Title

Open science resources for the discovery and analysis of Tara Oceans Data

Authors

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Abstract

The *Tara* Oceans expedition (2009-2013) sampled contrasting ecosystems of the world oceans, collecting environmental data and plankton, from viruses to metazoa, for later analysis using modern sequencing and state-of-the-art imaging technologies. It surveyed 210 ecosystems in 20 biogeographic provinces, collecting over 35000 samples of seawater and plankton. The interpretation of such an extensive collection of samples in their ecological context requires means to explore, assess and access raw and validated data sets. To address this challenge, the Tara Oceans Consortium offers open science resources, including the use of open access archives for nucleotides (ENA) and for environmental, biogeochemical, taxonomic and morphological data (PANGAEA), and the development of online discovery tools and collaborative annotation tools for sequences and images. Here, we present an overview of *Tara* Oceans Data, and we provide detailed registries (data sets) of all campaigns (from port-to-port), stations and sampling events.

Background & Summary

Over many centuries, global expeditions have led to major scientific breakthroughs, notably with the early voyages of the H.M.S. Beagle (1831-1836) and the H.M.S. Challenger (1872-1876). Ocean exploration now provides promising first steps towards understanding the role of the ocean in global biogeochemical cycles and the impact of global climate change on ocean processes and marine biodiversity. Recently, the *Sorcerer II* expeditions (2003-2010) (Ref. 1) and the Malaspina expedition (2010-2011) (Ref. 2) carried out global surveys of prokaryotic metagenomes from the ocean’s surface and bathypelagic layer (>1000 m), respectively. The *Tara* Oceans Expedition (2009-2013) complemented these surveys by collecting a wide variety of planktonic organisms (from viruses to fish larvae) from the ocean’s surface (0-200 m) and mesopelagic zone (200-1000m) at a global scale. Overall, *Tara* Oceans surveyed 210 ecosystems in 20 biogeographic provinces, collecting over 35000 samples of seawater and plankton. Organising such a knowledge base is essential to safeguard, discover and share *Tara* Oceans data. To address this challenge, *Tara* Oceans offers open science resources, including the use of open access data archives and the development of online tools for the collaborative annotation of sequences and images, and the discovery of *Tara* Oceans data.

*Tara* Oceans adopts the principle of open access and early release of raw and validated data sets. In the case of molecular data, raw short sequence reads are archived at the European Bioinformatics Institute short read archive (http://www.ebi.ac.uk/ena/) and made available immediately after manual curation of metadata. More advanced data (assemblies, annotations, etc.) will be released immediately after validation and before publication, and other versions will be released when available. In the case of environmental, biogeochemical, taxonomic and morphological measurements, data are published at PANGAEA, Data Publisher for Earth and Environmental Science (http://www.pangaea.de) and made available immediately after manual curation of metadata.

By combining modern sequencing and state-of-the-art imaging technologies, *Tara* Oceans is at the cutting edge of marine science (3). The amount of data generated by these technologies is unprecedented in the field of plankton ecology and requires adapted storage infrastructures and collaborative platforms to carry out manual and automated annotation of sequences and high throughput images. These open science resources are currently being developed by *Tara* Oceans.

A first series of publications has demonstrated the potential of *Tara* Oceans data to study the ecology of plankton and the structural and functional diversity of viruses, prokaryotes and eukaryotes in the global ocean (4-11). These publications are based on a fraction of the samples analysed so far and thus represent only the tip of the iceberg. The exploration of
Tara Oceans data by the scientific community will undoubtedly lead to new hypotheses and emerging concepts in domains unforeseen by the Tara Oceans Consortium. The current discovery portal of Tara Oceans offers a simple map interface that links each sampling location to available environmental and molecular data (http://www.taraoceans-dataportal.org/). It will however evolve to offer advanced search functionalities based on geospatial, methodological, environmental, morphological, taxonomic, phylogenetic and ecological criteria.

Here, we present an overview of the sampling strategy and size-fractionation approach of the Tara Oceans Expedition (Methods Section) and we explain the rationale behind the choice of sampling devices (Technical Validation Section). Most importantly, we provide registries (data sets) describing all campaigns (from port-to-port), stations and sampling events (Data Records Section). These registries contain geospatial, temporal and methodological information that will be essential for researchers to explore and assess the quality of Tara Oceans data. Environmental data sets are already available openly, in whole or in part, and additional data sets will be progressively released to the community. We intend to submit additional publications describing specific data types (e.g., Data Citations 1-5) in more detail, further extending the value of this resource as the data becomes available.

