolved organic matter (CDOM (ppb)), c,d) salinity and
 368 temperature (°C) and e,f) dissolved oxygen (mg L⁻¹) and turbidity (NTU) as function of
 369 *FChla_{AUTO}* (left) and time (right) from the USV AutoNaut.
 370

371

372 Values of corrected $FChla_{AUTO}$ varied as a function of wind during the period that the
373 USV AutoNaut was in Frohavet (Fig. 7). During a pre-bloom condition (~April 11th – 13th),
374 average wind speed was variable (from 1-10 m s⁻¹), followed by a blooming period (> 2 mg m⁻³),
375 where corrected $FChla_{AUTO}$ values from the USV AutoNaut rapidly increase in a few days
376 (~April 13th – 15th). During the blooming period average wind speed was low (< 6 m s⁻¹), while
377 DO and turbidity were high (> 10 mg L⁻¹ for DO and > 0.2 NTU for turbidity) (Fig 7c,d). After
378 this period, here referred as ‘transitional period’, winds started to pick up (average speed from 7
379 to 13 m s⁻¹), while chlorophyll concentration and turbidity were still high (> 4 mg m⁻³ for
380 corrected $FChla_{AUTO}$ and 0.25 NTU for turbidity) (Fig. 7). The transitional period was followed
381 by a post-bloom period, where chlorophyll values were low (< 2 mg m⁻³) but average winds were
382 still high (from 5 – 13 m s⁻¹) (Fig. 7b).

383



384
385 **Figure 7- Correlation plots of parameters from the USV AutoNaut.** Correlation plots
386 between a) year day and corrected chlorophyll fluorescence ($FChla_{AUTO}$ (mg m^{-3})), b) wind speed
387 (m s^{-1}) and corrected chlorophyll fluorescence ($FChla_{AUTO}$ (mg m^{-3})), c) wind speed (m s^{-1}) and
388 dissolved oxygen (mg L^{-1}) and d) wind speed (m s^{-1}) and turbidity (NTU).
389

390

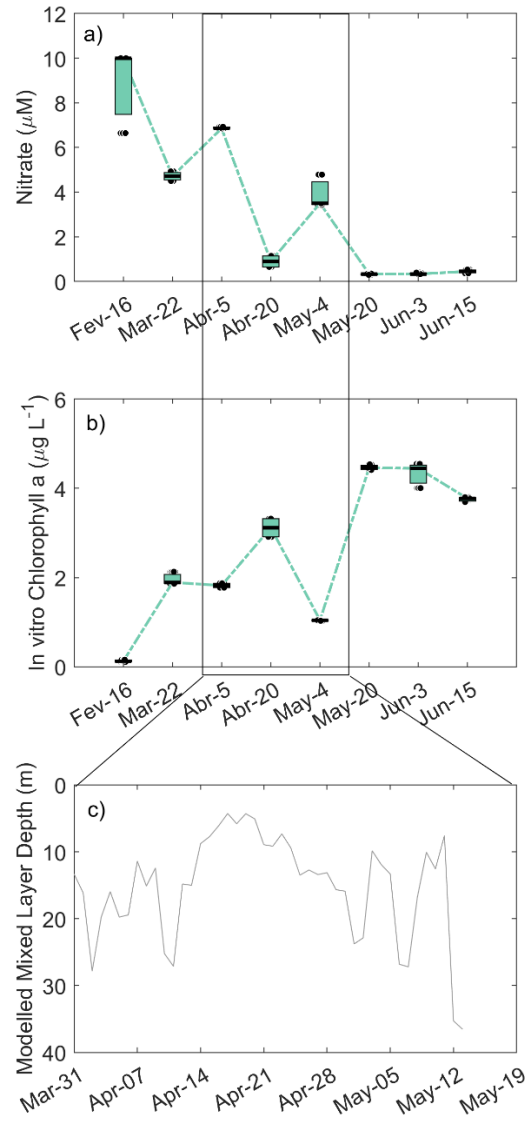
391 3.6 Discrete water sampling and modelled mixed layer depth

392

393 Nitrate concentrations from discrete water samples collected at the coast of Frøya, where the
394 buoy is located (Fig. 1), showed an overall continue decline from mid-February until mid-June
395 with pulses of nutrient occurring on April 5th and May 4th (6 and 4 µM, respectively, Fig. 8a).

396 Values of [*Chla*_{in-vitro}] increased with time, with a decline on May 4th (< 2 µg L⁻¹, Fig. 8b), which
397 coincided with the post bloom period observed after the USV AutoNaut was in the water.

398 Modelled mixed layer depth was shallow (< 15 m), particularly during the period of bloom
399 development, when the USV AutoNaut was in the water (Fig. 8c).



400
401 **Figure 8- Nitrate, in vitro chlorophyll *a* concentrations and mixed layer depth as a function**
402 **of time.** Time series of a) nitrate concentrations (μM) and b) *in-vitro* chlorophyll *a* from discrete
403 water samples collected at 3 m depth from near Frøya island ($\mu\text{g L}^{-1}$). c) Modelled average
404 mixed layer depth (MLD) from 31st March – May 13th derived from SINMOD simulations.
405 Location of Frøya and regions of modelled MLD simulation is shown in Figure 1 (circle
406 symbol).

