

1 **Reduced air–sea CO₂ exchange in the Atlantic Ocean due to biological surfactants**

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22 **Abstract**

23 Ocean CO₂ uptake accounts for 20–40% of the post-industrial sink for anthropogenic CO₂.
24 The uptake rate is the product of the CO₂ interfacial concentration gradient and its transfer
25 velocity, which is controlled by spatial and temporal variability in near surface turbulence.
26 This variability complicates CO₂ flux estimates and in large part reflects variable sea surface
27 microlayer enrichments in biologically derived surfactants that cause turbulence suppression.
28 Here we present a direct estimate of this surfactant effect on CO₂ exchange at the ocean basin
29 scale, with derived relationships between its transfer velocity determined experimentally and
30 total surfactant activity for Atlantic Ocean surface seawaters. We found up to 32% reduction
31 in CO₂ exchange relative to surfactant-free water. Applying a relationship between sea
32 surface temperature and total surfactant activity to our results gives monthly estimates of
33 spatially-resolved “surfactant suppression” of CO₂ exchange. Large areas of reduced CO₂
34 uptake resulted, notably around 20 °N, and the magnitude of the Atlantic Ocean CO₂ sink for
35 2014 was decreased by 9%. This direct quantification of the surfactant effect on CO₂ uptake
36 at the ocean basin scale offers a framework for further refining estimates of air-sea gas
37 exchange up to the global scale.

38 **Surfactants and gas exchange between the atmosphere and ocean**

39 Environmental control of the gas transfer velocity (k_w) of a sparingly soluble gas like CO₂ is
40 exerted through the modification of turbulent diffusion at the air-sea interface¹. Wind speed
41 is a fundamental control of near surface turbulence but relationships between k_w estimated in
42 situ using volatile tracers and corresponding wind speeds show considerable scatter, of which
43 only about 50% is attributable to procedural and/or measurement errors². Contrasting k_w -
44 wind speed parameterisations have consequently been derived for different ocean regions^{3,4},
45^{5,6}, the differences between them resulting from additional and variable k_w control by several
46 other turbulence related variables. These include atmospheric stability, wind fetch, sea state,
47 breaking waves, white caps, bubble transport, rain and the presence of surface active organics
48 (surfactants)¹.

49

50 Biologically-derived surfactants^{7,8} are ubiquitous in the open ocean sea surface microlayer
51 (SML)^{9,10}, where in addition to limiting the rate of air-sea gas exchange¹¹, they contribute to
52 the formation of marine boundary layer (MBL) aerosols involved in atmospheric chemistry
53 and climate regulation¹². The SML is <400 μm deep but is physically and biogeochemically
54 distinct from both the underlying water and the atmosphere¹³. SML enrichments in
55 surfactants, other dissolved components¹⁴ and buoyant particles predominantly occur via
56 bubble scavenging from subsurface water (SSW)¹³. Surfactants can be soluble or insoluble,
57 the former being most important to air-sea gas exchange^{7,8,15} by suppressing k_w through
58 modified surface hydrodynamics and subsequent capillary-gravity wave damping^{16,17}. By
59 contrast, insoluble surfactants that form visible “monolayer” slicks ~1-10 nm thick in calm
60 seas, break down and disperse under increasing turbulence¹⁸. Soluble surfactants
61 accumulating in the SML include transparent exopolymer particles (TEP)¹⁹, polysaccharides
62²⁰, lipid-like material e.g.²¹, amino acids²² and chromophoric dissolved organic matter

63 (CDOM)²³. Where concentrations of total soluble surfactants are high, parameterising k_w
64 solely in terms of bulk turbulence may involve significant errors^{9,24}.
65
66 Surfactant suppression of k_w by up to 50% may be typical but such estimates are mostly
67 based on artificial compounds used in laboratory experiments^{7,15} and/or deployed at sea^{11,18}.
68 In coastal regions, strong spatio-temporal gradients in k_w have been linked to natural
69 surfactants, both in the SML and in the underlying near-surface seawater^{8,24,25}. We recently
70 found persistent total surfactant activity (SA) enrichments in the Atlantic Ocean SML around
71 40°N and 10°N⁹. Air-sea gas exchange rates estimated for these regions without accounting
72 for such enrichments will almost certainly be too high. Previous estimates of the surfactant
73 effect utilised chlorophyll as a surfactant proxy¹⁷. We develop this further herein, providing
74 the first direct and systematic evaluation of surfactant suppression of air-sea gas exchange at
75 the ocean basin scale (Atlantic Ocean), based on observed changes in measured SA and in k_w
76 estimated experimentally with our unique laboratory gas exchange tank²⁶. Suppression
77 estimates were derived for individual biogeographical ocean provinces (“Longhurst
78 Provinces”) defined by phytoplankton distributions as regulated by hydrography²⁷. Our gas
79 exchange tank is a proven and effective tool for elucidating the relationships between k_w , SA
80 and biogeochemical indices of primary productivity²⁶. The data we present here were
81 obtained along Atlantic Meridional transect (AMT) #24 from 50°N to 50°S during
82 September-November 2014 (Figure 1).

