

Seasonal oceanography from Physics to micronekton in the south-west pacific

Christophe E. Menkès, Valérie Allain, Martine Rodier, Francis Gallois, Anne Lebourges-Dhaussy, Brian P.V. Hunt, Houssem Smeti, Marc Pagano, Erwan Josse, A. Daroux, et al.

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Seasonal Oceanography from Physics to Micronekton in the South-West Pacific.

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34	

Abstract

estimates and acoustic-derived estimates compared reasonably well. Estimates of
micronekton from net observations and the SEAPODYM model were in the same range. The
non-dedicated acoustics adequately reproduced trends observed in zooplankton from nets,
but the acoustics could not differentiate between zooplankton and micronekton and
absolute biomasses could not be calculated. Understanding the impact of mesoscale
features on higher trophic levels will require further investigation and patchiness induced by
eddies raises the question of how to best sample highly dynamic areas via sea experiments.

Keywords

.eys, mesost Zooplankton, nekton, acoustic data, oceanographic surveys, mesoscale eddies, oligotrophic,

primary production

1 Introduction

78	In the South Pacific Ocean fishing of apex predators, such as tuna and billfishes, represents a
79	major economic and food resource (Bell et al., 2013). Considerable variability in tuna catch
80	rates is observed in fisheries (Rouyer et al., 2008). Although much of this variability remains
81	unexplained, tuna abundance in space and time has been correlated with factors including
82	oceanographic conditions, physiological constraints (e.g. temperature, depth, oxygen
83	requirements), forage availability, and reproductive behavior (Farley et al., 2013; Senina et
84	al., 2008; Young et al., 2011).
85	Tuna forage predominantly comprises micronekton (Young et al., 2010; this issue).
86	Micronekton are defined as organisms in the 2-20 cm size range and are predominantly
87	distributed in the upper 1000 m of the water column. Micronekton play a key role as
88	intermediaries between plankton production, their prey, and top predators. Since
89	micronekton biomass is dependent on the availability of plankton prey, it is expected that
90	plankton production, its oceanographic drivers, and micronekton biomass would be tightly
91	coupled, and therefore act in concert in determining top predator distributions.
92	The New Caledonian Exclusive Economic Zone (EEZ), a region of more than 1.4 $10^6\mathrm{km^2}$, is
93	located in the Coral Sea, at the southeastern edge of the South Pacific (Figure 1). The
94	dominant feature of circulation across 0-150 m is the westward-flowing South Equatorial
95	Current (SEC) from $^{\sim}25^{\circ}\text{S}$ to the equator. The SEC flow bifurcates at the Australian
96	continental margin (Ridgway and Dunn, 2003) at $^{\sim}15^{\circ}$ S, with one branch connecting with
97	the southward flowing East Australian Current (EAC) (Qu and Lindstrom, 2002) and the
98	other forming the Gulf of Papua Current which flows northward along the coast of
99	Queensland. Within the Coral Sea, the SEC comprises narrow filaments and jets created by
100	the complex island, reef, seamounts and ridge topography (Gourdeau et al., 2008) namely
101	the North Vanuatu Jet at around 13-15°S, and the North Caledonian Jet at around 17-18°S
102	(Couvelard et al., 2008; Marchesiello et al., 2010). To the south of New Caledonia, the
103	surface flow returns from the EAC back into the central south Pacific (Figure 1) as the South
104	Tropical Counter Current (STCC) (Marchesiello et al., 2010). In this region, the structures of
105	the ocean currents are prone to shear instabilities and high eddy kinetic energy is observed

106	(Qiu et al., 2009). Excluding the very coastal areas, the New Caledonian EEZ is regarded as
107	oligotrophic (Dandonneau and Gohin, 1984) with a mean nitracline depth of $^{\sim}$ 110 m (Figure
108	1). South of 22°S, the region experiences higher productivity (Ceccarelli et al., 2013;
109	Dandonneau and Gohin, 1984).
110	Within this oceanographic context, the longline fishery for tuna represents approximately
111	30% of the total fisheries harvest in New Caledonia (Gillett, 2009). Catches are dominated
112	by albacore tuna (<i>Thunnus alalunga</i>) and exhibit two seasonal peaks in July - August and
113	December, and the highest catch rates occur in the north-western part of the EEZ (Briand et
114	al., 2011). The influence of temperature, primary production and micronekton density on
115	tuna catch rates has been demonstrated in New Caledonia (Briand et al., 2011), in American
116	Samoa (Domokos, 2009) and at the ocean basin scale in the Pacific Ocean (Lehodey et al.,
117	1998).
118	Large-scale observations of temperature, surface currents and surface primary production
119	derived from satellite data have allowed validation of the existing oceanographic models,
120	giving confidence in the use of modeled oceanographic parameters for such analyses.
121	However there are few observations of biological parameters, including micronekton, to
122	validate the model biological outputs.
123	At the scale of the South Pacific, nutrient and in situ phytoplankton data are sparse, as are
124	data on zooplankton (Carassou et al., 2010; Le Borgne et al., 2011; McKinnon, 2005; Young
125	et al., 2011). Knowledge of the micronektonic communities and their distributions is
126	somewhat more comprehensive, but is based primarily on top predator diet studies (Allain
127	et al., 2012; Olson et al., 2014; Young et al., 2011, 2010). Few data are available from in situ
128	sampling with nets in the South Pacific (Flynn and Paxton, 2012; McPherson, 1991) and in
129	general, none of the available micronekton data are coupled with information on
130	oceanographic conditions. These constitute important gaps in our knowledge and
131	understanding of the dynamics of the pelagic ecosystem.
132	Prior to this study, in situ data on micronekton in the New Caledonian region were derived
133	from a handful of studies conducted in the eastern part of the EEZ (Grandperrin, 1975,

1969; Legand et al., 1970; Roger, 1986, 1974). Overall, data from the New Caledonia region
are limited in both space and time, prohibiting a comprehensive description of the pelagic
ecosystem, including the main seasonal patterns of zooplankton and micronekton and their
relationships with the oceanography.
In 2011, we conducted two dedicated multi-disciplinary bio-oceanographic cruises
(Nectalis1 and Nectalis2) in an effort to fill some knowledge gaps highlighted above for the
New Caledonian and greater South Pacific region. Oceanography, nutrient and food web
components were sampled in areas of high (north-west) and low (north-east) albacore tuna
catch rates in the New Caledonia EEZ, during the austral cool and hot seasons when
oceanography is contrasted and tuna catches high. The primary aim of these cruises was to
provide insights into how phytoplankton, zooplankton and micronekton are coupled with
ocean dynamics in the upper water column (0-1000 m). Here we describe the overall
structure of the food web using in situ measurements of hydrodynamic parameters,
nutrients, phytoplankton distribution, primary production, and the biomass of zooplankton
and micronekton. We pay particular attention to the inter-comparability of the zooplankton
and micronekton sampling techniques (nets and acoustics) used, and the application of
acoustic techniques to improve estimates of micronekton in the future. We also use the
collected data to assess measures of micronekton estimated using the ecosystem model
SEAPODYM (Lehodey et al., 2010). Finally we interpret our findings in the context of the
broader southwest Pacific.

2 Methods and data

Two scientific cruises, Nectalis 1 and 2 were conducted onboard the R/V Alis from 29 July to
16 August 2011 (austral cool season - 18 sampling stations) and 26 November to 14
December 2011 (austral hot season - 23 sampling stations) within the New Caledonian EEZ
(Figure 2 and 3). The two cruises were conducted on approximately the same track, with
some differences due to weather conditions. The potential spatial variability introduced by
variability in station positions between cruises was considered minimal in view of the

162	variability of this highly dynamic pelagic system. Details of station sampling and continuous
163	measurements are summarized in Table 1 and detailed below. Comparisons of ensuing data
164	made between cruises included all stations.
165	2.1 Data collected during the cruises
166	2.1.1 S-ADCP currents
167	Five minute averaged ocean currents were acquired from 8 m bins across 16 to 200 m depth
168	using a ship-borne 153 kHz Acoustic Doppler Current Profiler (ADCP – Teledyne RD
169	Instrument, Seattle, USA). These velocity profiles were edited and processed using the
170	CODAS software,
171	(http://currents.soest.hawaii.edu/docs/adcp_doc/codas_setup/index.html) following the
172	procedure of Hummon and Firing (2003). Data presented are averages over the top 150 m.
173	2.1.2 Temperature and salinity
174	An on-board thermosalinograph continuously measured sea surface temperature (SST) and
175	salinity (SSS). At each station, Conductivity Temperature Depth (CTD) casts down to 500 m
176	recorded continuous vertical profiles of temperature and salinity. CTD data were checked
177	for spurious values using the Seasoft software (Sea-Bird electronics, Washington, USA),
178	binned at 1m intervals and presented for the top 200 m.
179	2.1.3 Water sampling
180	Water was sampled during the CTD casts using 8 L Niskin bottles to measure nutrients,
181	chlorophyll, phytoplankton cell counts, photosynthetic pigments and primary production.
182	Depth and frequency of sampling varied according to variables measured and associated
183	analyses (Table 1).
184	2.1.4 Nutrients
185	Nitrate, phosphate (Soluble Reactive Phosphorus: SRP) levels were measured in HgCl ₂ -
186	poisoned samples and analyzed in the laboratory within two months of the end of the
187	cruises using an Auto-analyzer AA3 (Bran+Luebbe, Norderstedt, Germany), as described in
188	Aminot and Kérouel (2007). Nitrate and nitrite (reported as NO ₃) concentrations were
189	determined at nanomolar precision (Raimbault et al., 1990). SRP concentrations (reported

190	as PO ₄) were analyzed according to Murphy and Riley (1962). Data were interpolated to plot
191	the 0-180 m vertical profiles using Dr Masson's SAXO package
192	(http://forge.ipsl.jussieu.fr/saxo/download/xmldoc/whatissaxo.html) based on IDL
193	(Interactive Data Langage, Exelisvis, Boulder, USA).
194	2.1.5 Phytoplankton: biomass and community structure
195	Phytoplankton composition and community structure were identified from water samples
196	collected (Table 1) and results were averaged across the depths: 0-50 m and 50-130 m.
197	2.1.5.1 Chlorophyll
198	In situ chlorophyll a (Chl-a) values were determined after methanol extraction (Le Bouteiller
199	et al., 1992), using a Turner Design fluorometer (Turner Designs, Sunnyvale, California, USA,
200	module # 7200-040, Chl-a extracted-acidification) calibrated with pure Chl-a standard
201	(Sigma). Total Chl-a concentrations were determined from 0.5 L water samples filtered onto
202	GF/F Whatman filters. Size-fractioned Chl-a across the size classes <3 μm , 3-10 μm and
203	>10 μm was determined from 2 L water samples collected onto 10 μm , 3 μm nucleopore
204	and GF/F filters by in-line serial filtrations, and represented proxies of pico-, nano and
205	microphytoplankton biomasses respectively. The mean and standard deviation of size-
206	fractionated Chl-a percentages were calculated for each cruise. Total Chl-a data were
207	interpolated to plot the 0-150 m vertical sections of each cruise using the SAXO package.
208	2.1.5.2 Cell counts by flow cytometry (FCM)
209	Water samples of 1.1 mL were fixed by adding paraformaldehyde solution (2% final
210	concentration) and then frozen in liquid nitrogen on board. Cell counts for pico and
211	nanophytoplankton (<3 μm , 3-10 μm respectively) were performed with a FACSCalibur flow
212	cytometer (BD Biosciences, San Jose, California, USA) at the Regional Flow Cytometry
213	Platform for Microbiology (PRECYM) (http://precym.com.univ-mrs.fr). Data were
214	normalized using both Fluoresbrite® Fluorescent Microspheres (Polysciences Inc. Europe)
215	and TruCountTM beads (BD) and the mean and standard deviation of cell count percentages
216	were calculated for each cruise.