Methods

As a research infrastructure, the Tara Oceans Expedition mobilised over 100 scientists to sample the world oceans on board a 36 m long schooner (SV Tara) refitted to operate state-of-the-art oceanographic equipment (Figure 1). On board the schooner, the team was consistently composed of five sailors and six scientists, including one chief scientist, two oceanography engineers in charge of deck operations, instrument maintenance and data management, two biology engineers preparing and preserving samples for later morphological and genetic analyses, and one optics engineer in charge of imaging live samples on board. A winch equipped with 2400 m of cable was installed to deploy sampling devices from the stern of the ship, and an industrial peristaltic pump was installed on starboard to sample large volumes of water from various depths down to 60 m. Peristaltic and vacuum filtration systems used to concentrate plankton on membranes of various pore sizes were setup in a laboratory container (wet lab) located outside on port side. Flow-through instruments connected to the continuous surface sampling system were installed in the fore peak and in a laboratory (dry lab) inside the schooner at the centre of the ship on port side.

The sampling strategy and methodology of the Tara Oceans Expedition is presented in six sub-sections. The first four describe why and how the environmental context was determined [1] at the mesoscale using remote sensing and meteorological data; [2] from sensors mounted on the continuous surface water sampling system; [3] from sensors mounted on the vertical profile sampling system; and [4] from discrete water samplers (Niskin bottles) mounted on the vertical profile sampling system. The last two Sub-Sections describe how [5] environmental features were selected and sampled; and how [6] plankton were collected for imaging and genetic analyses. These methods were also described briefly in (3).

[1] Atmospheric and oceanographic context at the mesoscale

The regular sampling programme was designed to study a variety of marine ecosystems and to target well-defined meso- to large-scale features such as gyres, eddies, currents, frontal zones, upwellings, hot spots of biodiversity, low pH or low oxygen concentrations. A total of 210 stations were characterised at the mesoscale to provide richer environmental context for the morphological and genomic study of plankton (Figure 2). In order to identify these
features before sampling but also to assess *a posteriori* if sampling events carried out during
a station were taken within a relatively homogeneous environment, the atmospheric and
oceanographic context were determined at the mesoscale, using climatologies, remote
sensing products and arrays of Argo profiling floats. Meteorological forecast services,
satellite observations (Chlorophyll *a*, sea surface temperature (SST) and altimetry) and real-
time ocean model outputs (Mercator Ocean) were also used on a daily basis to revise
sampling positions with respect to the selected oceanographic features.

Mapped altimetry from AVISO (Archiving Validation and Interpretation of Satellite Data in
Oceanography), mapped operational SST (OSTIA), and satellite ocean colour (ACR-ST
GlobColour service) were used to describe the spatial and temporal variability of key
environmental parameters at each sampling station. In addition, Temperature-Salinity
profiles available around sampling stations were compiled from the Argo autonomous
network array. Finally, a [BATOS] meteorological station mounted on-board *Tara*
continuously measured wind speed and direction, and air temperature, pressure and
humidity, which helped determine the variability of atmospheric conditions and vertical
mixing of surface waters.

In addition to the regular sampling programme, topical experiments were designed to study
ocean processes that operate at spatial and/or temporal scales larger or smaller than the
mesoscale (Figure 3). Topics included, for example, diurnal processes, storm-induced
perturbation of community structure and functions, latitudinal diversity gradients, oxygen
minimum zone (6), island effects on iron fertilization, and longitudinal transport by Agulhas
rings across the South Atlantic Ocean (11). For topical experiments, oceanographic context
was sometimes enriched by using automated underwater vehicles (e.g., gliders (12), ProvBio
(13)), surface-tethered Argo drifters, lowered ADCP mounted on the rosette, basin scale
eddy-field simulations and climatologies, and state-of-the-art physical models of global
ocean circulation with biogeochemistry and genome-informed models of microbial processes
(14). The specific sampling strategy of each topical experiment is available in the respective
cruise summary reports (see *Data Records Section*). *Tara* Oceans data corresponding to
methods described in this section are in part already open to the public at PANGAEA (Data
Citation 1).

[2] Properties of seawater and particulate matter from physical, optical and imaging
sensors mounted on the continuous surface water sampling system

Continuous measurements of surface water physical, chemical and biological properties
serve the dual purpose of a) assessing the boundaries and the homogeneity/heterogeneity
of an ecosystem studied during a station, and b) assessing the connectivity between stations.
Underway measurements were often used to fine tune the location of sampling stations that
were initially selected based on satellite images.