407

408 **4 Discussion**

409

410 4.1 Non-photochemical quenching

411

412 In vivo $FChla$ signal varied widely in this study as a function of daily irradiance. Non-
413 photochemical quenching clearly affected the $FChla_{AUTO}$ signal during daytime, appearing low
414 when irradiance is high (peak at noon), while being stable during nighttime. Methods for NPQ
415 correction from fluorometers installed on mobile (e.g. gliders and BCG-Argo floats) and moored
416 platforms (e.g. vertical profilers and buoys) have been established (Fragoso et al., 2021; Lucius
417 et al., 2020; Xing et al., 2018). Each of these methods follows their own set of assumption, given
418 that not all set of parameters are measured from distinct platforms. For semi-autonomous
419 observations (e.g. USVs), interpolation of unaffected nighttime $FChla$ signal has been applied in
420 this and other studies to correct for NPQ (Scott et al. (2020)). In Scott et al. (2020), daytime
421 $FChla$ from the USV Saildrone was corrected using the proportion of $FChla$ to Volume
422 Scattering Function (β , 124°, 650 nm) from the night before and after the NPQ event and
423 assumed that daytime β measurements are not influenced by NPQ . As suggested by the authors,
424 this method is only appropriate under a close examination of the types of water masses. That is
425 because it assumes a consistent temporal and spatial distribution of factors that influences β , such
426 as detritus, sediments and phytoplankton. In our study, this method was not suitable due the
427 noisy turbidity measurements (particularly when wind speed was high), suggesting the influence
428 of air bubbles and particles, such as sediments and detritus in the backscattering signal. Although
429 phytoplankton constitutes a significant portion of particles in productive waters of the coast of
430 mid-Norway, other particles, including zooplankton, fecal pellets, sediments and detritus can be

431 highly heterogeneous in space and time, influencing the backscattering signal (Fragoso et al.,
432 2019; Fragoso et al., 2021). Thus, for $FChla_{AUTO}$ corrections in this study, daily measurements
433 were based the percent reduction in $FChla_{AUTO}$ by modelled irradiance. This method is only
434 appropriate to observe daily trends in $FChla_{AUTO}$, rather than sub-mesoscale patchiness of
435 phytoplankton distributions occurring less than a day.

436

437 4.2 Phytoplankton bloom dynamics

438

439

440 The spring bloom (March-June) in Frohavet consisted of multiple peaks - a short one in
441 late March dominated by the haptophyte *Phaeocystis*, a second long one (2 weeks) during April
442 (where diatom *Skeletonema* spp. is the dominant) and a third ‘on-and-off’ bloom until late June
443 dominated again by *Skeletonema*. Multiple biomass peaks composed of the same taxa (in this
444 case *Skeletonema*) or assemblages of taxa are likely a response of the dynamic nature of this
445 region. In these places, distinct stochastic drivers (nutrient pulse, a period of calm weather,
446 heatwaves, grazing selection etc.) determine the timing of peaks over a short period of time
447 (Type 4 blooms as described in Isles and Pomati, 2021). This means that for each of these
448 blooms observed in during spring to summer in Frohavet, distinct abiotic (changes in nutrient
449 and light conditions) and biotic factors (grazing pressure) might have shaped the intensity,
450 composition and duration of those blooms.

451 The USV was in the water in April during the second long $FChla_{BUOY}$ peak (2 weeks) and
452 captured the temporal and spatial dynamics of *Skeletonema* bloom before, during and after its
453 recession. During this period, a rise in $FChla_{AUTO}$ from the USV AutoNaut occurred
454 concomitantly with a rise in SST and [CDOM] as well as with a decrease in salinity values and

455 wind speed, suggesting that sunny skies, calm winds and warmer temperatures promoted snow
456 melt and high freshwater input from river run off along the coast. Thus, light was the
457 environmental driver that likely promoted this bloom, since several days of clear skies and
458 relatively calm winds shoaled the mixing layer after intense wind mixing and prolonged period
459 of low light conditions of Norwegian winter. In the North Atlantic, similar conditions were
460 observed where a dramatic increase in cellular division rates (net population growth rate from
461 0.02 to 0.08 divisions d⁻¹) over a short period of time (9 days) occurs as a consequence of the
462 rapid shoaling of the mixed layer during calm weather periods, consistent with Sverdrup's
463 paradigm (Mignot et al., 2018). In Frohavet, *FChlaBUOY* started to accumulate since mid-March,
464 but it was not until mid-April that a proper bloom was observed, where concentrations changed
465 from 1 to 5 mg m⁻³ in 5 days.