217	2.1.5.3 Phycoerythrin
218	Water samples (4.5 L) were filtered onto 0.4 μm Nucleopore polycarbonate membrane
219	filters (47 mm diameter) and immediately frozen in liquid nitrogen until analysis. Using
220	methods described in Neveux et al. (2009), phycoerythrin (PE) was extracted in a 4 mL
221	glycerol-phosphate mixture (50/50) after vigorous shaking for resuspension of particles
222	(Wyman, 1992). Using a Perkin Elmer LS55 spectrofluorometer (PerkinElmer, Inc., Waltham
223	Massachusetts, USA) and emission and excitation slit widths adjusted to 5 and 10 nm,
224	respectively, the PE fluorescence excitation spectra were recorded between 450 and
225	580 nm (emission fixed at 605 nm). Quantitative estimates of phycoerythrin were obtained
226	from the area below the fluorescence excitation curve, after filter blank subtraction and the
227	mean and standard deviation calculated for each cruise.
228	2.1.6 Primary production
229	Net primary production (NPP, mgC m ⁻³ d ⁻¹ , Table 1) was measured using the ¹⁴ C tracer
230	technique (RochelleNewall et al., 2008). Water samples (76 mL) were inoculated with
231	0.40 MBq of a sodium ¹⁴ C bicarbonate solution (Perkin Elmer, initial concentration
232	37 MBq mL ⁻¹) and immediately placed in a thermoregulated (22-24°C) photosynthetron to
233	incubate samples at varying light levels (11%, 28%, 48%, 68%, 100%). After 1.5 h incubation
234	samples were filtered onto 0.4 μm polycarbonate filters (25 mm Whatman Cyclopore) which
235	were then placed into clean glass liquid scintillation counting vials and stored at -20 °C. In
236	the laboratory, 100 μL of 0.5N HCl was added to each sample, and the vial left open for 12 h
237	under a fume hood to remove unfixed ¹⁴ C. After acidification and drying, 5 mL of
238	scintillation cocktail (Ultima Gold MV, Parkard instruments) was added to each sample, and
239	the samples analyzed in a Packard Tri-Carb (1600TR) Liquid Scintillation Counter
240	(PerkinElmer, Inc., Waltham, Massachusetts, USA). The mean and standard deviation were
241	calculated for each cruise.
242	2.1.7 Zooplankton
243	Three methods were used to estimate zooplankton (organisms 2 μ m – 20 mm) biomass: a
244	Tracor Acoustic Profiling System (TAPS), net sampling and Ship-Borne Acoustic Doppler
245	Current Profilers (S-ADCP)

246	2.1.7.1 Tracor Acoustic Profiling System (TAPS)
247	The TAPS-6™ (BAE systems, San Diego, CA, USA) is a six frequency (265, 420, 710, 1100,
248	1850, 3000 kHz) profiler (Holliday and Pieper, 1980) used to acoustically detect the micro-
249	(20-200 $\mu\text{m})$ and meso- (200-2000 $\mu\text{m})$ zooplankton from the surface down to 200 m. The
250	TAPS-6 was used in "cast mode", profiling the water column in horizontal position with a
251	descent speed of 0.5 m s ⁻¹ , sampling a volume of about 5 L of water at each ping (ping rate:
252	$2.63 \ pings \ s^{-1}$). The TAPS-6 focused on small and abundant organisms such as copepods,
253	with larger and less abundant organisms such as euphausiids having less chance to pass
254	through this small volume (Pieper et al., 2001).
255	The Scattering Volume (Sv) signal (in dB) was transformed into biovolume estimates using
256	an inversion algorithm following the method applied by Lebourges-Dhaussy et al. (2014) and
257	successfully applied to small zooplankton (e.g. Holliday et al., 1989; Lebourges-Dhaussy et
258	al., 2009; Napp et al., 1993; Pieper et al., 1990). The algorithm provided vectors of
259	abundances per size range for each station, from which biovolumes were estimated in
260	$\text{mm}^3~\text{m}^{-3}$ and converted into mg m^{-3} using a density factor of ~1 kg L^{-1} (Simmonds and
261	MacLennan, 2005). The size range of organisms explored in the inversion process was 0.05-
262	3 mm (micro- and meso-zooplankton).
263	2.1.7.2 Zooplankton net sampling
264	Five layers of the water column were sampled from the surface down to 600 m depth (0-
265	100, 100-200, 200-400, 400-500, 500-600 m) using an Hydrobios MultiNet (Hydrobios, Kiel,
266	Germany). Each of the nets used were comprised of 200 μm nylon mesh and equipped with
267	a mechanical Hydrobios flowmeter. The volume filtered by each net was calculated using
268	the following equation:
269	V=d*k*A
270	where d is the number of revolutions of the flowmeter, k=0.3 m/revolution is the pitch of
271	the impeller of the flowmeter provided by the manufacturer (Hydro-Bios Apparatebau
272	GmbH. 2009) and A is the size of the net mouth area (0.25 m ²).

2/3	Samples collected by the nets were inimediately preserved in a 5 % buffered formalin-
274	seawater solution and processed for wet and dry weight analysis later in the laboratory. Dry
275	weights (DW) and wet weights (WW) were determined for the 0-200 m and the 0-600 m
276	layers respectively.
277	2.1.7.3 Ship-Borne Acoustic Doppler Current Profilers (S-ADCP) backscatter
278	The S-ADCP (see section 2.1.1), was also used to provide relative measures of acoustic
279	density, as a proxy for zooplankton to micronekton biomass (Flagg and Smith, 1989;
280	Heywood et al., 1991; Menkes et al., 2002; Radenac et al., 2010). At 153 kHz, this instrument
281	roughly detects organisms across the size ranges of a few millimeters to a few centimeters
282	(Sutor et al., 2005). The ADCP echo intensity (E_a) was converted into S_{ν} (in dB) using the
283	equation from Deines (1999) modified by Gostiaux and van Haren (2010):
284	$Sv = C + 10log_{10}[(T_x + 273.16)R^2] - L_{DBM} - P_{DBW} + 2\alpha R + 10log_{10}[10^{KcEa/10} - 10^{KcEnoise/10}]$
285	where T_x is the temperature of the transducer (°C), L_{DBM} is $10log_{10}$ (transmit pulse, in
286	meters), P_{DBW} is $10log_{10}$ (transmit power, in Watts), R is depth along the beam (m), α is the
287	sound absorption coefficient (dB/m) in water, K_c is a conversion factor for echo intensity
288	(dB/counts), E_a is the ADCP raw echo intensity (counts) and E_{noise} is the noise (counts). We
289	used the default parameters given in Deines (1999) for the constants C and $P_{\text{DBW}}.$ During the
290	time that the ship was stationary at each station, when ship noise is reduced, we selected
291	the minimum value of the echo intensities E_{a} in the vertical profiles and the minima were
292	then averaged over the entire cruise to obtain E _{noise} .
293	2.1.8 Micronekton
294	Three methods were used to estimate micronekton (organisms 2 - 20 cm) biomass and
295	species composition: using an EK60 echosounder, net sampling and the S-ADCP (see section
296	2.1.7.3).
297	2.1.8.1 EK 60 echosounder
298	Acoustic data were collected continuously during the cruise using a EK60 echosounder
299	(SIMRAD Kongsberg Maritime AS, Horten, Norway) with four hull-mounted split-beam
300	transducers (38, 70, 120 and 200 kHz). Echosounder calibration was performed according to

301	Foote et al. (1987) at the beginning of each cruise. Due to the presence of noise in
302	echograms, linked to the specificities of the installation of the sounder on the R/V Alis and
303	to rough seas during the cruises, the water column was only sampled down to depths of
304	100, 200, 250 and 600 m for the 200, 120, 70 and 38 kHz channels respectively. A data
305	cleaning step was performed with Matlab® (MathWorks, Natick, Massachusetts, USA)
306	filtering tools provided with the Movies3D software (IFREMER). The EK60 signal was
307	analyzed in terms of scattering volume (Sv) (MacLennan et al., 2002). It was not possible to
308	calculate micronekton biomass from echograms produced as the Sv to biomass conversion
309	requires knowledge of the acoustic properties of the detected organisms added to a
310	complex inversion of the signal and has not yet been performed for our dataset. The 38 kHz
311	frequency is commonly used as a proxy for micronekton (Bertrand et al., 1999; Kloser et al.,
312	2009; McClatchie and Dunford, 2003) and was used to represent micronekton over 0-600 m
313	To describe the spatial structure of the micronekton biomass derived from the 38 kHz EK60,
314	we removed the day/night signal from the data as the strong diurnal vertical migration of
315	micronekton might mask spatial patterns. The data were assigned to either day or night and
316	average values were calculated for each period for each cruise. The daytime (resp.
317	nighttime) mean was subtracted from the daytime (resp. nighttime) values to produce
318	anomalies for each period.
319	2.1.8.2 Micronekton net sampling
320	Micronekton were sampled at each station with a mid-water trawl with a 10 mm codend
321	mesh size. Vertical and horizontal mouth opening of ~10 m each were monitored with trawl
322	opening sensors (Scanmar, Åsgårdstrand, Norway). Horizontal tows were conducted to
323	target aggregations visually detected with the EK60 echosounder. Once the trawl net was
324	stabilised at the chosen depth, it was towed for 30 minutes at 3-4 knots. One or two tows
325	were conducted at each sampling station between 14 and 130 m at night and between 21
326	and 540 m during the day. Organisms were sorted on-board into groups and frozen. In the
327	laboratory, samples were identified at the lowest taxonomic level possible, counted,
328	measured and weighed. Gelatinous organisms (e.g. siphonophores, salps, pyrosomes) were

weighed frozen as a group. Biomass was expressed as mg of wet weight per m³ filtered. The 329 330 volume of water filtered by the net was calculated as: 331 V=S*D, 332 with S=h*v and D= R*c, $c = 2 * \arctan(\sqrt{a/(1-a)})$ 333 and $a=[\sin((lat_2-lat_1)/2)]^2 + \cos(lat_1)*\cos(lat_2)*[\sin((lon_2-lon_1)/2)]^2$ 334 where V is the volume filtered (m³), S is the net mouth opening (m²), h and v are the net 335 horizontal and vertical mouth opening (m), D is the distance covered by the trawl (m), 336 R=6371.e⁺³ m is the earth radius, lat₁, lat₂, lon₁, lon₂ are the latitude and longitude of the 337 338 start and the end of the set (radian). 339 2.2 Other in situ, satellite and model derived datasets 340 341 Estimated oceanographic and biological parameters derived from remote sensing and 342 physical and biological models were used to undertake direct comparisons between in situ data and satellite and model derived parameter estimates and investigate relationships of in 343 344 situ data collected during each cruise with broader scale regional ocean dynamics. 345 2.2.1 Ocean Currents 346 We used two datasets, the Kessler and Cravatte (2013) in situ dataset and the Ocean 347 Surface Current Analysis (OSCAR, http://www.oscar.noaa.gov/) satellite-derived dataset. 348 The first describes the time-averaged total geostrophic circulation of the top 1000 m. The 349 second provides surface currents estimated from a combination of data derived from drifting buoys and altimetry at a 5-day and 1/3° resolution. 350 351 2.2.2 Eddies: Okubo-Weiß parameter 352 Surface ocean dynamics were examined using an eddy detection algorithm. The Okubo-353 Weiß (OKW) parameter was calculated from the OSCAR surface currents. It describes the

354	deformation (shear and strain) and rotation (vorticity) of surface currents (Chelton et al.,
355	2011b; d' Ovidio et al., 2013; Dutrieux et al., 2008). This parameter allows discrimination of
356	regions where fluids circulate in a closed loop (OKW < 0, e.g. in the interior of eddies where
357	vorticity is high) from regions where shear and strain are high (OKW $>$ 0, e.g. on the edges of
358	eddies where strain is high). The OKW parameter is always negative within vortices whether
359	they are cyclonic or anticyclonic (Chelton et al., 2011b).
360	2.2.3 Sea Level Anomaly (SLA)
361	Sea level anomalies (relative to the long term mean across the period 1993-2010) were
362	extracted from http://www.aviso.oceanobs.com/en/data/products/sea-surface-height-
363	products/global/msla.html#c5122 at a resolution of 1/3° and 7 days. SLA was used to
364	identify downwelling (high values or ridges in SLA) versus upwelling eddies (low values or
365	troughs in SLA).
366	2.2.4 Sea Surface Temperature (SST)
367	Daily SSTs from the Group for High Resolution SST (GHRSST) were downloaded from the
368	website https://www.ghrsst.org/ and used to examine spatial patterns in SSTs in the New
369	Caledonian EEZ. This freely available product combines several satellite data sources and is
370	provided at 1/12° grid resolution.
371	2.2.5 Primary production
372	Depth-integrated primary production was estimated from satellite-derived chlorophyll,
373	Photosynthetically Available Radiation (PAR) fields and SST fields using the Vertically
374	Generalized Production Model (VGPM) (Behrenfeld and Falkowski, 1997). Primary
375	production was integrated across the euphotic layer, which was statistically derived from
376	satellite imagery (http://www.science.oregonstate.edu/ocean.productivity/). Satellite-
377	derived chlorophyll is calculated from ocean color data, which, for the period 2002-2009,
378	were derived from the Sea-viewing Wide Field Of View Sensor (SeaWiFS) satellite after
379	which data were computed at CLS (www.cls.fr) using the VGPM model and Moderate
380	Resolution Imaging Spectroradiometer (MODIS) and Medium Resolution Imaging
381	Spectrometer (MERIS) satellite data. PAR data were derived from the European Center for
382	Medium Range Forecast (ECMWF) analyses.