The in-line, Continuous Surface Sampling System [CSSS] installed on SV *Tara* (15,000 miles
long track) comprised a SeaBird [TSG] temperature and conductivity sensor, a WETLabs [AC-
S] spectrophotometer, a WETLabs chlorophyll [Fluorometer], and a Fast Repetition Rate
Fluorometer [FRRF] to assess photosynthetic efficiency. All data were recorded
simultaneously and archived daily in a single file, including navigation, date/time and GPS
position. Water was pumped at the front of the vessel from ~2m depth, then de-bubbled and
circulated through the [AC-S], [TSG], [Fluorometer], and [FRRF]. An automated switching
system provided periodic 0.2μm filtered samples to the [AC-S], such that its particulate
optical properties were not affected by instrument drift (15). Systems maintenance
(instrument cleaning, flushing) was done approximately once a week and in port between
successive campaigns. In the Arctic Ocean and Arctic Seas (2013 campaigns), additional
sensors for pH, PCO₂, optical backscattering (3 wavelengths), fluorescence emission [ALFA]
and surface Photosynthetically Active Radiation [PAR] were added to the in-line system. A
FlowCytobot also recorded images of microplankton every 20 minutes. Using daily discrete measurements of CDOM absorption with an [UltraPath] system, we calibrated the [AC-S] to also provide hourly CDOM absorption (besides particulate absorption and attenuation). Data were processed, quality-controlled, and are consistent with remote sensing. A total of 60 tracks of continuous measurements corresponding to methods described in this section are in large part already open to the public at PANGAEA (Data Citation 2-3).

Properties of seawater and particulate & dissolved matter from physical, optical and imaging sensors mounted on the vertical profile sampling system

Repeated deployments of a Rosette Vertical Sampling System [RVSS] during day and night also served the dual purpose of a) assessing the boundaries and the homogeneity/heterogeneity of mesoscale features during a station, and b) assessing the connectivity between stations. These deployments were essential to locate features that have a vertical component and have a signature below the surface, such as eddies, upwellings, fronts, deep chlorophyll maxima, and oxygen minimum zones. The [RVSS] was specifically designed with various sensors, comprising 2 pairs of conductivity and temperature sensors (Sea-Bird), chlorophyll and CDOM fluorometers (WETLabs), a 25 cm transmitter for particles 0.5-20 μm (WETLabs), a one-wavelength backscatter meter for particles 0.5-20 μm (WETLabs), and a Underwater Vision Profiler (16) for particles >100 μm and zooplankton >600 μm (Hydroptic). A sbe43 oxygen sensor (Sea-Bird) and an In Situ Ultraviolet Spectrophotometer (ISUS) nitrate sensor (SATLANTIC) were also mounted on the Rosette. In the Arctic Ocean and Arctic Seas (2013 campaigns), a second sbe43 oxygen sensor (Sea-Bird) and a four frequency acoustic profiler (Aquascat) were added. Each component was powered on specific Li-Ion batteries and CTD data were self-recorded at 24Hz. All sensors were calibrated in factory before, during and after the four year programme. Oxygen data were validated using climatologies. Nitrate and Fluorescence data were adjusted with discrete measurements from Niskin bottles mounted on the Rosette, and dark calibrations of the optical sensors were performed monthly on-board. A total of 837 vertical profiles were made during the Expedition. Additional stand-alone Sea-Bird components [sbe19] and [sbe9S] were exceptionally mounted directly on the oceanographic cable during harsh sea conditions, when the deployment of the rosette was not safe. In addition, apparent optical properties of sea water were measured using a surface tethered [RADIOMETRE-TSRB] in 2009-2012 and a profiling [RADIOMETRE-COPS] in 2013. Finally, a total of 101 deployments of the Secchi disk [SECCHI] provide a valuable, world-wide contribution to historical records of this very simple and yet fundamental sampling device. Tara Oceans data corresponding to methods described in this section are already open to the public at PANGAEA (Data Citation 4).

Properties of seawater and particulate & dissolved matter from discrete water samples

In addition to sensors mounted on the Rosette Vertical Sampling System [RVSS], seawater was collected using Niskin bottles [NISKIN] (6 x 8-L Niskins and 4 x 12-L Niskins) in order to further characterise environmental conditions in the ecosystem under study. Measurements include pigment concentrations from HPLC analysis (10 depths per vertical profile; 25 pigments per depth), the carbonate system (Surface and 400m; pH, CO₂, pCO₂, fCO₂, HCO₃⁻, CO₃⁻, Total alkalinity, Total carbon, OmegaAragonite, OmegaCalcite, and quality Flag), nutrients (10 depths per vertical profile; NO₃, PO₄, NO₂/NO₃, Si, quality Flags), DOC, CDOM, and dissolved oxygen isotopes. More than 200 vertical profiles of these properties were made across the world ocean. DOC, CDOM and dissolved oxygen isotopes are available only for the Arctic Ocean and Arctic Seas (2013 campaigns). Tara Oceans data corresponding to methods described in this section are already open to the public at PANGAEA (Data Citation 5).