83

84 **Surfactants and gas exchange in the Atlantic Ocean**

85 Our data synthesis sets the Atlantic Ocean contribution to the 2014 global marine CO₂ sink at
86 ~25% (1433 Tg C). Our gas exchange tank experimental results show that in comparison to
87 surfactant-free water used as a control, k_w was suppressed in Atlantic Ocean waters by 2 - 32

88 % (Table S1), which is generally greater than a previous estimate of 5% suppression for the
89 open ocean²⁸. Using sea surface temperature (SST) and salinity as a conservative tracer of
90 Atlantic Ocean water bodies (Figure 1) we found a consistently smaller degree of k_w
91 suppression in the South Atlantic (3-18%) than in the North Atlantic (2-24%), and highest k_w
92 suppressions in the Western Tropical Atlantic (WTRA, 6°S–11°N). We related these results
93 (Table S1 and Figure S1) directly to our previously published measurements of SA in the
94 SML (SA range for all sites used in this study = 0.15 to 0.67 mg L⁻¹ T-X-100, n = 13; Table
95 S1)⁹ and derived a linear relationship (% suppression = 32.44 x SA_{SML} + 2.51, $r^2 = 0.51$, p =
96 0.009, n = 13) that is similar to what we previously found for the coastal North Sea²⁵. The
97 range in SML SA relevant to our gas exchange tank experiments is comparable to the ranges
98 we reported previously⁹ for individual Longhurst Provinces, consistent with our derived
99 relationship for the Atlantic Ocean being representative of the wider SA distribution we
100 observed (Table S2). However, the SA variability that we found within each individual
101 Longhurst Province implies a corresponding variability in province-wide values of k_w , which
102 prompts a need for further gas exchange experiments of this type to reduce the attendant
103 uncertainties. Additionally, we note that the SA vs k_w relations for our North Sea²⁵ and
104 Atlantic Ocean datasets have contrasting slopes and intercepts. These imply that in addition
105 to differences in the total concentrations of SML surfactants (SA) in these two areas,
106 variability in the chemical compositions of their total surfactant pools will likely impact k_w
107 variability²⁹. This brings into question approaches that employ proxies such as chlorophyll-a
108 to quantify the suppression of k_w by surfactants¹⁷, and this is supported by our recent North
109 Sea²⁵ and Atlantic Ocean⁹ datasets that show no clear relationships between SA in the SML
110 and either total chlorophyll, total pigments or size-fractionated primary production.

111

112 To further interrogate the effect of surfactants on Atlantic Ocean CO₂ exchange, we adjusted
113 wind-speed derived CO₂ fluxes with an empirically-derived, non-linear % Suppression-SST
114 relationship ($r^2 = 0.61$, $n = 13$; see methods, Figure S2, and Table S3). Our estimated mean k_w
115 suppression due to surfactants is spatially and temporally variable (Figure S3) and the wind
116 speed relation of Nightingale et al. ⁴ that we used to calculate k_w derives from spatially and
117 temporally heterogeneous coastal seas for which no corresponding SA measurements are
118 available. Consequently, the influence of surfactants on its wind speed-derived estimates of
119 k_w are unknown. We note that all available k_w parameterisations ^{3,4,5} are likely derived from
120 data that are influenced by variable SA. We evaluated the percent k_w suppressions by
121 surfactant for each of the three coastal locations included in the Nightingale et al. ⁴ synthesis
122 (Southern North Sea, Georges Bank, Florida Shelf) by applying our above % suppression-
123 SST relationship to the published temperature ranges for these experiments (5.3-16.9 °C) ^{4,30,}
124 ³¹. The resulting range in k_w suppression was 0.3-6.5 %. We also note that the majority of the
125 data used in the Nightingale et al. ⁴ analysis (those from the Southern North Sea and Georges
126 Bank) are from locations bordering regions of very low surfactant suppression (Figure S3).
127 Based on this we applied the central limit theorem (see methods) and so assume that the data
128 used to develop the Nightingale et al. ⁴ parameterisation comprise of an equal number of
129 surfactant-influenced and surfactant-free data points and that the upper range of this
130 surfactant influence is likely low (i.e. $\leq 6.5\%$ suppression). Following this reasoning, the
131 Nightingale et al. ⁴ relation thus identifies the central estimate of the k_w -wind speed relation,
132 which tends towards surfactant-free conditions. We therefore contend that our use of the
133 Nightingale et al. ⁴ relation as a reference is entirely reasonable given that this study is the
134 first systematic attempt to evaluate the surfactant effect at the ocean basin scale. We selected
135 SST based on its use in satellite algorithms, in conjunction with photosynthetically active
136 radiation and chlorophyll-a, for estimating primary production integrated over time and

137 space. This integration has the advantage of smoothing out any process-related time lags that
138 are manifested in the phytoplankton photosynthetic response and subsequent surfactant
139 production ^{29,32}. Such time lags are consistent with our inability to observe any clear
140 relationship between SA and chlorophyll-a, and likely account for at least some of the
141 otherwise unexplained variance in our data. Applying this relationship to our experimental
142 and observational results yields a basin-wide, spatio-temporal estimate of the impact of
143 surfactants (see Figure S3). The resulting range in k_w suppression for individual grid-squares
144 ($1^\circ \times 1^\circ$; see methods) was 2 to 24 % and the corresponding net effect on the 2014 Atlantic
145 Ocean CO₂ sink was to reduce it by 9 % (131 Tg C), from 1433 Tg C (derived using wind
146 parameterisation based estimates alone⁴) to 1302 Tg C (Table S3 expressed as R_{660} ' values
147 and Table S4), with monthly values of % suppression ranging from 2.6 to 11.2 %, peaking in
148 November (Table S3). Recent work in the Atlantic Ocean between 40°N and 60°N
149 highlighted the importance of bubble-mediated CO₂ exchange during wave breaking at high
150 wind speeds ^{33,34}. In our earlier work we found surfactants to be ubiquitous in the Atlantic
151 Ocean SML up to wind speeds of at least 13 m s⁻¹ ⁹, beyond the typical threshold for wave
152 breaking ³³, and that the SML surfactant enrichments were maximal between 40°N and the
153 northern boundary of our study at 50°N ⁹. A co-existence of bubbles and surfactant in these
154 high wind regions will act to impact air-sea gas exchange in two ways. First, we contend that
155 irrespective of the important solubility-driven enhancement of k_w by bubbles ³⁵, a separate
156 “surfactant effect” at these wind speeds will to some extent attenuate the increase in gas
157 exchange rates that would otherwise be exerted through this bubble effect alone. Second, by
158 coating the surfaces of rising bubbles surfactants act to reduce their rise speeds and thereby
159 force an additional inhibition of air-sea gas exchange ³⁵. As we have shown previously ²⁵, the
160 influence of surfactants on k_w , whether in the presence of bubbles or not, is spatially and
161 temporally variable, due to corresponding variability in both SA and near-surface turbulence

162 (e.g. as driven by wind speed), which will also impact bubble production. Consequently, in
163 regions of high and variable winds uncertainty over the fractional contribution of surfactants
164 to k_w will be maximal, being compounded by the presence of bubbles, and both spatially and
165 temporally variant.