383	2.2.6 Micronekton from the SEAPODYM model
384	The end-to end spatial ecosystem model SEAPODYM (Lehodey et al., 2008) describes the
385	interactions of tuna species with the environment and incorporates external forcings
386	associated with fishing and the environment. It includes environmental parameters such as
387	temperature, currents, oxygen and primary production as well as a micronekton sub-model
388	describing the transfer of energy from primary production to tuna species through mid-
389	trophic levels. The sub-model comprises six functional groups of micronekton occupying
390	different water layers according to day and night (diel migration model). Modelled
391	micronekton is advected by currents and assimilates carbon from primary production
392	produced three months earlier (Lehodey et al., 2010).
393	The micronekton sub-model is driven by satellite-derived primary production (see section
394	2.2.5) and by the outputs of the GLobal Ocean ReanalYsis and Simulations (GLORYS2.V1) of
395	currents and temperature produced by the French Groupe Mission Mercator Coriolis
396	(Barnier et al., 2006; Ferry et al., 2012) across two reanalysis periods 2002 – 2008 and 2009
397	2012. The 2002 – 2008 reanalysis was conducted at a daily and 1/4° resolution and was
398	performed by the MERCATOR-OCEAN operational oceanography center. It is forced by daily
399	surface meteorological data from the European Centre for Medium-Range Weather
400	Forecasts (ECMWF). By assimilating satellite-derived sea level anomalies, sea surface
401	temperatures and in situ measurements of vertical temperature and salinity profiles, the
402	model estimates realistic mesoscale activity with eddy field variability in good agreement
403	with altimetric data. Reanalysis across 2009-2012 included temperature and currents
404	provided by the same numerical ocean model, while altimetry, SST and temperature/salinity
405	profiles were also assimilated in their operational configuration (http://www.mercator-
406	ocean.fr/)(Abecassis et al., 2013).
407	The biomass distribution of micronekton functional groups along the cruise track and in the
408	south-west Pacific at the time of the cruises was estimated with the SEAPODYM ecosystem
409	model using a revised definition of vertical biological layer boundaries, at a spatial
410	resolution of 1/4° averaged over 7 days. Vertical biological layers comprised the epipelagic
411	layer, which lies between the surface and the euphotic depth (derived from ocean color

412	satellite data and the VGPM model), the mesopelagic layer located at 1-3 times the euphotic
413	depth, and the bathypelagic layer located at 3-7 times the euphotic depth. The model
414	simulates a diel behavior of micronekton by considering that during the night, daily
415	migratory species can move from one layer to another, thus adding to the residing biomass
416	of non-migratory species of the layer. Because we used 7-day outputs to compare with the
417	continuous Nectalis data, we reconstructed a SEAPODYM time series with a day/night signal
418	that mimicked the Nectalis data. We interpolated the Nectalis tracks into the SEAPODYM
419	model. This interpolation was temporally referenced so that day and night estimates from
420	SEAPODYM could be extracted.
421	To describe the spatial structure of the micronekton biomass derived from SEAPODYM, we
422	removed the day/night signal from the data by calculating anomalies following the same
423	procedure than for 38 kHz EK60 (see section 2.1.8.1).
424	
425	2.3 Statistics and comparison of methods
426	2.3.1 Primary production
427	The non-parametric rank-sum Wilcoxon-Mann-Whitney test at $\alpha \text{=-}5\%$ was used to test the
428	seasonal difference in in situ primary production values and in VGPM satellite-derived
429	primary production along the cruise track. Spatial auto-correlation was eliminated from the
430	VGPM dataset by building a new dataset of independent points before conducting the
431	seasonal comparison.
432	To build this new dataset of independent points, we determined the distance ("d" in km) at
433	which two points are independent. The initial auto-correlated dataset was then resampled
434	selecting a point every "d" kilometers creating a dataset of independent points. Shifting the
435	start position of this subsampling by 1 km, another dataset of independent points was then
436	created. This procedure was repeated until the number of resampled datasets of
437	independent points was equal to "d", and contained a number "n" of independent points
438	which was the length of the cruise track divided by "d". In the statistical tests the "d"
439	resampled datasets of independent points were all tested and the result of the test

440	(difference or no difference) comes with the percentage of the number of "d" tests
441	producing this result.
442	To estimate the distance "d" between two independent points, empirical spatial variograms
443	with 10km bins were used. The variogram of the VGPM dataset was compared to the noise
444	constructed from a series of variograms of 100 randomly re-ordered VGPM datasets by a
445	Monte Carlo procedure. The distance "d" at which the dataset points become uncorrelated
446	was estimated when the variogram of the dataset reached the noise.
447	For the VGPM dataset "d" was estimated at ~50 km during Nectalis1 and 100 km during
448	Nectalis2. Overall a conservative value of 100 km between two successive data points was
449	considered, prompting $^{\sim}$ 100 possible VGPM datasets of 30 independent points (or,
450	equivalently degrees of freedom) for Nectalis1 and 35 independent points for Nectalis2.
451	These 100-ensemble datasets were used for seasonal comparisons.
452	We estimated that in situ primary production points were independent (no spatial auto-
453	correlation) on the basis that the minimum distance between two sampling stations where
454	in situ primary production was measured (144 km) is greater than the estimate at which 2
455	points were determined to be independent using the VGPM dataset (100 km).
456	The Wilcoxon-Mann-Whitney test at α =5% was also used to compare <i>in situ</i> primary
457	production to VGPM data at the location of the <i>in situ</i> measures (12 data points). For small
458	sample sizes, values for significance were read in classical tables, while where sample sizes
459	were n>20 degree of freedom, the test was calculated using IDL's routines
460	(imsl_wilcoxon.pro).
461	2.3.2 Zooplankton
462	A Wilcoxon-Mann-Whitney test at α =5% was used to test for seasonal difference in
463	zooplankton biomasses estimated from the zooplankton net (in DW and WW), the TAPS and
464	the S-ADCP backscatter. We accounted for the spatial auto-correlation of the S-ADCP by
465	following the procedure described in section 2.3.1. For the S-ADCP dataset, the distance "d"
466	at which two points were considered independent was 30 km for both cruises.
467	Consequently, 30 datasets of 100 and 116 independent points were built for Nectalis1 and

468	Nectalis2 respectively for statistical analyses. We estimated that <i>in situ</i> zooplankton
469	biomass estimates (zooplankton net and TAPS) points were independent (no spatial auto-
470	correlation) on the basis that the minimum distance between two sampling stations (67 km) $$
471	is greater than "d" (30 km).
472	The potential for the S-ADCP to provide a proxy of zooplankton biomass was evaluated
473	against the log-transformed biomass measurements of zooplankton derived from the TAPS
474	and net sampling using a Spearman's correlation.
475	For this general overview, no detailed examination of zooplankton spatial distribution and
476	composition were conducted, but will be conducted in a separate study (Smeti pers.
477	comm.).
478	2.3.3 Micronekton
479	A Wilcoxon-Mann-Whitney test at α =5% was used to test for seasonal differences in
480	micronekton biomass estimated by the EK60 and by the SEAPODYM ecosystem model. No
481	seasonal comparison was conducted on net sampling because of differences in sampling
482	strategies between the two cruises (non-comparable depth or day-night tows). We
483	accounted for the spatial auto-correlation of the 38 kHz EK60 Sv and SEAPODYM
484	micronekton biomass estimates by following the procedure described in section 2.3.1. For
485	the 38 kHz EK60 Sv dataset the distance "d" at which two points were considered
486	independent was 30 km for both cruises. Consequently, 30 datasets of 100 and 116
487	independent points were built for Nectalis1 and Nectalis2 respectively for statistical
488	analyses. For the SEAPODYM dataset, "d" was estimated at 100 km for Nectalis1 and 50 km
489	for Nectalis2. A conservative value of "d"=100 km was used for both cruises to build 100
490	datasets of 30 and 35 independent points for Nectalis1 and Nectalis2 respectively.
491	The potential for the S-ADCP to provide a proxy of micronekton biomass was evaluated
492	against the Sv values of the four frequencies of the EK60 echosounder. To do this, the EK60
493	high-resolution time and vertical profiles were averaged to the ADCP time/vertical
494	resolution. The Sv was calculated for a 5-minute Elementary Sampling Unit (ESU) and 8 m
495	height layers. Correlations between data provided by the S-ADCP and the four frequencies

496	of the EK60 were investigated using a Spearman's correlation. We accounted for the spatial
497	auto-correlation by computing 30 correlation coefficients from the 30 resampled datasets of
498	independent points distant of 30 km, as explained above. The range of correlation
499	coefficients from the 30 correlation ensemble was provided as well as the percentage of
500	these correlations significant at α =5% level.
501	Estimates of epi and mesopelagic micronekton biomass (mg m ⁻³) derived from SEAPODYM
502	were compared to the estimates of micronekton biomass derived from the 38 kHz EK60
503	echosounder. The high-resolution EK60 data were averaged across ¼ degree grid squares
504	along the cruise track to correspond with the spatial resolution of the SEAPODYM
505	ecosystem model. Correlation between the two data series was investigated applying the
506	same procedure as for EK60 $\emph{vs.}$ S-ADCP. and accounting for the Spatial auto-correlation was
507	accounted for by calculating the correlation on 100 resampled datasets of independent
508	points distant of 100 km from the two biomass series

3 Results

3.1 Physical oceanography and biogeochemistry

512 3.1.1 Surface features

During the cool season (Nectalis1) thermosalinograph measurements showed that surface waters in the southern part of the cruise track (south of 19°S) had an average temperature of 23.6 ± 1.0 °C and salinity of 35.2 ± 0.2 while the northern part of the cruise was characterized by waters of 25.3 ± 0.7 °C and 35.0 ± 0.1 (Figure 2). During the hot season (Nectalis2), overall SST and SSS patterns were similar, although temperatures were warmer by ~3°C. Salinity was very similar in the south but lower by ~0.1 in the north (Figure 3). During both cruises, salinity varied across similar gradients to temperature, but in the opposite direction at both larger-scales and smaller scales. For example, high temperature and low salinity waters were observed during Nectalis1 at stations 4 and 5 with waters with