Environmental features and sampling stations
During the *Tara* Oceans Expedition (2009-2013), plankton were sampled from 5-10-m thick layers in the water column, corresponding to specific environmental features that were characterised on-board from sensor measurements. Environmental features are defined by controlled vocabularies in the environmental ontology (EnvO; http://environmentontology.org/) (17).

The *surface water layer* (ENVO:00002042), sometimes labelled in the literature and databases as “surface”, “SRF”, “SUR”, “SURF” or “S”, was simply defined as a layer between 3 and 7 m below the sea surface. The *deep chlorophyll maximum layer* (ENVO:01000326), often labelled in the literature and databases as “DCM” or “D”, was determined from the chlorophyll fluorometer (WETLabs optical sensors) mounted on the Rosette Vertical Sampling System [RVSS]. The presence of a DCM may indicate a maximum in the abundance of plankton bearing chlorophyll pigments, or it may result from the higher chlorophyll content of plankton living in a darker environment (18). This can be assessed *a posteriori* using water samples analysed for pigments by HPLC methods and from plankton counts. The *mesopelagic zone* (ENVO:00000213), also labelled in the literature and databases as “MESO” or “M”, corresponds to the layer between 200 and 1000 m depths. The sampling depth within the *mesopelagic zone* was selected based on vertical profiles of temperature, salinity, fluorescence, nutrients, oxygen, and particulate matter. The selected depth varied from station to station, targeting for example a nutrient line, a minimum concentration of oxygen, a maximum concentration of particulate matter, or a fixed depth of ca. 400 m when no particular feature could be identified. Other environmental features of special scientific interest include the *oxygen minimum zone* (ENVO:01000065), often labelled in the literature and data sets as “OMZ” or “O”, and the *epipelagic mixing layer* (ENVO:01000061), also labelled in the literature and data sets as “ML”, “MIX” or “X”.

A complete sampling station consisted of collecting plankton from three distinct environmental features, typically the *surface water layer*, *deep chlorophyll maximum layer*, and *mesopelagic zone* (Figure 4). Such a station lasted typically 24-48 hours and special care was taken to reposition SV *Tara* in order to remain within a radius of 10 km and sample a homogeneous ecosystem as much as possible (see previous two sub-sections). The sequence of sampling deployments varied but generally followed the order illustrated in Figure 4, i.e. *surface water layer* and *mesopelagic zone* during daytime on the first day, night sampling over fixed depths, and *deep chlorophyll maximum layer* during daytime on the second day. Sampling devices consisted essentially of a High Volume Peristaltic pump [HVP-PUMP], a Rosette Vertical Sampling System [RVSS] equipped with sensors and [NISKIN] bottles, instrumented plankton nets [NET-TYPE-MESH], a [SECCHI] disc, and a [RADIOMETRE] (Table 1).

Plankton were sampled from a total of 210 stations, of which: 51 stations did not target a specific environmental feature and conducted classical vertical profiles of physical and optical sensors and depth integrated net tows; 57 stations sampled only the *surface water layer*; 62 stations sampled the *surface water layer* and a second depth-specific feature; and 40 stations sampled the *surface water layer*, the *deep chlorophyll maximum layer* and a third depth-specific feature. *Tara* Oceans data corresponding to methods described in this section are already open to the public at PANGAEA (Data Citations 6-8) and are described in the *Data Records Section* of the present paper.