166

167 We hypothesise that the increases in % suppression we observed for both the North and
168 South Atlantic are in large part likely driven by total sunlight hours that influence primary
169 productivity at the sub-basin scale. If so, the observed temporal variability may be associated
170 with seasonality and/or the movements of air such as the trade winds, and atmospheric
171 moisture and cloud formation in the inter-tropical convergence zone (ITCZ). Partitioning of
172 the total net air-sea CO₂ flux between the North and South Atlantic in our analysis was
173 similar with and without surfactant suppression (54 % vs 46 %). At the Longhurst Province
174 scale there were notable ‘hotspots’ of CO₂ flux suppression that were sustained throughout
175 the year (Figure 2 and Table S4). The largest reduction was found for the North Atlantic
176 Tropical Gyral (NATR) province, where purely wind-based CO₂ exchange may be
177 overestimated by up to 4.6 Tg C per month, whereas the smallest impact was found for the
178 North Atlantic Subtropical Gyre - West (NAST(W)) and Eastern Tropical Atlantic (ETRA)
179 provinces (0.03 Tg C per month). The most prevalent surfactant effects were in WTRA and
180 NATR and occurred throughout the year, with other hotspots in the North Atlantic
181 Subtropical Gyral – East (NAST(E)) and NAST(W) provinces from October to June (Figure
182 2). These hotspots imply that at these times the Atlantic Ocean contribution to the global CO₂
183 sink is reduced by 10-24%. However, when NAST(E) (August-September), NAST(W) (July-
184 September), and ETRA (February-June) all became net CO₂ sources the ‘surfactant effect’
185 also reduced CO₂ emissions. Further minor reductions in CO₂ emissions were notable off the

186 West African coast in June, south of the equatorial Atlantic from January to May and along
187 30°N from July to October (Figure 2).

188

189 **Global implications of the surfactant suppression effect**

190 Our analysis shows that SML surfactants impact the air-sea exchange of CO₂ at the oceanic
191 scale. Consequently, spatio-temporal variability in SA and hence in k_w suppression likely
192 accounts for a substantial fraction of the observed uncertainty in basin-wide CO₂ air-sea
193 fluxes. More tightly constraining this suppression and its variability in space and time should
194 therefore aid in reducing the uncertainty inherent in constraining net oceanic CO₂ uptake. The
195 surfactant suppression effect is particularly important given our recent work that suggests a
196 persistence of surfactants in the SML beyond wind speeds at which they were previously
197 thought to disperse⁹. The occurrence of “hotspots” of CO₂ exchange in the Atlantic Ocean
198 during warmer periods, as revealed by our analysis, highlights a need to more effectively
199 constrain the temporal and spatial scales of k_w suppression by surfactant and the distributions
200 of SA and related physical and biogeochemical variables, not only in the Atlantic but in all
201 major ocean basins. Such studies will substantially improve the confidence of CO₂ uptake
202 estimates that we necessarily up scaled from limited data and should lead to the development
203 of more effective proxies for estimating the spatio-temporal variability of surfactant control
204 on air-sea gas exchange.

205

206 Specifically, the use of Atlantic Ocean SST as a spatial and temporal integrator of the
207 variability in SA requires additional verification via further measurements in all major ocean
208 basins, where we currently lack any clear supporting evidence of a link between SA and SST
209 or between SA and other potential SA proxies. If SST indeed proves to be a universally
210 robust predictor of SML surfactants resulting from primary production, this would be an

211 important step towards refining the “surfactant suppression effect”. This important positive
212 feedback mechanism to ocean carbon uptake should be taken account of in future evaluations
213 of the global carbon cycle and the attendant implications for future global climate.

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334 **Contributions**

335 R.P. performed the gas exchange experiments. B.S. provided the surfactant measurements.

336 I.A. and J.D.S developed the FluxEngine analysis and ran the model. R.P. and R.U.G.

337 conceived the study. All authors discussed the results and developed the project and

338 manuscript.

339

340 **Competing financial interests**

341 The authors declare no competing financial interests

342

343 **Figures and Figure Captions**

344 Figure 1: Left: Atlantic Meridional Transect 24 (AMT24) from the UK to the Falkland

345 Islands, 2014. The cruise transect (blue line) crosses the following Longhurst

346 Biogeographical Provinces: North Atlantic Drift (NADR); North Atlantic Subtropical

347 Gyral—East (NAST(E)); North Atlantic Subtropical Gyral—West (NAST(W)); North

348 Atlantic Tropical Gyral (NATR); Western Tropical Atlantic (WTRA); Eastern Tropical

349 Atlantic (ETRA); South Atlantic Gyral (SATL); and the South Subtropical Convergence

350 (SSTC). AMT24 start and end shown as blue diamonds. Sample locations shown in black

351 circles. Right: Scatterplot of % suppression of k_w as a distribution of Atlantic Ocean

352 temperature and salinity in 2014.

353

354 Figure 2: Monthly maps of the estimated difference in air-water CO₂ flux caused by

355 surfactants. FluxEngine outputs presented as suppressed flux – original flux, with lighter

356 colours showing decreased CO₂ emission and darker colours showing decreased CO₂ uptake.