522	particularly low temperature and high salinity observed at stations 1 and 2 during both
523	cruises. This gradient was observed in particular during Nectalis2.
524	During both cruises, ADCP surface layer (0-150 m) currents varied in a similar way across
525	large and small spatial scales to surface-only currents derived from OSCAR. North of 19°S,
526	the currents were predominantly directed westward, while in the south they were
527	predominantly directed eastward. Along the western coast of the main island of New
528	Caledonia, currents were flowing predominantly south-eastward during Nectalis1 and
529	southward during Nectalis2. In addition to these broad scale patterns, high current
530	variability was observed, for example at stations 6-7 during Nectalis1 and stations 7-8-9
531	during Nectalis2.
532	Satellite-derived SST clearly showed the large-scale north-south gradient in observed during
533	both cruises. Smaller scale meandering of the SST field centered at ~19-20°S was observed
534	with strong association between the thermohaline patterns and the currents. Meanders
535	were noted through intrusions of warmer waters from the north which were advected
536	south (e.g. Nectalis1 station 8, 17, Nectalis2 station 9) and intrusions of cooler waters from
537	the south which were advected north (e.g. Nectalis1 stations 6, 7, 16, Nectalis2 stations 17-
538	18).
539	Values of OKW and SLAs along with current vectors from both OSCAR and S-ADCP described
540	an important turbulent eddy activity. Cyclonic eddies ("upwelling" type eddy) corresponding
541	to sea level depression (or equivalently, thermocline uplifting) were observed during the
542	cool season, for example at $^{\sim}23^{\circ}S$ 161 $^{\circ}E$ (eddy A on Figure 2), at stations 6-7 (eddy B) and at
543	~16.5°S 159°E (eddy C) the south-eastern edge of which was sampled at station 10 (Figure
544	2). During the hot season, strong cyclonic eddies were observed at $^{\sim}24^{\circ}\text{S}$ 156°E (eddy D on
545	Figure 3), ~24°S 164°E (eddy E) and ~25°S 172°E (eddy G) and a series of energetic eddies
546	were observed between stations 7 and 18 (eddies H, I, J). The edge and the center of an
547	anticyclonic eddy ("downwelling" type eddy with thermocline deepening and sea-level
548	ridge) were sampled during the hot season at stations 8 and 9 respectively (eddy K, Figure
549	3). Overall, lower eddy activity was observed during the cool season (Figure 2) than during
550	the hot season (Figure 3).

551	3.1.2 Vertical structures
552	During the cool season, stations 8 - 18 in the north and east were characterized by relatively
553	warm, low salinity waters with a mixed layer depth of $^{\sim}60\text{m}$ and low values of nitrate
554	(0.05 \pm 0.07 $\mu M)$ to a depth of ~90 m (Figure 4). The Deep Chlorophyll Maxima (DCM)
555	($^{\sim}0.25$ -0.3 mg m $^{^{-3}}$) and the nutricline were located at $^{\sim}90$ m depth (Figure 4). By contrast,
556	stations 1 - 7 in the south and west were comparatively cooler with higher surface salinity.
557	At these stations, higher concentrations of nitrate (0.13 \pm 0.12 $\mu\text{M})$ and chlorophyll content
558	occurred with more frequent maxima at the surface. Phosphate concentrations varied with
559	an average of 0.067 \pm 0.038 μM from the surface to 100 m depth and were occasionally
560	lower than 0.05 μM in the surface layer. Within this general pattern, a number of stations
561	demonstrated unique characteristics. Cool, highly saline waters which were homogeneous
562	down to 100 m with a shallow nutricline and enhanced chlorophyll were recorded at
563	stations 1 and 2 (Figure 4). Waters with a deep mixed layer, elevated chlorophyll from the
564	surface down to 100 m were also recorded at stations 6 and 7, contrasting with the
565	surrounding waters (Figure 4). High surface (0-20 m) chlorophyll levels (0.23 $\mu g \ l^{-1}$) were
566	recorded at station 10 (Figure 4).
567	The north-south gradient in temperature and salinity observed during the cool season was
568	also evident at depth during the hot season, with warmer and fresher waters north of ~20°S
569	observed at stations 8 - 19 (Figures 3 and 4) and cooler and saltier waters south of $^{\sim}20^{\circ}\text{S}$
570	observed at stations 1 - 7 and 20 to 23 . The mixed layer across all stations was shallower
571	during the hot season, located at $^{\sim}25$ m, denoting stronger surface stratification in the
572	water column than during the cool season. Surface waters were low in nitrate
573	(0.03 \pm 0.02 $\mu\text{M})$ across almost the entire cruise track (Figure 4) and the DCM was often
574	centered at around 100 m with mean values of $^{\sim}0.41\pm0.16$ mg m $^{^{-3}}$. In general, above the
575	DCM in the top $^{\sim}$ 50 m, chlorophyll concentrations were slightly lower than during the cool
576	season (Figure 4), particularly in the southern part of the survey area. Phosphate tended to
577	be low (0.05 \pm 0.03 $\mu M)$ for stations 1 - 7 and 20 - 23 in the southern part of the cruise track
578	in comparison to stations in the northern part of the cruise (0.09 \pm 0.03 $\mu\text{M}).$ A few stations
579	had unusual characteristics: high temperatures and low salinities down to 100 m were

580	observed at station 9 and surface nitrate was slightly enhanced at stations 7 and 8
581	(0.09 \pm 0.01 $\mu\text{M})$ compared to other stations (0.02 \pm 0.02 $\mu\text{M}).$
582	
583	3.2 Primary production
584	Depth- integrated measurements of in situ primary production in the photic layer and
585	similar satellite-derived net primary production (NPP) from VGPM along the cruise track
586	were significantly higher during the cool season (352±160 mgC m ⁻² d ⁻¹ and
587	301±62 mgC m ⁻² d ⁻¹ on average respectively) than during the hot season
588	(231 \pm 133 mgC m ⁻² d ⁻¹ and 199 \pm 55 mgC m ⁻² d ⁻¹) (Figure 5 and Table 2)
589	The NPP pattern (Figure 2) demonstrated a strong gradient during the cool season with
590	values of $^{\sim}350~\text{mgC}~\text{m}^{^{-2}}~\text{d}^{^{-1}}$ in the southern part of the survey area (south of 20°S and west
591	of the main island) and values lower than 200 mgC m ⁻² d ⁻¹ in the northern part of the survey
592	area (north of 20°S and east of the main island). During the hot season the entire region was
593	more oligotrophic, with a weaker north-south gradient, and average values of
594	$^{\sim}200~\text{mgC}~\text{m}^{^{-2}}~\text{d}^{^{-1}}$ in the survey area (Figure 3). Within this large-scale gradient, specific
595	patterns linked to mesoscale structures were observed. For example, the center of some
596	eddies were characterized by enhanced primary production (e.g. Nectalis1 eddy A; Nectalis2
597	eddy D; Nectalis2 eddy G), while primary production was enhanced at the edge of others
598	(e.g. Nectalis1 station10 eddy C; Nectalis2 series of eddies H, I, J).
599	No significant differences were found between in situ production estimates and VGPM
600	satellite values at the in situ sample locations (Figure 5).
601	
602	3.3 Phytoplankton
603	During the cool season, size fractionated chlorophyll was dominated by picophytoplankton
604	(< 3 μ m) across all stations (mean=75.9% ± SD=17.2% in biomass); nano and micro-
605	phytoplankton represented 12.8% ± 9.6% and 11.3% ± 12.6% respectively of chlorophyll
606	biomass. The cyanobacteria <i>Prochlorococcus</i> were the dominant species of the

607	picophytoplankton group (91.9% \pm 6.3% in abundance; Figure 4) with cell abundances of up
608	to 250 x 10 ³ mL ⁻¹ . Remaining abundances of picophytoplankton across stations were
609	comprised of Synechococcus (6.3% \pm 6.2%) and picoeukaryotes (1.8% \pm 0.8%). Overall
610	phytoplankton composition did not vary latitudinally or longitudinally, with the exception of
611	particular features observed at stations 2 and 10 (eddy C). In comparison to other stations,
612	higher proportions of large cells were observed at station 2 from the surface to 150 m
613	(~30% nano- and ~28% microphytoplankton in abundance) and at station 10 from the
614	surface to 50 m ($^{\sim}17\%$ nano- and $^{\sim}31\%$ microphytoplankton in abundance).
615	The fractionated chlorophyll and community structure during the hot season was similar to
616	that observed in the cool season with picophytoplankton and <i>Prochlorococcus</i> dominating
617	the communities (83.4% \pm 10.4% in biomass and 92.3% \pm 7.8% in abundance respectively;
618	Figure 4). Nano and micro-phytoplankton represented $8.6\% \pm 5.4\%$ and $7.9\% \pm 7.1\%$
619	respectively. However, cell abundance was much lower during the hot season, with
620	maximum cell counts of <i>Prochlorococcus</i> of 160 x 10 ³ mL ⁻¹ . Remaining cell abundances were
621	comprised of Synechococcus (5.0% \pm 6.8%) and picoeukaryotes (2.7% \pm 1.7%). Again, the
622	phytoplankton structure was relatively homogeneous along the cruise track with the
623	exception of station 9 (eddy K) which had a higher proportion of larger cells from the
624	surface to 50 m ($^{\sim}7\%$ nano- and $^{\sim}39\%$ microphytoplankton in abundance) than the rest of
625	the stations (~13% \pm 3.1% nano- and ~11% \pm 5.0% microphytoplankton in abundance; Figure
626	4). Phycoerythrin (PE) concentration was also much higher at this station (1836 fluorescence
627	unit $vs.$ 389 ± 359 fluorescence unit for the other stations).
628	3.4 Zooplankton
629	Diurnal variability in zooplankton biomass was observed with all methods, with enhanced
630	biomass at night in the top 200 m during both cruises (Figure 6 and 7). Zooplankton WWs
631	during the hot season however were relatively similar during the day and at night, which is
632	at odds with dry weight (DW) estimates where diurnal variability is evident (Figure 6 and 7).
633	Zooplankton wet weight (WW) (Figure 6) vertical profiles showed that the majority of the
634	biomass concentrated in the top 100 m, deeper biomass rapidly decreased.

635	Mean biomass estimates from the TAPS ($^{\sim}$ 100 mg m $^{-3}$) were more than one order of
636	magnitude higher than WW estimates from net samples (< 6.5 mg m ⁻³) and DW estimates
637	from net samples (< $6~\text{mg m}^{-3}$) (Table 2). Zooplankton biomasses derived from TAPS and
638	WW estimates across all sampling stations were not significantly different between the two
639	cruises, while significant differences were observed in DW (Nectalis1 <nectalis2) and="" s-adcp<="" td=""></nectalis2)>
640	(Nectalis1>Nectalis2) estimates (Table 2).
641	Correlations between acoustic biomass proxies and net biomass measures for zooplankton
642	were all significant, with the TAPS and S-ADCP having the highest correlation overall (Table
643	3). Estimates derived from net samples demonstrated similar correlations with those
644	derived from the S-ADCP and those derived from the TAPS (Table 3). Correlation values
645	between zooplankton measurements were roughly similar across the two depth ranges
646	explored: 0-100 m and 0-200 m.
647	
648	3.5 Micronekton
649	Preliminary examination of the micronekton composition indicated that micronekton net
650	catch was dominated by gelatinous organisms (e.g. siphonophores, salps, pyrosomes),
651	which represented 53.8% of the overall wet weight biomass. Fish, molluscs and crustaceans
652	represented 36.5%, 7.6% and 2.1% of the biomass, respectively. In total, approximately 480
653	taxa were identified, including $^{\sim}240$ fish taxa, $^{\sim}95$ crustacean taxa, $^{\sim}85$ mollusc taxa and $^{\sim}60$
654	gelatinous organism taxa. Of those species able to be identified, those species with the
655	highest biomasses in each taxa group were the lanternfish Ceratoscopelus warmingii,
656	Hygophum hygomii and Diaphus perspicillatus; the molluscs Sthenoteuthis oualaniensis,
657	Abraliopsis sp. and Abralia omiae; and the crustaceans Thysanopoda tricuspidata,
658	Thysanopoda cristata and Euphausia mucronata. Of the gelatinous organisms the most
658 659	

Biomass estimates from in situ measurements from the micronekton nets for the 0-600 m

and from SEAPODYM model were in the same range: $^{\sim}$ 4 mg m $^{^{-3}}$ (Table 2).