Marine plankton

Plankton sampled during the *Tara* Oceans Expedition cover six orders of magnitude in size ($10^{-2}$-$10^{7}$ μm) and correspond to viruses, giant viruses (giruses), prokaryotes (bacteria and archaea), unicellular eukaryotes (protists), and multicellular eukaryotes (metazoans). These five groups form the bulk of biomass throughout the oceans and drive the global biogeochemical cycles that regulate the Earth system (19) (20) (21). Ocean viruses play an
important role in plankton ecology by inducing mortality, horizontal gene transfer, and modulating microbial metabolism (22). They are thought to target diverse prokaryotic and eukaryotic hosts including microalgae and heterotrophic protists, and to play a role in the evolution of their hosts (23). Small viruses (<0.2 µm) are known to be ubiquitous and the most abundant plankton in seawater, while the larger giant viruses or giruses (0.2-1 µm) were discovered more recently and are increasingly observed in marine samples (24) (25). Prokaryotes are believed to be responsible for 30% of primary production and 95% of community respiration in oceans (26) and are thus a fundamental component of marine food webs and biogeochemical processes. They are often divided and studied as two size fractions: the free-living prokaryotes range in size from 0.22-3 µm, and those that are attached to larger cells, particles, or aggregates are found in the 3-20 µm size-fraction (27). In most cases, they are very difficult to culture. Unicellular eukaryotes, or protists, cover a broad range of cell size (0.8-2000 µm). They are taxonomically very diverse with representatives in all of the 8 super-groups of the eukaryotic tree of life (28), whose roles in marine and Earth systems ecology are largely unexplored. Only the most abundant groups, such as diatoms and dinoflagellates, have been studied extensively in the field and cultured successfully (29). Meso-zooplankton (metazoans; multicellular eukaryotes) range in size from 50 µm to tens of metres in colonial forms, and play a pivotal role in both the transfer of energy to higher trophic levels such as fish and other large predators, and in the vertical export of particulate matter produced at the surface of the ocean (30). Their life history (e.g., metabolism, development, locomotion, reproduction, feeding) and their body-size are important properties affecting these two processes (31).

Various sampling methods were used to capture the diversity of both the dominant and less abundant organisms described above (see Technical Validation Section). These methods effectively separated organisms into 10 size fractions: <5 µm (or <3 µm), 5-20 µm (or 3-20 µm), <20 µm, 20-180 µm and 180-2000 µm for planktonic viruses, prokaryotes and unicellular eukaryotes, and >50 µm, >200 µm, >300 µm, >500 µm and >680 µm for large planktonic unicellular eukaryotes and metazoans. Whenever possible, replicate sampling was performed to assess plankton natural variability and to ensure long-term storage of samples in view of future re-analysis using new technologies, notably in the fields of high throughput imaging and –omics which are evolving extremely rapidly.

Detailed protocols concerning the filtration, preservation and storage of plankton samples will be described in detail as a separate publication. Morphological data will be openly released at PANGAEA (http://www.pangaea.de) and nucleotides data will be openly released as they become available at the European Nucleotide Archive (http://www.ebi.ac.uk/ena/).

[6a] Sampling planktonic viruses, prokaryotes and unicellular eukaryotes

Sampling devices used to collect small size organisms (<20 µm size fractions) include Niskin bottles [NISKIN] mounted on the rosette [RVSS] or occasionally attached individually on the oceanographic cable, a High Volume Peristaltic Pump [HVP-PUMP], and exceptionally a 10-L plastic bucket [BUCKET] (Table 1). Waste waters from Tara were not released automatically at sea and were not purged during stations. The pump [HVP-PUMP] was fixed on deck and connected to a 40 mm diameter flexible tube, lowered from either starboard to sample surface water (3-7m) or from the stern to sample other depths down to 60 m. At the beginning of each sampling activity, the tube was rinsed by letting seawater flow overboard for 10 minutes. A single deployment of the pump [HVP-PUMP] brought back up to 1000 L, whereas 4-6 deployments of the rosette [RVSS] [NISKIN] were necessary to collect 400-600 L of seawater.

The choice of the sampling device was determined by weather conditions and the depth of the targeted environmental features. The surface water layer was systematically sampled using the pump [HVP-PUMP], and exceptionally (at 3 stations) using a 10-Litres plastic bucket
[BUCKET]. The deep chlorophyll maximum layer was sampled preferentially with the pump [HVP-PUMP] or alternatively using multiple deployments of the rosette [RVSS] [NISKIN] when the sampling depth was >60 m. A WETLabs three-optical-sensor [ECOTriplet], measuring pressure, temperature, CDOM and chlorophyll fluorescence, was attached to the intake of the pump tube to monitor sampling depth, temperature and chlorophyll concentrations in real time during pumping. The mesopelagic zone was systematically sampled using multiple deployments of the rosette [RVSS] [NISKIN].