466 While the spring bloom in this study developed exponentially with 5 days, the collapse of
467 the bloom was rather quick, 1-2 days. *Skeletonema* sp. is an ubiquitous fast-growing diatom
468 (Lundsør et al., 2022), and it is possible that nutrient limitation after intense growth could have
469 caused of the collapse of the bloom. However, due to the intermittent changes and dynamic
470 nature of weather conditions of the coast of mid-Norway, particularly from winter to spring
471 transition, it is likely that strong wind speed and deepening of the mixing layer (after 2 weeks of
472 calm weather) paused the bloom development. This could initially dilute the accumulated
473 biomass and consequently reduced the availability of light for the phytoplankton growth. Similar
474 conditions were observed in the Northwestern Mediterranean Sea, where subsequent storms and
475 vertical mixing due to the intermittent changing weather conditions from winter to spring cause
476 the collapse of spring blooms (Keerthi et al., 2021). In this study, the evidence of a storm surge
477 shown by the sudden increase in average wind speed and increased nutrient availability at the

478 surface in early May indicates that vertical mixing likely promoted the rapid decline of the
479 bloom.

480 Zooplankton abundance and top-down pressure is evident in the coast of mid-Norway,
481 particularly during summer (Fragoso et al., 2022). Although light might have induced
482 phytoplankton cellular division to optimum levels, slight lags (few days) in zooplankton grazing
483 might have been another reason why phytoplankton accumulation reached a peak (3-4 days),
484 where grazing rates outbalanced phytoplankton division and growth. This suggests that loss
485 processes, such as grazing, can also be rather fast, highlighting the need of highly temporal and
486 spatial-resolved measurements in dynamic regions. The copepod *Calanus finmarchicus* is the
487 dominant mesozooplankton species in the Norwegian Sea and abundance has strongly been
488 correlated with chlorophyll *a* concentration, suggesting tight coupling in other regions of the
489 coast of Norway (Dong et al., 2022). Microzooplankton, such as ciliates and heterotrophic
490 dinoflagellates, have a short generational time and high rates of predation, where slight lags in
491 the response of these predators might have allowed for the positive accumulation rates during the
492 initial state of the bloom (Mojica et al., 2021). Thus, it is likely that grazing might have slowed
493 down further accumulation of phytoplankton biomass ([Chl a]) and bloom development,
494 particularly in shallow mixed layers, which favors predator-prey encounter rates. Although the
495 short-scale (few days) mismatch between phytoplankton and zooplankton abundances might
496 have some influence in the development and collapse of the bloom, it is still likely that increase
497 in average wind speed was the main obvious cause, given the fast (1-2 days) decline in
498 chlorophyll concentrations from 5 to 1 mg m⁻³.

499

500 4.3 Submesoscale patchiness

501

502

503 High frequency-resolved mobile platforms such as the USV AutoNaut is a great tool to
504 examine the sub-mesoscale variability of water masses, revealing the patchiness of the system
505 (Dallolio et al., 2021). The wide range in salinity (31.5 - 34°C), temperature (6 - 9°C) and
506 CDOM (1 - 2.5 ppb) reveals that the USV AutoNaut travelled through many distinct water
507 masses, some with more or less influence of riverine input from Trondheimsfjord. Satellite
508 images revealed that phytoplankton spatial distribution in Frohavet is highly patchy over the
509 bloom period, but temporal changes, such as the start ~ April 7th, peak on April 21st, suppress on
510 April 24th and collapse on April 26th were more prominent. The phytoplankton bloom was
511 patchy, particularly on April 24th but did not appear to be confined to a certain water mass, rather
512 transitioning in time and associated to wind conditions (low wind speed - accumulation and high
513 wind speed – disruption of the bloom). This suggests that in spite of some degree of spatial
514 variability observed in environmental factors (salinity and temperature, for example), the
515 formation and decline of the bloom (from <1 to 6 mg m⁻³) was rather temporal due the fast
516 changes in the environment (increase of wind strength).

517

518

519 **5 Conclusions**

520

521 Here we showed the detailed dynamics of a phytoplankton spring bloom in a complex productive
522 region of the coast of Norway using an integrative observational approach. We observed a rapid

523 (5 days) bloom development (from 1 to 5 mg m⁻³) dominated by the diatom *Skeletonema*
524 occurring in spells of ‘good weather’, meaning few days of sunny, clear skies and weak winds in
525 the middle of ‘stormy spring’, typical of high latitudinal regions. The collapse of this bloom was
526 even faster, occurring in 1-2 days, concomitant with increase in wind speed, suggesting the
527 strong influence of environmental conditions in the spring bloom.

528 Integrative approaches using of multiple ocean observation platforms (referred as the
529 observational pyramid in Williamson et al., 2023) is essential to capture the short-term changes
530 of phytoplankton in space and time. As climate continues to change, fluctuations in the
531 environmental conditions (e.g. storm, floods, heatwaves) will likely become more and more
532 frequent, and thus, the response of plankton communities, will likely become more extreme.
533 Therefore, the combination of sensor-based technology and traditional methods for validation
534 and monitoring of the ocean is fundamental to understand of the interlink and tipping points of
535 phytoplankton dynamics to multiple environmental stressors related to climate change.

536

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538

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547

548 **Open Research**

549

550 Our observational data of the parameters measured with the AutoNaut in Frohavet are available
551 at 10.5281/zenodo.8283108.

552

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554

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