357 **Methods**

358

359 **Sample Collection**

360 Samples were collected and gas exchange experiments executed during Atlantic Meridional
361 Transect Cruise (AMT) 24 on board R.R.S *James Clark Ross* (*JCR*; Figure 1). The cruise
362 track ran southbound from Immingham, UK to Port Stanley, Falkland Islands, between 22
363 September and 6 November 2014. The cruise crossed seven biogeographical provinces
364 defined by Longhurst ²⁷: North Atlantic Drift (NADR, 44–58°N); North Atlantic Subtropical
365 Gyral—East (NAST(E)), 26–44°N); North Atlantic Tropical Gyral (NATR, 11–26°N);
366 Western Tropical Atlantic (WTRA, 6°S–11°N); South Atlantic Gyral (SATL, 42–6°S); and
367 the South Subtropical Convergence (SSTC, 45–42°S).

368

369 Sea surface microlayer (SML) and sub-surface water (SSW) samples were collected during
370 midday CTD casts following a well-established protocol ³⁶. To minimise contamination from
371 *JCR* all samples were collected following best practices detailed in Sabbaghzadeh, et al. ⁹.
372 Briefly, the SML was sampled using a Garrett Screen ³⁷ (mesh 16, wire diameter 0.36 mm,
373 opening 1.25 mm), with an effective surface area of 2025 cm², transferred to 50 mL high-
374 density polyethylene (HDPE) bottles and stored in an on-board refrigerator at 4 °C ^{13,25}. SSW
375 was collected from the ship's underway non-toxic seawater supply (located approximately 3-
376 5 m below the water surface) after flushing of the line with copious amounts of sample
377 seawater. 93 litres of this seawater were directly loaded into our gas exchange tank for gas
378 exchange experiments that commenced immediately and were completed within 1.5 hours of
379 sample collection. 50 mL sample aliquots were also collected in HDPE bottles and stored in
380 an on-board refrigerator at 4 °C for subsequent SA analysis that was completed within 24
381 hours. We used SSW in the tank experiments following the procedure outlined by Pereira, et

382 al.²⁵ as there is no practical procedure for collecting a large volume sample of surface
383 seawater that preserves the integrity of the SML. However, we have shown (i) that following
384 its disturbance by vigorous mixing in a laboratory tank the SML becomes re-established on a
385 time scale of seconds with respect to surfactants and other SML components¹³; (ii) that a
386 new SML is similarly established when sub-surface coastal waters are pumped into large
387 mesocosm tanks³⁸.

388

389 **Sample Analysis**

390 Surfactant activity (SA) in the SML and in SSW was measured on board *JCR* by phase-
391 sensitive hanging mercury drop AC voltammetry³⁹ (797 VA Computrace: Metrohm,
392 Switzerland), within 24 hours as previously reported by Sabbaghzadeh, et al.⁹. In brief, the
393 polarograph was situated on a gimbal table to minimise any vibration due to ship movement
394 that might otherwise affect the integrity of the mercury drop. All samples were analysed in
395 triplicate, with their salinities pre-adjusted to 35.0 via the addition of surfactant-free 3 mol
396 L⁻¹ NaCl solution or by dilution with Milli-Q 18.2 Ohm deionised water (Millipore System
397 Inc., USA). Calibration was against the non-ionic soluble surfactant Triton T-X-100. The SA
398 of Milli-Q water was continually analysed throughout the cruise and was always found to be
399 below the method detection limit. All equipment was acid-washed (10% HCl) and rinsed in
400 Milli-Q 18.2 Ohm deionised water prior to use.

401

402 The SSW samples were used to estimate the variability in k_w using a fully automated, closed
403 air–water gas exchange tank, the design, operation and routine rigorous cleaning of which are
404 described in detail elsewhere^{25,26}. Briefly, the system generates controllable and
405 reproducible water-side turbulence with an electronically operated baffle whilst measuring
406 the partial pressures of artificially enriched gaseous tracers. To achieve this the gas exchange

407 tank is coupled to two gas chromatographs (GC's) in a continuous gas-tight system, which
408 allows temporal changes in the partial pressures of the gaseous tracers to be measured
409 simultaneously in the tank water (via an integral equilibrator) and headspace, thereby
410 facilitating independent estimates of k_w for each turbulence setting applied. Due to the
411 dependence of k_w on the Schmidt Number (Sc : the ratio of kinematic viscosity of water to gas
412 diffusivity) raised to the power n , the k_w estimates were converted to k_{660} , the value of k_w for
413 $Sc = 660$ (the value for CO_2 in seawater at 20 °C), assuming $n = 0.5$ for a wavy surface¹. The
414 setup of the gas exchange tank system on board *JCR* differed slightly from our previous
415 studies²⁵ in that only CH_4 and the lowest water-side baffle setting of 0.6 Hz were selected.
416 Using only one tracer gas allowed for simpler setup in a relatively confined space and our
417 previous work demonstrated significantly better analytical precision for CH_4 than for SF_6 ²⁵.
418 With a baffle setting of 0.6 Hz, bubble generation in our gas exchange tank is precluded²⁵. A
419 critical consideration of our experimental setup was to account for the movement of the ship,
420 which had the potential to create turbulence additional to that generated in the experiments.
421 We therefore ran a surfactant-free Milli-Q water sample (herein referred to as a sample
422 blank) immediately prior to each seawater sample. The sample blank k_{660} ($cm\ hr^{-1}$) was
423 normalised to the k_{660} of an 'installation blank', a surfactant-free Milli-Q water sample that
424 was run whilst the ship was stationary in port prior to the cruise. SA analysis of the Milli-Q
425 water ensured that the gas exchange tank was surfactant-free prior to the experiment. The
426 resulting correction factor was then applied to k_{660} derived for each sample (i.e. $k_{660\ Sample}' =$
427 $k_{660\ Sample} \times (k_{660\ Installation\ Milli-Q} / k_{660\ Sample\ Milli-Q})$). To ascertain the comparative SA effect on k_w
428 we normalised our seawater k_{660}' values for each site to the k_{660} value of the installation blank
429 (i.e. $R_{660}' = k_{660\ Sample}' / k_{660\ installation\ blank}$)²⁶. The uncertainty in each k_{660} measurement was
430 derived via Gaussian error propagation⁴⁰ and was always less than $\pm 0.6\ cm\ hr^{-1}$ ($n = 13$). To
431 gauge the possibility of artefacts arising from biological production or consumption of CH_4 ,