Whole seawater collected by these devices was then pre-filtered successively on nylon conical sieves with a mesh of 200 µm and 20 µm, and additionally 5 µm for protists (Table 1). The filtrate was collected in four to six 100-L polyethylene containers, which were thoroughly washed with 0.1% bleach, rinsed twice with fresh water and rinsed again twice with the filtrate. Depending on protocols, the <5 µm and <20 µm filtrates were further fractionated on-board using one or a combination of membranes with pore sizes 0.1 µm, 0.2 µm, 0.45 µm, 0.7 µm, 0.8 µm, 1.6 µm or 3 µm. The retention efficiency of meshes, pore-membranes and fibre-filters is a constant debate in plankton ecology. Organisms display various shapes, including high length-to-width ratios, some may easily “squeeze” through pores smaller than their “normal” size, and others may form colonies or tend to aggregate into particles much larger than their individual size. We do not intend to assess the efficiency of the various meshes and filters in retaining the different groups of organisms targeted during the Tara Oceans Expedition. We simply picked commonly used size-fractions and accept the fact that organisms or parts of organisms from the different groups may be present in several size-fractions.

The choice of size thresholds used to collect small eukaryotes, and prokaryotes associated with small particles or with eukaryotes varied during the Tara Oceans Expedition, between 3-20 µm and 5-20 µm. Plankton from that size fraction comprise organisms that are often not abundant enough in whole seawater and often too fragile to be collected with plankton nets that are themselves too delicate to be deployed in rough seas. The sampling method was therefore weather-dependent and often a combination of using either the pump [HVP-PUMP] or Niskin bottles [RVSS] [NISKIN] for whole seawater, and using a plankton net with a mesh size of 5 µm [NET-SINGLE-5] when sea conditions allowed.

When the pump [HVP-PUMP] or Niskin bottles [RVSS] [NISKIN] were used to collect the 3-20 µm or 5-20 µm size-fraction, whole seawater was pre-filtered successively on 200 µm and 20 µm, and either collected on 3 µm membrane filters (1 to 5 replicates) or concentrated in a conical, 5 µm mesh nylon sieve [SIEVE-5]. The volume of water passing through each 3 µm pore size membrane was consistently 100 L, whereas the volume passing through the [SIEVE-5] was estimated by recording the pumping rate and the start and end time of pumping. Material collected in the [SIEVE-5] was rinsed into an 8-L polyethylene bottle using seawater pre-filtered on 0.1 µm, up to a final volume of 3 L. The 3-20 µm method was used throughout the Tara Oceans Expedition (2009-2013), whereas the 5-20 µm method was used only in 2009-2012, i.e. not in the Arctic.

When nets [NET-SINGLE-5] were used for the 5-20 µm size-fraction, they were lowered to the selected environmental feature and towed horizontally for 5-15 min at a speed of 0.3 m/s. Net samples from the cod-end were sieved through 20 µm [SIEVE-20] and poured into an 8-L polyethylene bottle, which was thoroughly pre-washed with 0.1% bleach, rinsed twice with fresh water and rinsed again twice with seawater pre-filtered on 0.1 µm. The volume of net sample was adjusted to 3 L with 0.1 µm pre-filtered seawater. After each use, nets, cod-ends, and sieves were rinsed with fresh water and checked for holes. That method was used only in 2009-2012, i.e. not in the Arctic.

Plankton from the 20-180 µm size-fraction were collected using a double plankton net with a 20 µm mesh size [NET-DOBLE-20]. Nets were lowered to the selected environmental
feature and towed horizontally for 5-15 min at a speed of 0.3 m/s. Net samples from the two
cod-ends were sieved through 180 µm [SIEVE-180] and poured into an 8-L polyethylene
bottle, which was thoroughly pre-washed with 0.1% bleach, rinsed twice with fresh water
and rinsed again twice with seawater pre-filtered on 0.1 µm. The volume of net sample was
adjusted to 3 L with 0.1 µm pre-filtered seawater. After each use, nets, cod-ends, and sieves
were rinsed with fresh water and checked for holes.

Plankton from the 180-200 µm size-fraction were collected using a 180 µm Bongo Net [NET-
BONGO-180]. Nets were lowered to the selected environmental feature and towed
horizontally for 5-15 min at a speed of 0.3 m/s. Net samples from the two cod-ends were
sieved through 2000 µm [SIEVE-2000] and poured into an 8-L polyethylene bottle, which was
thoroughly pre-washed with 0.1% bleach, rinsed twice with freshwater and rinsed again
twice with seawater pre-filtered on 0.1 µm. The volume of net sample was adjusted to 3 L
with 0.1 µm pre-filtered seawater. After each use, nets, cod-ends, and sieves were rinsed
with fresh water and checked for holes.