660

662	During both seasons, the EK60 and SEAPODYM signal anomalies indicated that the region
663	north of ~19°S-20°S had lower micronekton biomass than the region south of this latitude
664	(Figure 8). Smaller scale variability was also apparent in both datasets, most prominently
665	south of 19°S-20°S where patches of higher biomass were observed; for example along the
666	west coast of the main island and at ~20.5°S 161°E during Nectalis1 and ~20.5°S 158°E
667	during Nectalis2.
668	Micronekton abundance estimated from the S-ADCP, the nets, the EK60 and the SEAPODYM
669	model exhibited a clear maximum at night (Figure 7). Vertical profiles of the micronekton
670	estimated from the 38 kHz EK60 Sv (Figure 6) demonstrated a bimodal distribution with
671	higher micronekton biomass estimates occurring at 0-200 m and 400-600 m than at other
672	depths during both the day and night.
673	Seasonal differences observed in micronekton biomass estimated by the EK60 Sv and
674	SEAPODYM were not statistically significant (Table 2). Conversely, estimates derived from
675	the S-ADCP were different with higher values during the cool season (Table 2).
676	Micronekton estimates derived from the EK60 Sv and S-ADCP Sv were highly correlated, and
677	the highest correlation was observed with the 70 kHz EK60 (correlation range = 0.87-0.96)
678	(Table 4). Micronekton biomass estimates calculated by the 38 kHz EK60 Sv were highly
679	correlated with estimates derived from the SEAPODYM model (correlation range = 0.73-
680	0.79) (Table 4).
681	
682	4 Discussion
683	4.1 Oligotrophic waters and water masses
684	The physical, biogeochemical and biological data collected during the two Nectalis cruises,
685	in two contrasting seasons, have provided new insights into the spatial and temporal
686	dynamics of the pelagic ecosystem in the waters around New Caledonia. Observations
687	collected from the two cruises support prior characterization of the region as oligotrophic.
688	The vertical nutrient profiles, low nitrate and sometimes low phosphate, low primary

689	production and chlorophyll biomass, and a phytoplankton composition dominated by small
690	size cells (picophytoplankton), were consistent with previous studies in South Pacific region
691	(Campbell et al., 2005; Jacquet et al., 2006; Young et al., 2011) and are typical of a Low
692	Nutrient Low Chlorophyll (LNLC) system. Although it is generally thought that nitrate is the
693	main limiting nutrient in this oligotrophic region (Le Borgne et al., 2011), some
694	phytoplankton species may be limited by phosphate (Moutin et al., 2005) and this can
695	induce higher contributions of diazotrophs such as Trichodesmium sp. in this area.
696	Trichodesmium sp. was not observed in the samples we collected, but it was seen at the
697	surface of the water along the track at one occasion. Examination of isotope values
698	calculated from biological samples collected during the Nectalis cruises (Hunt et al., this
699	issue) suggests the contribution of diazotrophs to phytoplankton composition as previously
700	observed in the area (Campbell et al., 2005; Dupouy et al., 2011).
701	Two distinct water masses were encountered in the studied area. North of 19°S-20°S,
702	waters in the top 200 m were characterized by warm temperature, low salinity, low nitrate,
703	lower primary production and lower micronekton biomass estimates. These characteristics
704	are representative of the "Coral Sea" oligotrophic regime (Ceccarelli et al., 2013), and are
705	largely influenced by the warmer and fresher waters of the south Pacific convergence zone
706	(SPCZ) where the SEC predominantly flows,.
707	South of 19°S-20°S, waters are characterized by colder temperature, higher salinity, a
708	shallower nitracline, higher nitrate content in the surface layer, higher primary production
709	and higher micronekton biomass estimates and are under the influence of the South
710	Tropical Counter Current branches (Marchesiello et al., 2010).
711	Although water masses were variable latitudinally, phytoplankton compositions were very
712	similar throughout the whole area.
713	4.2 Horizontal advection, mesoscale and submesoscale phenomena
714	Large regional-scale organization of surface currents, SST, SSS and primary production was
715	observed to be strongly distorted by meanders and smaller scale phenomena under the
716	influence of horizontal advection from highly variable currents. The similarity of

717	temperature and salinity variations suggested the action of advection processes in
718	modifying salt and temperature at small scales. Numerous processes such as upwellings,
719	mesoscale (20-100 km, Lévy, 2008) eddies and submesoscale (2-20 km) fronts were
720	observed influencing the biological distributions in complex manners.
721	In the south/south-westward flowing the ALIS currents observed during the Nectalis cruises
722	along the west coast of New Caledonia (Marchesiello et al., 2010), an example of coastal
723	upwelling was observed at stations 1 and 2 during the two seasons. This coastal upwelling
724	was characterized by cool temperatures and high salinities observed to be homogeneous
725	down to 100 m during the cool season and down to 50 m during the hot season. During the
726	cool season the upwelling was also characterized by a shallow nutricline, enhanced
727	chlorophyll at the surface and higher proportion of large phytoplankton cells, which were
728	not observed during the hot season. A coastal upwelling induced by south-east trade winds
729	particularly during the hot season has been reported in a number of other studies
730	(Ganachaud et al., 2010; Marchesiello et al., 2010).
731	Observations during each season described quite turbulent ocean circulation with myriads
732	of small cyclonic and anticyclonic eddies of ~50-100 km in size. Such observations have also
733	been reported by Chelton et al. (2011b).
734	The region is known for its strong interactions between the SEC, which enters from the east,
735	the STCC flowing from the west and the tortuous topography of island masses and ocean
736	floor ridges. These interactions between the large scale currents and topography produce
737	non-linearities in the ocean currents (Couvelard et al., 2008; Marchesiello et al., 2010) which
738	can favor eddy developments. Eddies can also be associated with incoming Rossby waves
739	(Killworth et al., 2004) as well as barotropic instabilities resulting from the sheared
740	westward and eastward currents in the northern region of the EEZ (Figure 1). South of
741	~22°S, Rossby waves and baroclinic instabilities between the surface flowing STCC and the
742	deeper flowing SEC are also known to generate eddy activity as depicted in strong ocean
743	eddy kinetic energy which peaks during the hot season (Qiu et al., 2009).

744	Primary production and phytoplankton composition within eddies can differ depending
745	upon the oceanographic processes and underlying trophic mechanisms operating in time
746	and space. At the mesoscale, cyclonic eddies (southern hemisphere) induces upwellings
747	near eddy centers and "eddy pumping" (Martin and Richards, 2001; McGillicuddy et al.,
748	2007, 1998) of nutrients into the photic layer. Its effects are most commonly observed near
749	eddy centers where enhanced chlorophyll can be found. Conversely in downwelling eddies
750	(anticyclonic in the southern hemisphere) poorer waters are expected. Lateral advection of
751	pre-existing primary production gradients by eddies (Chelton et al., 2011a) or advective
752	concentration/dispersion of floating materials (Dandonneau et al., 2003) are also common
753	mechanisms and linked to mesoscale phenomena. Maximum impacts on phytoplankton are
754	expected at the eddy edge or out of eddies in association with the frontal submesoscale
755	dynamics. Vertical pumping may also occur within submesoscale structures produced by
756	eddy-eddy interactions through frontal and ageostrophic mechanisms ($e.g.$ Klein and
757	Lapeyre, 2009; Lévy, 2008).
758	A number of eddies and frontal oceanographic processes were observed during the Nectalis
759	cruises. The sampling resolution of both <i>in situ</i> and satellite data during the Nectalis cruises
760	was sufficient to observe mesoscale eddies (20-100 km scale). However the sampling
761	resolution was insufficient to differentiate between submesoscale fronts (2-20 km scale,
762	Lévy, 2008) and lateral advection.
763	Enhanced primary production was observed mainly south of ~20°S in the New Caledonia EEZ
764	at the center of several cyclonic eddies (e.g. Nectalis1 eddy A; Nectalis2 eddy D; Nectalis2
765	eddy G) suggesting the occurrence of eddy pumping. Lower primary production was
766	observed in the downwelling (anticyclonic) eddy at station 9 during Nectalis2 (eddy K). In
767	downwelling areas lateral advection from fluid convergence can concentrate floating
768	organisms (Dandonneau et al. 2003) such as the diazotrophic cyanobacterium
769	Trichodesmium which can be quite frequent in the region (Dupouy et al., 2011). At station 9
770	(Nectalis 2) a higher proportion of large phytoplankton cells with higher concentration of
771	phycoerythrin suggested the presence of <i>Trichodesmium</i> , consistent with lateral advection
772	accumulation.

773	Enhanced primary production and chlorophyll were observed more commonly at the edge
774	of several eddies to the north of $^{\sim}20^{\circ}\text{S}$ in the more oligotrophic regions. Enhanced primary
775	productivity may have resulted from chlorophyll advected into the area either from the
776	north via a series of cyclonic eddies (e.g. Nectalis2 series of cyclonic eddies H, I, J; Nectalis1
777	station 10 eddy C) or from the south (Nectalis1 stations 6-7 eddy B). The increased
778	proportion of larger phytoplankton cells at Nectalis1 station 10 (eddy C) is consistent with
779	evolution in composition of eddies with time. The phytoplankton community may have
780	developed in the north and aged along the eddy streamlines as it was advected to the south
781	by eddy currents. However this observation was not consistent for all eddies observed with
782	no specific phytoplankton composition observed at some eddies (e.g. Nectalis1 station 6
783	eddy B).
784	Overall, the primary production patterns around New Caledonia appeared to be more highly
785	dominated by horizontal advection rather than by vertical processes (direct eddy pumping).
786	The patchy and high frequency signal complicated the general understanding of the
787	ecosystem organization, as is often the case in oligotrophic waters. More generally, how
788	mesoscale eddies and the submesoscale structures affect primary production is still under
789	debate (Chelton et al., 2011a; Gruber et al., 2011; Klein and Lapeyre, 2009; Lévy, 2008) and
790	the Nectalis data suggests that there is not one particular mechanism at work during the
791	period of the cruises in this region of the South Pacific.
792	The effect of primary production dynamics at these scales on upper trophic levels are also
793	poorly understood because of the difficulty of accessing datasets spanning a wide range of
794	trophic levels at the scales relevant to eddies and submesoscale structures. Similarly to
795	primary production, the few examples of zooplankton organization around eddies (e.g.
796	Lebourges-Dhaussy et al., 2014; Menkes et al., 2002; Roman et al., 1995) show a variety of
797	organizations. The S-ADCP backscatter (not shown), EK60 Sv and SEAPODYM micronekton
798	data showed strong patchiness, especially in the south, indicating the influence of
799	mesoscale features on the organization of zooplankton and micronekton. Similarly to
800	zooplankton, the relationship between mesoscale features and micronekton distribution is

801

considered to be complex and not yet well understood (Béhagle et al., 2014; Domokos, 802 2009; Potier et al., 2014). 803 4.3 Seasonality 804 Observations collected during the Nectalis cruises reflected strong seasonality in 805 hydrodynamics and water column characteristics in response to the seasonal migration of 806 the solar heating and convective system of the SPCZ. The hot season was characterized by 807 warmer and fresher ocean conditions, increased eddy activity, lower NPP and 808 phytoplankton biomass, and higher stratification as modelled by Marchesiello et al. (2010), The cool season was characterized by lower eddy activity, higher NPP and phytoplankton 809 810 biomass. The NPP latitudinal gradient during the cool season mimicked the SST gradient, 811 indicating a tight coupling between ocean dynamics and phytoplankton growth. There was a 812 stronger decoupling between the surface temperature patterns and primary production 813 during the hot season, as expected in oligotrophic waters (Le Borgne et al., 2011). 814 Primary production almost doubled during the cool season compared to the hot season. 815 Contradictory seasonal signals for zooplankton and micronekton biomass, however were 816 provided by various sampling methods, resulting in an inability to determine seasonality in 817 mid-level organisms. Two hypotheses, possibly acting in combination, may explain the observations. Firstly, enhanced net primary production during the cool season may have 818 819 been largely due to enhanced recycling, with a small portion of the primary production 820 transmitted to higher trophic levels. Secondly, different turn-over times between 821 phytoplankton and zooplankton/micronekton may have induced a time decoupling and a 822 delay in transmission of primary production to secondary and tertiary levels. 823 It should be noted that, at the time of the cruises in 2011, the South Pacific was considered 824 to be in a weak La Niña state 825 (http://iri.columbia.edu/climate/ENSO/currentinfo/archive/201110/technical.html). In the 826 New Caledonia region, the expected response of the ocean to La Niña is a weakening of the 827 trade winds during the hot season and slightly warmer SST conditions (~+0.5°C in average)