432 from losses due to leakage or arising from GC analytical drift, the total masses of CH₄ were
433 continually estimated over the duration of the experiments, from their measured partial
434 pressures and the known water-phase and air-phase volumes (mass balance; Schneider-Zapp,
435 et al. ²⁶, Eq. 14). All experimental data with a total mass balance error of more than ±5 %
436 were excluded ²⁵.

437

438 Supporting biogeochemical and meteorological data were obtained from the AMT data pool,
439 accessible via the British Oceanographic Data Centre (BODC) (<http://www.bodc.ac.uk>).

440 Physical parameters including *in-situ* salinity and temperature were logged from the ship's
441 underway non-toxic seawater systems and calibrated against discrete samples using benchtop
442 instruments. Wind speed and direction were measured using an on-board meteorological
443 package and post processed following BODC guidelines. Chlorophyll a concentrations were
444 determined on samples collected at 2 m depth from the midday CTD cast, by acetone
445 extraction and fluorometric detection according to Welschmeyer ⁴¹.

446

447 **Ocean - Atmosphere Fluxes**

448 Ocean - atmosphere exchange of CO₂ was calculated using the FluxEngine toolbox, an
449 established set of open-source Python tools that enable user defined inputs and gas flux
450 parameterisations in the estimation of regional and global fluxes ^{42, 43, 44}.

451

452 In this work, ocean - atmosphere CO₂ fluxes were calculated following equation 2 of Shutler,
453 et al. ⁴³ and were consistent with the rapid model methods and temperature handling of
454 Woolf, et al. ⁴⁵. The wind speed relation of Nightingale, et al. ⁴ implemented within
455 FluxEngine was used to calculate k_w , such that $k_w = (0.222(U_{10})^2 + 0.333U_{10}) (Sc/600)^{-1/2}$,
456 where Sc is the Schmidt number of CO₂. Net integrated air-sea CO₂ fluxes were calculated by

457 FluxEngine with adjustments for sea ice (using the ice normalisation of Takahashi, et al. ⁴⁶),
458 varying pixel areas (assuming Earth is an ellipsoid) and sub grid cell variations of land and
459 sea as described in Appendix A of Shutler, et al. ⁴³.

460

461 All gas flux calculations used CO₂ fugacity (fCO₂) as inputs and calculated the CO₂
462 concentration at the top and bottom of the mass boundary layer. Earth observation sea surface
463 temperature (SST_{find}) data from the Reynolds data set ⁴⁷ are calibrated to depth and so can be
464 used to represent the temperature at the bottom of the mass boundary layer, allowing the CO₂
465 concentration at this point to be calculated. A constant cool-skin difference of -0.17°C ⁴⁸ was
466 applied to the SST_{find} to represent the skin (interface) temperature SST_{skin}, allowing the CO₂
467 concentration at the top of the mass boundary layer to be calculated.

468

469 Surface-water fCO₂ (fCO_{2w}) data were obtained from the Surface Ocean CO₂ Atlas
470 (SOCAT) version 4 observations. These were re-analysed to a common satellite-derived sea
471 surface temperature (SST) data set that represents the temperature at the bottom of the mass
472 boundary layer ⁴⁷, using the method of Goddijn-Murphy, et al. ⁴⁹. A global climatology,
473 normalised to the year 2010, was then created from these data ⁴⁹.

474

475 Atmospheric pCO₂ (pCO_{2A}) was calculated by FluxEngine using modelled air pressure (P)
476 from the European Centre for Medium-Range Weather Forecasts (ECMWF) ERA-interim
477 model and the in-situ CO₂ dry air mole fraction (XCO₂) from the National Oceanic and
478 Atmospheric Administration Earth System Research Laboratory (NOAA ESRL) ⁵⁰. pCO_{2A}
479 was converted to fCO_{2A} (µatm) using Weiss ⁵¹: equation 9,

480

481
$$b11 = -1636.75 + (12.04SST_{skin} - 0.033SST_{skin}^2 + 3.17e^{-5}SST_{skin}^3)$$

482
$$d_{12} = 57.7 - 0.12SST_{skin}$$

483
$$fCO_{2A} = \exp \left[1e^{-6}(b_{11} + d_{12}) \left(\frac{P}{R SST_{skin}} \right) \right] pCO_{2A}$$

484

485 Where the gas constant, $R = 82.06, \text{ cm}^3 \text{ atm (mol K)}^{-1}$

486

487 Global wind speed at 10 m (U_{10}), was taken from the European Space Agency (ESA)

488 GlobWave (GlobWave 2015) data set. The salinity dataset was from the World Ocean Atlas

489 salinity climatology provided in Takahashi, et al. ⁴⁶ and global percent sea-ice cover data

490 were from the Centre de Recherche et d' Exploitation Satellitaire (CERSAT) Special Sensor

491 Microwave Imager (SSM/I) dataset. Where they were not originally provided as such, input

492 datasets were re-gridded onto a $1^\circ \times 1^\circ$ grid where each grid value was the statistical mean of

493 all contributing data within that cell.