[6b] Sampling large planktonic unicellular eukaryotes and metazoans

Sampling devices used to concentrate and collect the larger and less abundant organisms
(>50 µm size fractions) consisted of plankton nets with mesh sizes ranging from 50 to
680 µm [NET-TYPE-MESH] and metal pan-shaped sieves [SIEVE-MESH] to remove large
organisms as needed (Table 1). All nets were equipped with a flow meter and a temperature-
depth recorder, and their depth was monitored and adjusted during deployments using an
acoustic SCANMAR system. Upon recovery, all nets were rinsed from the outside with
running seawater. Cod-ends and metal sieves used to size-fractionate samples were rinsed
with running seawater pre-filtered successively on 25 µm and 0.1 µm, using Polygard-CR
Cartridge Filters (CR2501006, CRK101006). After each use, nets, cod-ends, and sieves were
rinsed with fresh water and checked for holes.

Organism-selectivity and capture-efficiency of plankton nets depend on the mesh size and
deployment methods, i.e., depth, tow method (oblique/vertical/horizontal), tow speed and
time of day (32). During the Tara Oceans Expedition, plankton nets were deployed during
day and night in order to capture the nycthemeral vertical migrations. Both [NET-WPII-50]
and [NET-WPII-200] were towed vertically or obliquely from a depth of 100 m to the surface,
during night and daytime, at a speed of 0.3-0.5 m/s depending on weather conditions. Both
[NET-BONGO-300] and [NET-REGENT-680] were towed obliquely from a depth of 500 m to
the surface, during night and daytime, at a speed of 0.5 m/s depending on weather
conditions. Net samples were preserved on-board with buffered formaldehyde, ethanol or
RNA-Later for later morphological and/or molecular analyses.

Where time and weather allowed, a multiple opening-closing net equipped with 5 nets of
300 µm mesh size [NET-MULTI-300] was deployed preferentially at night or during daytime
to study the vertical distribution of zooplankton. Nets opened and closed at selected depths
between 1000 m and the surface, according to water column features identified from
vertical profiles of temperature, salinity, fluorescence, nutrients, oxygen, and particulate
matter. Samples were preserved on-board in buffered formaldehyde. The Underwater Vision
Profiler (UVP) mounted on the rosette was also used to study the vertical distribution of
zooplankton >600 µm during day and night.

In 2011-2013, a neuston net [NET-MANTA-500] was towed at the surface for about 1h at a
speed of 0.7 m/s in order to collect plastic particles and associated organisms. Samples were
preserved in ethanol for later morphological and molecular analyses. Finally, a Continuous
Plankton Recorder (CPR) was deployed between stations in 2013. Samples were preserved in
formaldehyde and sent to the Sir Alister Hardy Foundation for Ocean Science (SAHFOS) for
later morphological and molecular analyses.
Data Records

Tara Oceans developed best practices for the standardisation and interoperability of data generated across environmental, morphological and molecular analyses. This effort contributed to the publication of a set of standards for reporting and serving data in Marine Microbial Biodiversity, Bioinformatics and Biotechnology (M2B3) (33). Here we describe three levels of the M2B3 reporting standard: campaigns, stations and events. For each level, we provide a registry of all campaigns/stations/events, pdf documents describing each campaign/station/event, and universal resource locator (URL) queries to access related nucleotides and environmental data.

Labels

Labels that identify each campaign, station or event are built using a consistent syntax that has the following format:

- **Campaign** labels: TARA_date(yyyyymmddZ), e.g., TARA_20110401Z
- **Station** labels: TARA_station(#)001-210, e.g., TARA_100
- **Event** labels: TARA_datetime(yyyyymmddThhmmZ)_station#(001-210)_EVENT_TYPE, e.g., TARA_20110416T1306Z_100_EVENT_CAST

Registries

The Tara Oceans Expedition (2009-2013) comprised 60 campaigns (from port-to-port), 210 stations and over 3200 sampling events. Three registries listing the campaigns, stations and events are published at PANGAEA, Data Publisher for Earth and Environmental Science:

- **Campaigns** registry (Data Citation 6): [http://doi.pangaea.de/10.1594/PANGAEA.842191](http://doi.pangaea.de/10.1594/PANGAEA.842191)
- **Stations** registry (Data Citation 7): [http://doi.pangaea.de/10.1594/PANGAEA.842237](http://doi.pangaea.de/10.1594/PANGAEA.842237)
- **Events** registry (Data Citation 8): [http://doi.pangaea.de/10.1594/PANGAEA.842227](http://doi.pangaea.de/10.1594/PANGAEA.842227)

The **campaigns** registry provides details about the scientific interest of each campaign, a list of scientists on board, and URLs for the corresponding campaign summary report (pdf), environmental data sets and nucleotides data sets.