828 with a slightly deeper thermocline (Menkes, 2012). This weak effect of ENSO in the New 829 Caledonian area however, is unlikely to bias the seasonal view from the two cruises. 4.4 Diel migration 830 831 Classical diel behavior of organisms migrating towards the surface at night and to deep 832 waters during the day was observed in both zooplankton and micronekton using acoustic 833 methods. Using net sampling however, the day-night difference in zooplankton DW was not 834 observed in the zooplankton WW for the 0-200 m depth during the hot season. This may be 835 explained by the increase of gelatinous organisms (mainly salps and doliolids) at the surface 836 during the hot season (H. Smeti, pers. com.) and their representation in WW and DW 837 estimates. Because gelatinous organisms comprise the largest group in WW estimates, their 838 diel behavior will dominate any diel signal for zooplankton. As they rarely migrate vertically, 839 little day-night differences in WW would be expected for zooplankton. The proportion of 840 DW biomass contributed by gelatinous organisms however is much smaller. The 841 predominance of diel vertical behavior in the other taxa groups then results in a diel signal 842 being evident in zooplankton DW. 4.5 Measuring primary production, zooplankton and micronekton 843 844 Despite the high variability observed in our primary production in situ measurements, 845 particularly during the cool season, and despite their small number (12 measures out of 41 846 stations for both cruises), we estimated they were reasonably representative of the entire 847 cruise. In situ measures of primary production during both Nectalis cruises were similar to previous studies in the region (300 – 1000 mgC m⁻² day⁻¹ in September 2004 at 28°S and 848 849 \sim 155°E-162°E according to Young et al. (2011). These in situ measurements were also in the range of values estimated by satellite for the season, providing some validation and allowing 850 851 confidence in using the satellite VGPM values for a larger scale assessment of primary 852 production. 853 The biomass estimates of microzooplankton and mesozooplankton (~20-2000 μm) provided 854 by the S-ADCP were significantly correlated to the estimates using acoustics (TAPS) and

nets. Estimates derived from the S-ADCP however, are likely to include other organisms and

856	therefore the S-ADCP provides a proxy which is not only reflective of zooplankton biomass
857	(e.g. Burd and Thomson, 2012; Chereskin and Tarling, 2007). The strong correlation
858	between the 153kHz S-ADCP Sv and the EK60 signal, the frequencies of which detect a range
859	of organisms from zooplankton to micronekton and larger, indicate that estimates provided
860	by the S-ADCP also include estimates of micronekton biomass. Although the results did not
861	allow us to define precisely which size-range of organisms was detected by the S-ADCP, and
862	despite the difficulties in calibrating the S-ADCP onboard vessels (Gostiaux and van Haren,
863	2010), our results suggest that this instrument may provide a useful proxy of relative
864	biomass of zooplankton and micronekton confirming its use as a functional tool for this
865	purpose (Brierley et al., 1998; Flagg and Smith, 1989; Heywood et al., 1991; Lee et al., 2008;
866	Radenac et al., 2010). Given that the S-ADCP has been used routinely for more than two
867	decades to sample ocean currents, the data provided by these instruments could potentially
868	be useful for mapping zooplankton/micronekton biomass distributions.
869	Correlations between net and acoustic (TAPS) estimates of zooplankton biomass observed
870	in this study have also previously been observed (e.g. Lebourges-Dhaussy et al., 2014;
871	2009a). Preliminary comparisons between the target micronekton trawls and the EK60
872	acoustic signal however, were not correlated. These data need to be further explored. Low
873	but significant correlations between micronekton net sampling and acoustic EK60 Sv have
874	previously been found for micronekton south of New Caledonia using standardized oblique
875	tows from 600 m to the surface at night (Young et al., 2011). Net sampling at targeted depth
876	and selectivity/catchability/avoidance biases are some of the classic issues that may explain
877	the often low correlation between micronekton net sampling and acoustic estimates (Kloser
878	et al., 2009; Koslow et al., 1997).
879	Zooplankton biomass estimates provided by the TAPS were observed to be more than one
880	order of magnitude higher than the biomass estimates provided by net sampling. These
881	results are at odds with previous comparisons in other regions (e.g. Lebourges-Dhaussy et
882	al., 2014, 2009) and at present we do not have a clear explanation for this discrepancy.
883	Several hypotheses can be proposed which might explain such a disagreement. The TAPS
884	detects organisms in the 50-3000 µm size range, while the nets used during the cruises only

collected organisms larger than 200 $\mu m.$ The smaller sized organisms detected by the TAPS
but not collected by the net may lead to smaller biomass estimates from net samples.
Exploration of the data suggested that differences in the size ranges sampled by each
method cannot explain the large difference in the two estimates. A plausible explanation
could be the inadequacy of the parameterization of the model used in the inversion
algorithm to calculate biomass from the TAPS signal. This would induce an overestimation of $% \left\{ 1\right\} =\left\{ 1\right\}$
the biovolume by the TAPS if, for example, the density contrast between organisms and the
water is underestimated. Further exploration of both the data and sampling of the water
column by the TAPS is required. Moreover, vertical net sampling of the water column may
be insufficient for representatively sample the oligotrophic waters around New Caledonia as
the quantity of zooplankton collected was very low. Additional cruises are needed in which
alternative sampling methods such as oblique tows which filter larger quantities of water
and collect larger quantities of samples can be trialed and investigated.
Biomass estimates of micronekton provided by net sampling and those provided by the
SEAPODYM ecosystem model were observed to be in the same range. Because the net
sampled at specific depths thereby providing estimates which were not representative of
the whole water column, and because of the relative simplicity of the way in which
micronekton are estimated in SEAPODYM (Lehodey et al., 2010) this was not necessarily
expected. Additionally, micronekton biomass estimates provided by SEAPODYM were
significantly correlated to the 38 kHz EK60 acoustic signal. Again this was not expected as
the ocean currents used to force SEAPODYM and determine micronekton spatial
distribution in the sub-model come from a model reanalysis and not observations. Hence
good space/time coherence between observed and modeled mesoscale structures at the
time of the observations would not necessarily be expected. Given that the SEAPODYM
simulation is simply eddy permitting, these results are encouraging and indicate that the
influence of mesoscale features on micronekton biomass is adequately captured by
SEAPODYM in this region.

913	4.6 A broader view of the south-west Pacific
914	The correlations observed between observations and both VGPM satellite-derived primary
915	production and SEAPODYM estimates of micronekton biomass provide some confidence in
916	using these products in describing the ocean dynamics of the broader south-west Pacific
917	Ocean (~15°S-35°S and 150°E-175°E) encompassing the Coral Sea.
918	Looking more broadly than the region in which the cruises were conducted, horizontal
919	advections also shaped the horizontal structure of primary production in the. (Figure 9).
920	Using VGPM satellite-derived primary production data and SEAPODYM micronekton
921	estimates, higher primary production and micronekton values were observed south of 23°S
922	during the cool season and south of 31°S during the hot season than areas further north.
923	Across the overall south west Pacific Ocean region, primary production was much stronger
924	(by approximately a factor 2 to 3) during the cool season compared to the hot season as the
925	SPCZ regime weakened and stronger winds promoted replenishment of surface nutrients
926	(Figure 9). A similar seasonal signal was observed in micronekton biomass as derived from
927	SEAPODYM however, the contrast between seasons was smaller than that observed in
928	primary production estimates (micronekton was higher during the cool season by a factor
929	<2).
930	Micronekton biomass south of 20°S as provided by SEAPODYM was globally organized in
931	very patchy structures and the primary production maxima provided via remote sensing did
932	not necessarily match the micronekton maxima. One good example can be found in the
933	"downwelling" anticylonic eddy in the EAC at 32°S-155°E, where primary production was
934	organized in a strong band around the eddy whereas micronekton biomass was organized
935	along a filament at the edge of the eddy. Estimates of micronekton biomass provided by
936	SEAPODYM indicated that the southern region was richer in biomass, but was also much
937	more variable than the northern region (Figure 9).
938	Presence of patchy structures and decoupling between different trophic levels raises
939	uncertainty associated with using snapshot surveys to understand the coherence of an

ecosystem in turbulent regions. Additional observations in the region will be needed to confirm the nature of the ecosystem organization at (sub) mesoscales.

5 Conclusions and perspectives

By collecting new data extending from the ocean dynamics to micronekton in the top
600 m, the two Nectalis cruises conducted in the south-west Pacific Ocean in austral cool
and hot season of 2011 have provided a better understanding of the pelagic offshore
ecosystem of this oligotrophic region. Multiple methods were used to measure zooplankton
and micronekton (S-ADCP, TAPS, zooplankton net, SIMRAD EK60, micronekton net).
Correlations were found between methods, however, net biomass estimates and acoustic-
derived estimates did not compare very well. On the other hand, estimates of micronekton $% \left\{ 1,2,,n\right\}$
provided from net sampling and SEAPODYM were in the same range. The S-ADCP
reproduced adequately the trends observed in micronekton and zooplankton, but was
unable to distinguish zooplankton from micronekton and absolute biomasses could
therefore not be calculated. Calibration of the different methods used to estimate
zooplankton and micronekton will require additional and more specifically designed studies
Based on large existing S-ADCP datasets, the demonstrated relation between the S-ADCP
signal and the zooplankton/micronekton biomass estimates provides the opportunity to
estimate relative zooplankton/micronekton biomasses on much larger scales than those
available from dedicated instruments such as EK60 or TAPS. Such effort will be undertaken
in the New Caledonia region using the available S-ADCP database spanning the past 20
years. In line with this work, we believe that the development of on-board calibration
methods for the S-ADCP similar to those for echosounders (e.g. EK60) would be of great
interest, particularly in providing absolute measures of abundance. Models such as
SEAPODYM would benefit from absolute biomasses to better calibrate energy transfer
parameterizations.
Based on our limited dataset and the resolution of our data, we could not examine the
systematic effects of submesoscale phenomenon such as eddies and fronts on ocean

biochemistry and planktonic/nektonic communities structures during the Nectalis cruises.
Data collected however, did suggest that horizontal advection was dominant over eddy
pumping. Our study highlights the difficulty of understanding the impact of eddies in
oligotrophic conditions without a full three dimensional dataset. We were also unable to
explore the role that spatial variability might have at the submesoscale (frontal) level, a
scale at which ecosystems have been shown to organize in some cases (e.g. Lebourges-
Dhaussy et al., 2014; Lévy et al., 2012; Tew Kai et al., 2009). This remains an open question
of wide scientific interest. Further, two cruises in two seasons are not sufficient to fully
describe the role of seasonality on the ecosystem. Additional <i>in situ</i> measurements will be
required to further understand the magnitude of the spatial distribution and seasonal cycle
of zooplankton/micronekton biomass in the region, as planned in the coming years within
the framework of the Nectalis program.
The synoptic Nectalis cruises and the SEAPODYM model at the regional scale indicated that
the micronekton structure south of 20°S was remarkably patchy during both seasons in
relation to the mesoscale dynamics of the region. This patchiness raises the question of how
to best sample the region with dedicated cruises. At present, we have chosen to broadly
sample the New Caledonian EEZ. We believe that given the large uncertainty in
understanding of the ecosystem organization and species, it is still useful to pursue this
effort and will be carried out in a series of two additional cruises in the coming years. We
also do note that is it extremely difficult to interpret ecosystem signals at the mesoscale
level using transects organized to cover wide spatial areas. We therefore aim to design
dedicated cruises to follow a number of eddies in the region and understand the time
dynamics of such evolving systems.