494

495 **Surfactant suppression effect modelling**

496 The surfactant suppression effect was determined by fitting a non-linear relationship ($R_{660}' =$

497 $0.0046 \times SST_{skin}^{2.5673}$) to our R_{660}' and SA dataset presented in this manuscript. This approach

498 recognises that at lower temperatures, primary production effects may be less influential than

499 some other processes (e.g. sea ice formation) ^{52, 53}. As a result, at lower temperatures, the

500 predicted suppression effect is 0.25% below 5 °C and 0.05% below 3 °C. For this study, we

501 did not apply an upper temperature limit but recognise that further work is needed to

502 constrain the threshold and upper limit at which SST ceases to exert a strong control on

503 primary productivity and hence on SA. In our conceptual approach, we assume that SST_{skin} is

504 the driving factor in primary production and thereby influences the presence of surfactants in

505 the SML. For each Longhurst biogeographical province, we applied our R_{660}' suppression

506 factor to the wind parameterisation of Nightingale et al. ⁴ to estimate the reduction in CO_2

507 flux across the Atlantic Ocean air-sea interface. The wind speed relation of Nightingale et al.,
508 ⁴ used to calculate k_w , derives from spatially and temporally heterogeneous coastal seas for
509 which no corresponding SA measurements are available and for which the influence of
510 surfactants on k_w are thus unknown. We therefore assume that the effect of SA inherent in the
511 Nightingale et al. ⁴ data is variable. Consequently, we applied the central limit theorem.
512 Similar assumptions apply when handling 'Type A' uncertainties, i.e. the analysis of N
513 repeated independent measurements of a quantity, as described by the Joint Committee for
514 Guides in Metrology ⁵⁴. This suggests that the Nightingale et al. ⁴ dataset should contain
515 approximately equal numbers of surfactant-free and surfactant influenced data points. We
516 therefore assume that the Nightingale et al. ⁴ analysis identifies the central estimate of the k_w -
517 wind speed relationship and thus tends towards a condition of a small amount of surfactant
518 influence. Given this, we adopted the Nightingale et al. ⁴ parameterisation as our reference.
519
520 Uncertainties in the non-linear relationship between SST and R_{660} were estimated as the
521 RMSE of the residuals when fitted to the in-situ measured data. For the power relationship,
522 the errors from the residuals, $dR_{660} = d(e^{SST})$ and so, $dR_{660} = e^{SST} dSST$. The observed
523 variability is likely to be caused by sea-surface conditions that persist spatially. As such, a
524 spatial correlation length would be expected in the variability. This will serve to increase the
525 variability in net-flux estimates as a purely random spatial signal would be largely removed
526 in a spatial total. It follows that the upper limit in uncertainties will arise when the error is the
527 same simultaneously across the entire area considered. In this work, perturbations were
528 drawn at random from dR_{660} for 20 ensemble runs and applied to all values in an area
529 simultaneously. The uncertainty values were calculated as the RMSE of the relevant value
530 across all runs and represent the largest expected uncertainties.

531

532 Uncertainty in the in-situ R_{660} / SST relationships could result from the influence of other
533 factors that affect the flux, and which are not accounted for in the above. However, without
534 additional measurements, we chose to use the largest possible uncertainty estimate. Further
535 data and a more detailed understanding of the relationship between R_{660} and other relevant
536 parameters would likely reduce these uncertainties.

537

538 **Data Availability**

539 The authors declare that the data supporting the findings of this study are available within the
540 article and its supplementary information files. FluxEngine outputs are available from the
541 corresponding author upon request. Supporting biogeochemical and meteorological data were
542 obtained from the AMT data pool, accessible via the British Oceanographic Data Centre
543 (<http://www.bodc.ac.uk>). The FluxEngine documentation is available online:
544 <http://www.oceanflux-ghg.org/Products/FluxEngine> or the open-source code can be
545 downloaded from github (<https://github.com/oceanflux-ghg/FluxEngine>). Atmospheric air
546 pressure data were obtained from the European Centre for Medium-Range Weather Forecasts
547 (ECMWF) ERA-interim model ([http://apps.ecmwf.int/datasets/data/interim-full-](http://apps.ecmwf.int/datasets/data/interim-full-daily/levtype=sfc/)
548 [daily/levtype=sfc/](http://apps.ecmwf.int/datasets/data/interim-full-daily/levtype=sfc/)). The CO₂ dry air mole fraction (XCO₂) data were from the National
549 Oceanic and Atmospheric Administration Earth System Research Laboratory
550 (https://www.esrl.noaa.gov/gmd/ccgg/globalview/co2/co2_intro.html). Global wind speed at
551 10m (U₁₀) was obtained from the European Space Agency (ESA) GlobWave (GlobWave
552 2015: <http://globwave.ifremer.fr/>) data set and the global percent sea-ice cover data were
553 from the Centre de Recherche et d' Exploitation Satellitaire (CERSAT) Special Sensor
554 Microwave Imager (SSM/I) dataset available at
555 <http://cersat.ifremer.fr/data/products/catalogue>.

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The effect of biological surfactants on air-sea CO₂ exchange in the Atlantic Ocean

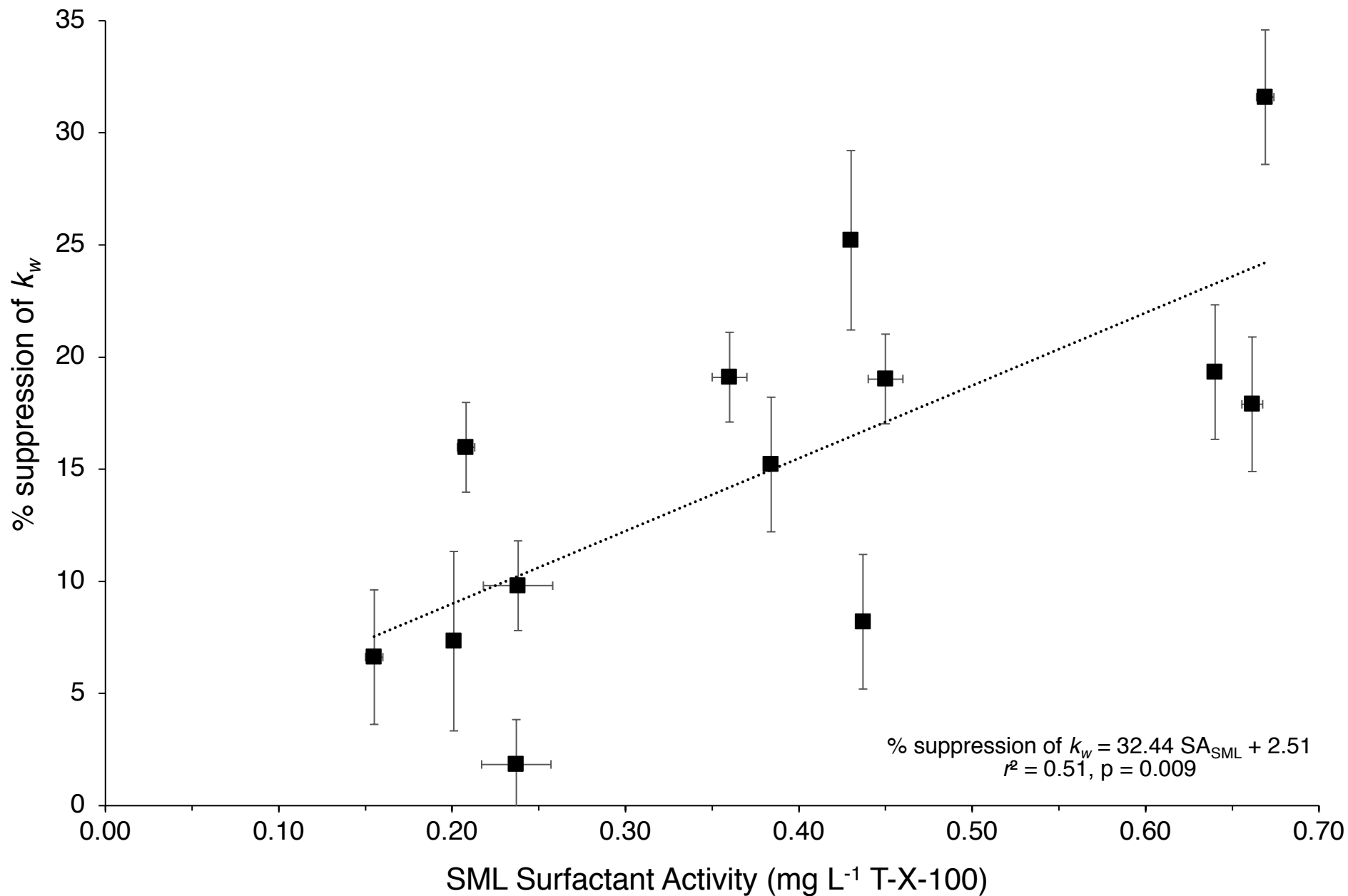


Figure S1: Scatter plot of % suppression of k_w ($R_{660}' = (R_{660}' = k_{660 \text{ Sample}}' / k_{660 \text{ installation blank}})$) and surfactant activity (SA) in the SML for the Atlantic Ocean September to October 2014. Error bars are the standard errors of SA and R_{660}' .

The effect of biological surfactants on air-sea CO₂ exchange in the Atlantic Ocean