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1321 Table 1. Summary of the cruise activities.

	Nectalis1	Nectalis2
Number of sampling stations	18	23
Physics: Temperature, salinity, fluorescence, oxygen 0-500 m: CTD sensors Currents: 153 kHz ship borne-ADCP (16-200 m) Sea surface salinity and temperature:	All stations Along the track Along the track	All stations Along the track Along the track
thermosalinograph		
Nutrients: Nutrients, NOx, SRP, 8 depths: 180 m, 150 m, 130 m, 90 m, 70 m, 40 m, 3 m; CTD water sampling	All stations	All stations
Phytoplankton & pigments: Total Chlorophyll, 8 depths, fluorometry Size fractionated chlorophyll (<3 μm, 3-10 μm, >10 μm): 8 depths, fluorometry Pigments: Phycoerythrin, 4 depths in the euphotic zone depending of the stratification (3m, between 20-40m, DCM, below DCM) spectrofluorometry.	All stations Stations 2, 4, 6, 8, 10, 12, 14, 16 Every other station	All stations Stations 1, 2, 6, 9, 11, 14, 17, 20, 23 Every other station
Cell counts: CTD water sampling and flow cytometry	Stations 1, 3, 5, 7, 9 11, 13, 14, 15, 17, 18	Stations 1, 3, 5, 7, 9, 11, 15, 17, 19, 21, 23
Primary production: 3 depths (surface, between 20-40m, DCM), ¹⁴ C tracer technique	Stations 1, 4, 7, 9, 13, 16, 17	Stations 4,8,12,15, 21
Zooplankton:		
Net sampling: Hydrobios 5 layer (0-600 m) mesh > 200 μm	All stations	All stations
Acoustics: 1 frequency 153 kHz S-ADCP (16-200 m) Acoustics: 6 frequency zooplankton profiler TAPS (0-200 m)	Along the track All stations	Along the track All stations
Micronekton:		
Net sampling: micronekton net between 14 and 540 m depth (10 mm codend mesh size)	All stations	All stations
Acoustics: 4 frequency EK60 SIMRAD echosounder (0-600 m)	Along the track	Along the track
Acoustics: 1 frequency 153 kHz S-ADCP (16-200 m)	Along the track	Along the track

Table 2. Mean and standard deviation (SD) of primary production, biomass estimates and acoustic signal of zooplankton and micronekton during the cool season (Nectalis1) and the hot season (Nectalis2). Results of the Mann-Whitney statistical test (for α =5%) comparing Nectalis 1 (N1) and Nectalis2 (N2) and percentage of the number of tests producing this result for datasets with spatial auto-correlation (see section 2.3.1 for detailed explanation). Seasonal difference between micronekton biomass estimates derived from net sampling was not undertaken because different times and depths were sampled during each survey. DW: dry weight; WW: wet weight.

	Necta	Nectalis1		alis2	Seasonal difference (Mann- Whitney) and percentage of tests producing the result
	Mean	SD	Mean	SD	
<i>In situ</i> primary production (mgC m ⁻² d ⁻¹)	352	160	231	133	N1>N2
Satellite derived primary production along the cruise track (mgC m ⁻² d ⁻¹)	301	62	199	55	100% N1>N2
Sv ADCP (dB)	-82.2	3.5	-83.4	2.8	80% N1>N2; 20% No difference
TAPS biovolume (mg m ⁻³)	107.7	37.3	106.7	22.6	No difference
Zooplankton DW. 0-200 m (mg m ⁻³)	3.9	2.6	5.8	2.3	N1 <n2< td=""></n2<>
Zooplankton WW. 0-600 m (mg m ⁻³)	6.3	3.4	5.6	1.8	No difference
Micronekton (net) mg m ⁻³)	3.4	3.0	7.1	6.8	
Sv EK60 0-600 m (dB)	-77.8	2.7	-77.7	2.6	No difference at 100%
Micronekton (SEAPODYM 0-600 m mg m ⁻³)	4.3	1.2	4.3	0.9	No difference at 100%

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Table 3. Correlations between estimates of zooplankton biomass derived from each of the sampling methods deployed during the cruises. Grouped Nectalis1 and 2 Spearman's correlations and p-values between zooplankton dry and wet weight measurements (in mg m⁻³) from net sampling and their acoustic proxies, S-ADCP Sv (in decibels, dB) and TAPS biovolume (in mg m⁻³) for the averaged top 100 m and top 200 m. Statistics involving S-ADCP Sv is performed by calculating log₁₀ of TAPS biovolume, zooplankton DW and WW.

Variables	Spearman's correlation coefficient		p-v	alue
	0-100 m	0-200 m	0-100 m	0-200 m
S-ADCP Sv vs. log ₁₀ (TAPS biovolume)	0.58	0.64	7e-5*	7e-6*
S-ADCP Sv vs. log ₁₀ (zoopl. dry weight from net)	0.53	0.66	6e-4*	1e-5*
S-ADCP Sv vs. $log_{10}(zoopl.$ wet weight from net)	0.44	0.36	5e-3*	0.03*
TAPS biovolume vs. zoopl. dry weight from net	0.52	0.61	9e-4*	6e-5*
TAPS biovolume vs. zoopl. wet weight from net	0.46	0.55	3e-3*	4e-4*

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Table 4. Correlations between estimates of micronekton biomasses. Grouped Nectalis1 and 2 Spearman's correlations and significance between the four frequencies of S-ADCP Sv (dB) and the corresponding EK60 Sv (dB) averaged across 0-200m, and between estimates derived from \log_{10} (SEAPODYM) and the corresponding 38 kHz EK60 Sv (dB) average across 0-350 m. Statistics involving S-ADCP Sv is performed by calculating \log_{10} of TAPS biovolume, zooplankton DW and WW.

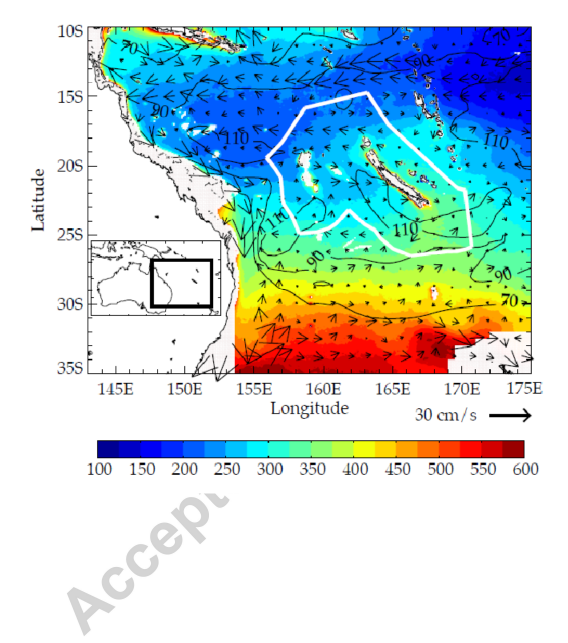
Variables	Range of Spearman's correlation coefficient	Percentage of significant correlations at α =0.05		
S-ADCP Sv vs. 200 kHz EK60 Sv	0.83-0.90	100%		
S-ADCP Sv vs. 120 kHz EK60 Sv	0.85-0.92	100%		
S-ADCP Sv vs. 70 kHz EK60 Sv	0.87-0.96	100%		
S-ADCP Sv vs. 38 kHz EK60 Sv	0.73-0.79	100%		
Log ₁₀ (SEAPODYM) vs. 38 kHz EK60 Sv	0.66-0.80	100%		
Log ₁₀ (SEAPODYM) vs. 38 kHz EK60 Sv 0.66-0.80 100%				

1355	rigures
1356	Figure 1: Mean 1998-2007 primary production estimated from satellite (VGPM) in mgC m ⁻²
1357	d ⁻¹ (shading). Regions of ocean depth shallower than 200 m have been blocked out. Mean
1358	depth of the 1 μM nitrate isopleth (proxy for the nitracline depth) was extracted from CARS
1359	climatology (http://www.marine.csiro.au/~dunn/cars2009/) (Ridgway et al., 2002) in meters
1360	(contour lines) and mean 0-150 meter total geostrophic currents sourced from Kessler and
1361	Cravatte (2013) (vectors). The New Caledonia Exclusive Economic Zone is delineated by the
1362	white line.
1363	Figure 2: Mean surface in situ (left column) and satellite-derived (right column)
1364	oceanographic conditions in the New Caledonian region during the cool season (Nectalis1 -
1365	29 July to 16 August 2011): cruise track and station numbers with those sampled at night in
1366	bold red and those sampled during the day in regular black (top left); thermosalinograph
1367	sea-surface temperature in °C (SST) and salinity (SSS) (middle left); 0-150 m averaged
1368	currents in m $\rm s^{-1}$ (vectors) from the S-ADCP with blue and red arrows indicating eastward
1369	and westward currents respectively (bottom left); surface currents (vectors) from OSCAR
1370	(right column), scale identical to S-ADCP scale; MODIS-VGPM derived depth-integrated net
1371	primary production in mgC m ⁻² d- ¹ (top right); GHRSST satellite sea surface temperature in
1372	°C (top-middle right); sea level anomalies (SLA) referenced to the mean geoid in cm
1373	(bottom-middle right), letters indicate eddies identified in the text; eddy depiction index:
1374	Okubo-Weiß parameter (day ⁻²) (bottom right). The cruise track is plotted in black on the
1375	right column.
1376	Figure 3: Mean surface in situ (left column) and satellite-derived (right column)
1377	oceanographic conditions in the New Caledonian region during the hot season (Nectalis2 -
1378	26 November to 14 December 2011). See Figure 2 caption for details.
1379	Figure 4. Biogeochemical parameters across 0-200 m along cruise tracks during the cool
1380	season (Nectalis1, left panel) and the hot season (Nectalis2, right panel), from CTD sensors
1381	and bottle water analyses. The x-axis labels denote station numbers. From top to bottom:
1382	temperature (°C), salinity, nitrate (NO ₃ μM), phosphate (PO ₄ μM), chlorophyll (mg m ⁻³) and