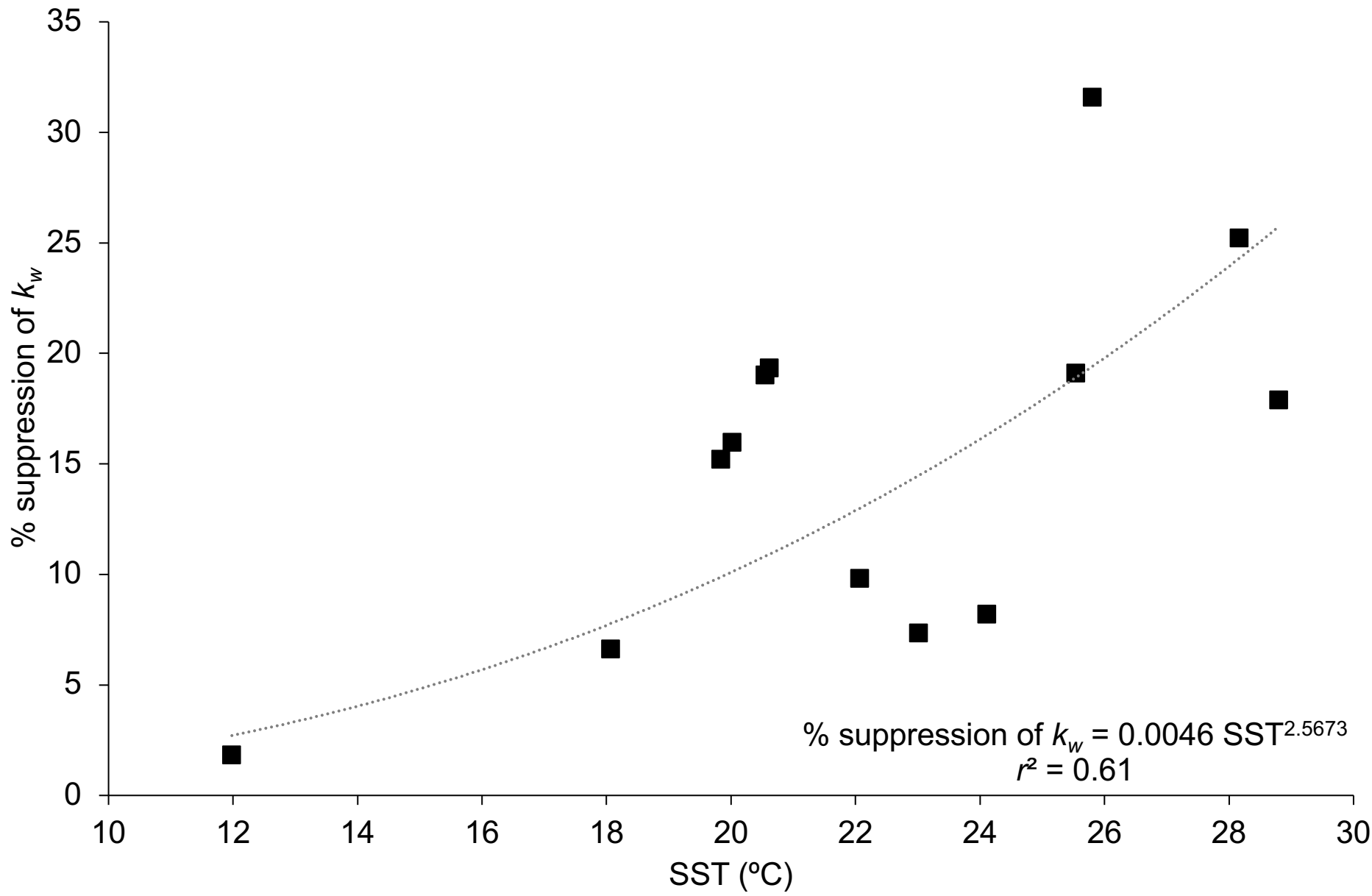


Figure S2: Scatter plot of the SST (°C) and R_{660}' (expressed as % suppression of k_w) non-linear relationship of the Atlantic Ocean from September to October 2014.

The effect of biological surfactants on air-sea CO₂ exchange in the Atlantic Ocean

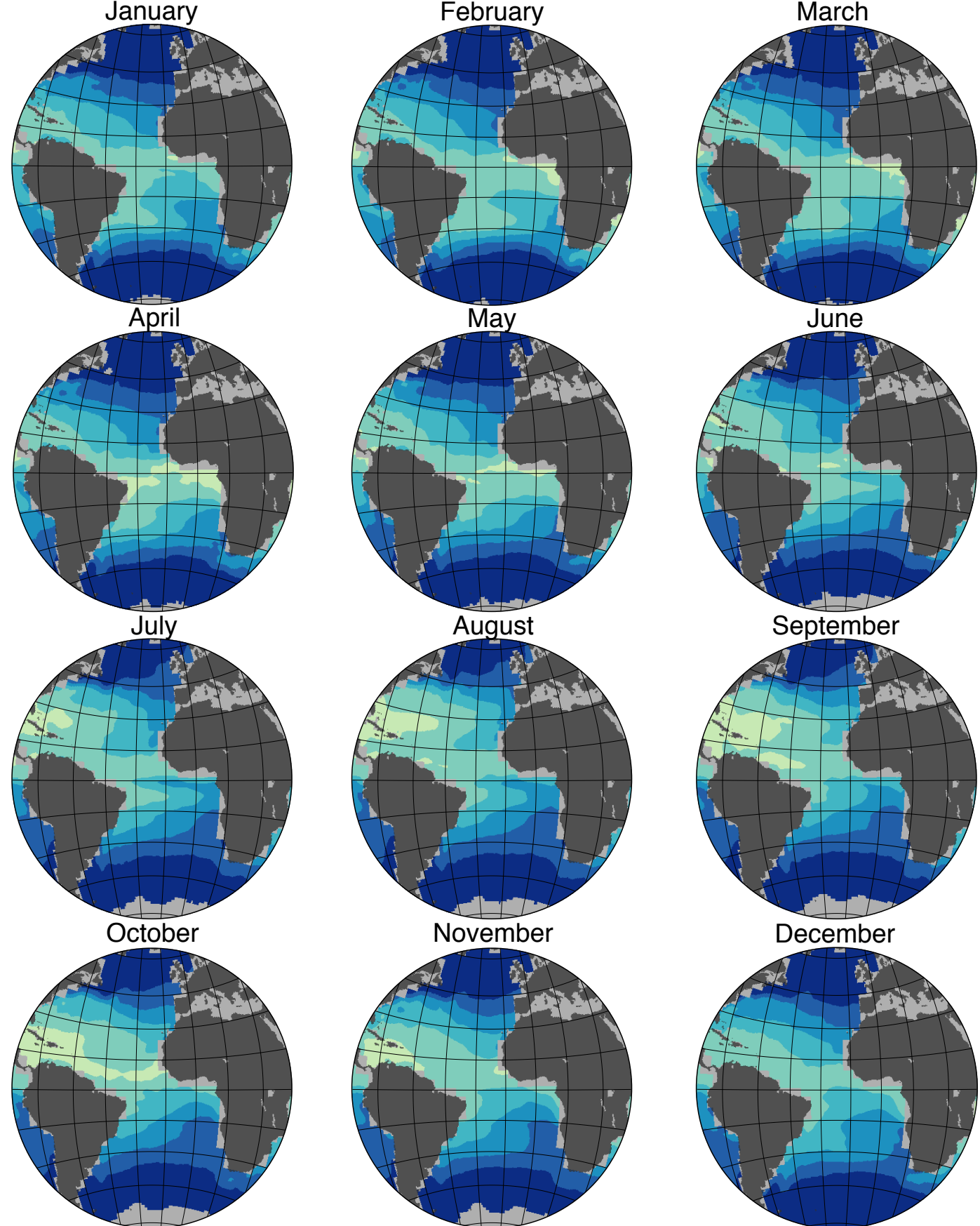


Figure S3: FluxEngine monthly surfactant effect maps of calculated R'_{660} as a function of SST (°C). Light colours show estimates of increased surfactant suppression and darker colours show areas decreased surfactant suppression.