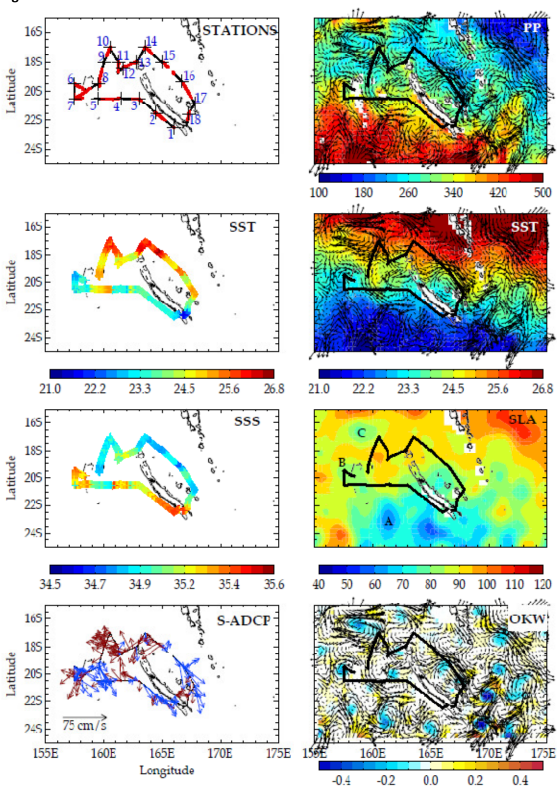
1383	phytoplankton composition. The connected filled circles on the temperature and salinity
1384	panels represent the mixed layer depth, calculated as the depth at which the density equals
1385	the surface density \pm 0.03 kg m $^{-3}$ (de Boyer Montégut et al., 2004). The connected filled
1386	diamonds on the chlorophyll panel represent the depth at which nitrate reaches 1 $\mu\text{M}\textsc{,}$ a
1387	proxy for the nitracline depth. Phytoplankton composition is described as a percentage of
1388	picoplankton (< 3 μ m, black), nanoplankton (3 μ m to 10 μ m, blue) and microplankton
1389	(> 10 μm , red) biomass; orange symbols represent the ratio of $\textit{Prochlorococcus}$ cells to total
1390	picoplankton cells (in % abundance); the dots represent the average value of the top 50 m
1391	and the crosses represent the average value of the 50-130 m layer.
1392	Figure 5: Box plots of the distribution of in situ (In situ) primary production and satellite-
1393	derived (VGPM) primary production recorded at the points where in situ production
1394	measurements were performed (Sat.) and along the cruise track (Sat. full). Estimates are
1395	given for the cool (Nectalis1) and the hot season (Nectalis2). The boxplots denote mean
1396	values and 25% and 75% interquartiles (IQ25 and IQ75 respectively); the whiskers represent
1397	IQ25-1.5x(IQ75-IQ25) and IQ75+1.5x(IQ75-IQ25); dots represent outliers.
1398	Figure 6: Day (plain line) and night (dashed line) 0-600 m mean vertical profiles of
1399	zooplankton wet weight (mg m ⁻³) and mean vertical profiles of 38kHz EK60 scattering
1400	volume (dB) during the cool season (Nectalis1, thick line) and the hot season (Nectalis2, thin
1401	line).
1402	Figure 7: Estimates of zooplankton and micronekton biomass during the day (D) and night
1403	(N) during the cool season (Nectalis1, 1) and the hot season (Nectalis2, 2) using the different
1404	methods employed during the cruises. From left to right: distributions of mean S-ADCP Sv
1405	(dB) across 0-150 m, mean TAPS biovolume (mg m^{-3}) across 0-200 m, mean zooplankton dry
1406	weight (DW, mg $\mathrm{m^{\text{-}3}}$) across 0-200 m, mean zooplankton wet weight (WW, mg $\mathrm{m^{\text{-}3}}$) across 0-
1407	200 m, micronekton wet weight (mg m ⁻³) from cumulated net samplings at discrete depths
1408	between 14 and 540 m, mean 38 kHz EK60 Sv (dB) across 0-350 m, and corresponding
1409	depth-averaged mean (epi- and mesopelagic layers) of micronecton biomass estimates from
1410	the SEAPODYM model. The boxplots denote mean values and 25% and 75% interquartiles
1411	(IQ25 and IQ75 respectively); the whiskers represent IQ25-1.5x(IQ75-IQ25) and

1412	IQ75+1.5x(IQ75-IQ25); dots represent outliers. Note that biomass estimates from
1413	SEAPODYM and EK 60, have been identically averaged over three euphotic depths (~350 m)
1414	and day/time periods (see text for further details).
1415	Figure 8. Spatial distribution of the epi- and mesopelagic micronecton biomass (mg m ⁻³)
1416	estimated from SEAPODYM at the stations and periods of the cruises (top panels) and the
1417	corresponding observed 38 kHz Sv from the EK60 echosounder (bottom) during the cool
1418	season (Nectalis1, left panel) and the hot season (Nectalis2, right panel). The day/night
1419	signal was removed from the data (see text for details). For the sake of clarity the EK 60 Sv
1420	data were arbitrarily re-transformed into a linear scale by computing $10^{-\text{Sv/100}}$, but the unit
1421	by itself has no significance. The EK 60 data have been vertically averaged over the same
1422	depths as the micronekton model incorporated into SEAPODYM (3 euphotic layers ~350 m)
1423	and the data was resampled onto the model ¼° grid resolution.
1424	Figure 9: Satellite primary production (VGPM in mgC m ⁻² day ⁻¹) and euphotic layer currents
1425	from GLORYS (top panel). Averaged micronekton biomass (mg $\mathrm{m}^{\text{-3}}$) estimated by SEAPODYM
1426	and averaged currents from GLORYS across the water column (0 $-$ 1000m) (bottom panel).
1427	Cool season (Nectalis1, left panel), hot season (Nectalis2, right panel).
1428	Cool season (Nectalis1, left panel), hot season (Nectalis2, right panel).

Figure 1

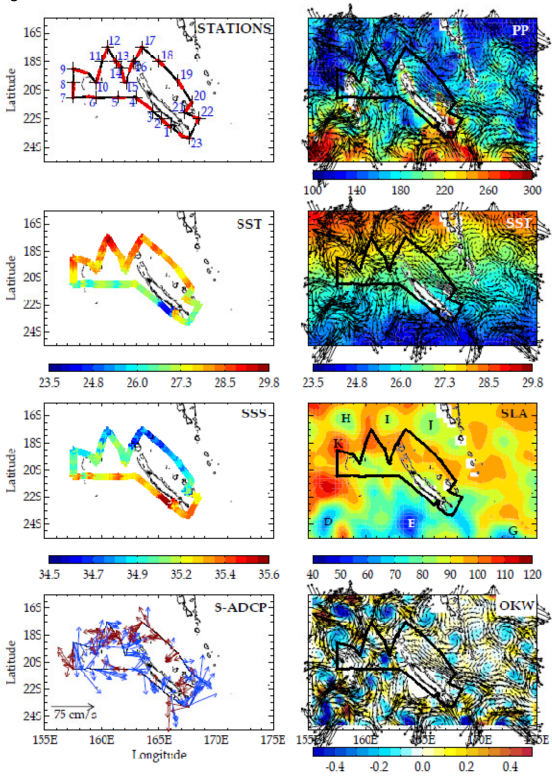


1432 Figure 2

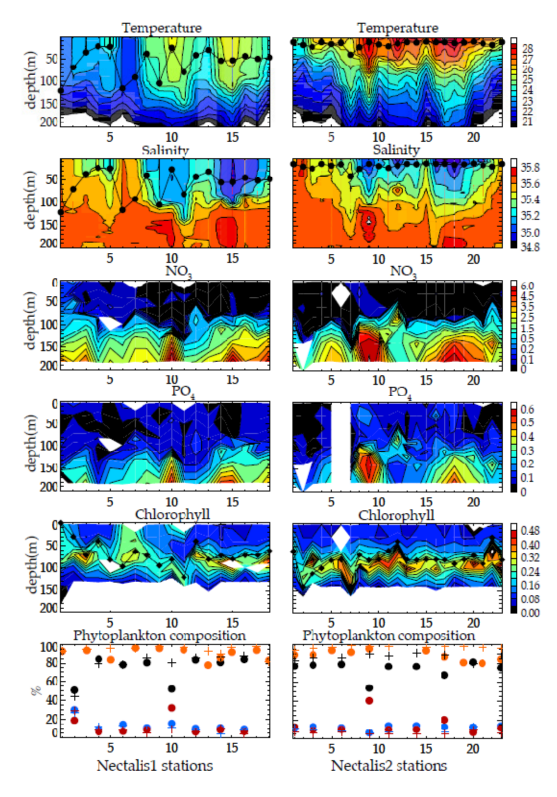


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Figure 3

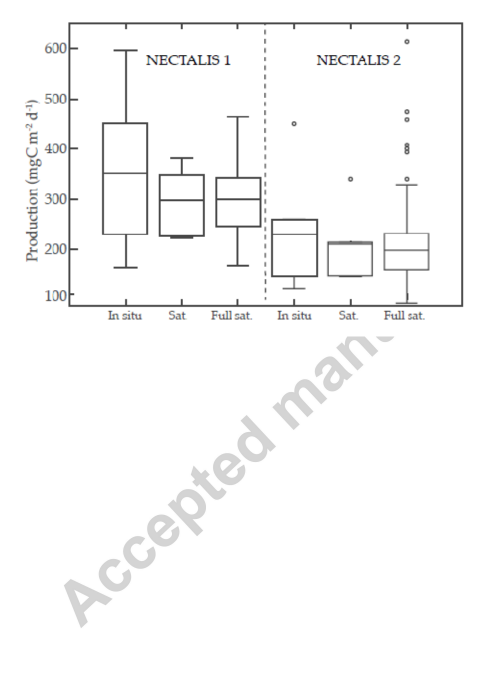


1437 Figure 4



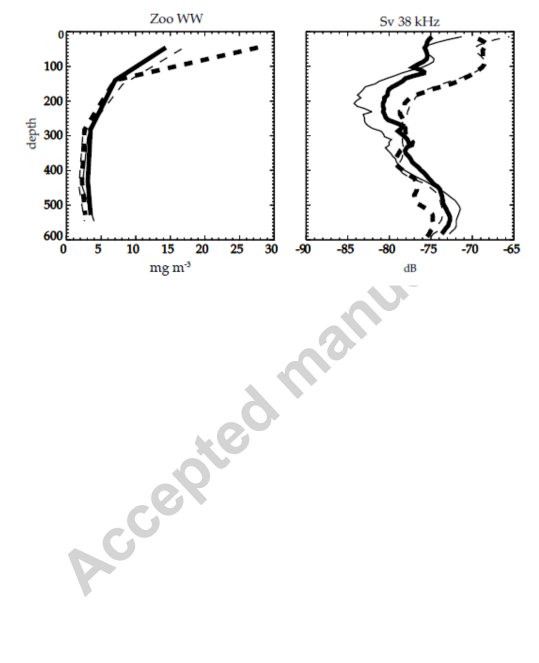
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1440 Figure 5



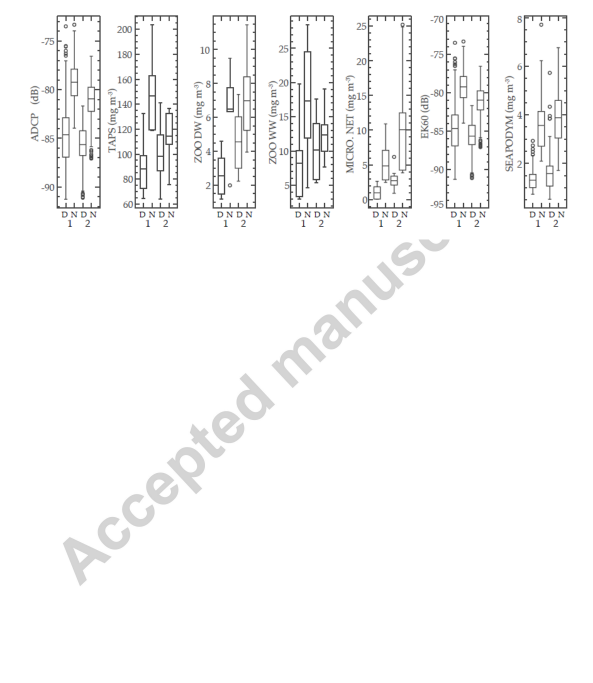
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14431444 Figure 6

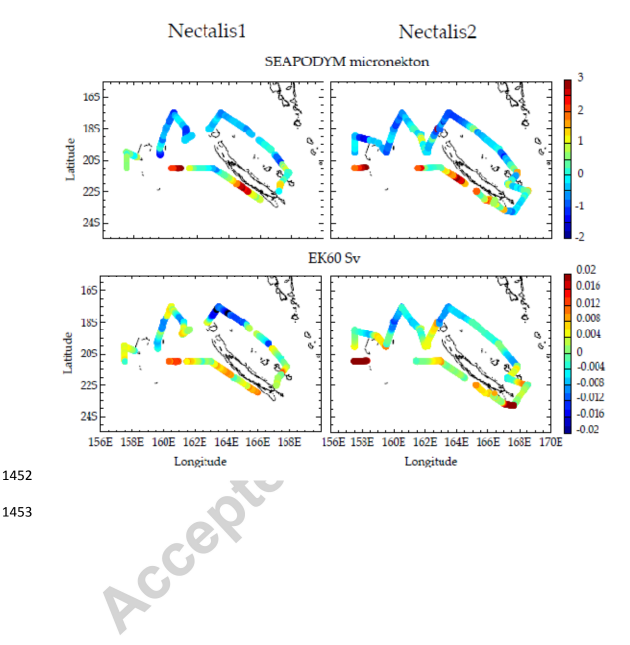


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1448 Figure **7**



1451 Figure 8



1454 Figure 9