The **stations** registry provides details about the geographic context of each station, including mean and maximum bathymetric depth (extracted from the General Bathymetric Chart of the Oceans; GEBCO), minimum distance from the coast, the corresponding marine biomes and biogeographical provinces defined by Longhurst (34), and when applicable information about the corresponding exclusive economic zone and related legal aspects. Additionally for each station, we provide information about the environmental features that were sampled and their depth, and the number of deployments carried out with the different sampling devices listed in Table 1. URLs provide access to the corresponding oceanographic context report (pdf), environmental data sets and nucleotides data sets.

The **events** registry provides details about the sampling date, time, location and methodology of each event. URLs provide access to the corresponding event log sheet (pdf), environmental data sets and nucleotides data sets. Sampling events occurring outside the context of a station were assigned the station label TARA_999. Such events include for example underway measurements of the on-board meteorological station [BATOS] and of the continuous surface sampling system [CSSS], and exceptional deployments of plankton
nets [NET-TYPE-MESH] or of the rosette vertical sampling system [RVSS].

All available metadata information about a single campaign/station/event can be extracted from the respective registries using a web service (description: https://ws.pangaea.de/dds-fdp/). Here we provide an example for each registry:

**Campaign**: specific metadata query: https://ws.pangaea.de/dds-fdp/rest/panquery?datasetDOI=doi.pangaea.de/10.1594/PANGAEA.842191&filterParameter=Campaign&filterValue=TARA_20110401Z

**Station**: specific metadata query: https://ws.pangaea.de/dds-fdp/rest/panquery?datasetDOI=doi.pangaea.de/10.1594/PANGAEA.842237&filterParameter=Station&filterValue=TARA_100

**Event**: specific metadata query: https://ws.pangaea.de/dds-fdp/rest/panquery?datasetDOI=doi.pangaea.de/10.1594/PANGAEA.842227&filterParameter=Event&filterValue=TARA_20110416T1306Z_100_EVENT_CAST

Additionally, the same web service can be used to extract information about all stations from a campaign, all events from a campaign, or all events from a station as shown in these examples:

**Stations-from-a-campaign**: specific metadata query: https://ws.pangaea.de/dds-fdp/rest/panquery?datasetDOI=doi.pangaea.de/10.1594/PANGAEA.842237&filterParameter=Campaign&filterValue=TARA_20110401Z

**Events-from-a-campaign**: specific metadata query: https://ws.pangaea.de/dds-fdp/rest/panquery?datasetDOI=doi.pangaea.de/10.1594/PANGAEA.842227&filterParameter=Campaign&filterValue=TARA_20110401Z

**Events-from-a-station**: specific metadata query: https://ws.pangaea.de/dds-fdp/rest/panquery?datasetDOI=doi.pangaea.de/10.1594/PANGAEA.842227&filterParameter=Station&filterValue=TARA_100

**Reports and log sheets**

Campaign reports were written by the chief scientist and scientific crew in order to document the objectives and main achievements of each campaign, as well as any deviations from the regular sampling programme (e.g., TARA_20110401Z_report.pdf). The station reports were written by Flavian Kokoszka, Rémi Laxenaire and Sabrina Speich to provide background knowledge of the physical oceanography at each station (e.g., TARA_100_oceanographic_context_report.pdf). Finally, event log sheets were filled on board each time a sampling device was deployed, recording the position, date, time, type of device, sampling depths, volume sampled, ID and filename of sensor outputs, operator’s comments, and unique identifiers (barcodes) of samples collected during the event (e.g., TARA_20110415T1312Z_100_EVENT_CAST.pdf). Most of the information found in reports and log sheets was extracted manually, quality checked using controlled vocabularies, and archived in the campaigns, stations and events registries. Nevertheless, these narrative documents remain a valuable and complementary source of information. The three registries contain universal resource locators (URLs) pointing to the reports and log sheets of each campaign/station/event. Reports and log sheets can also be browsed directly in the PANGAEA store:

**Campaign** reports: http://store.pangaea.de/Projects/TARA-OCEANS/Campaign_Reports/

**Station** reports: http://store.pangaea.de/Projects/TARA-OCEANS/Station_Reports/
Event log sheets: http://store.pangaea.de/Projects/TARA-OCEANS/Logsheets_Event/

Up-to-date lists of available nucleotides and environmental data sets

Tara Oceans data will progressively be released openly following their analysis and validation. Up-to-date lists of nucleotides and environmental data sets can be obtained from universal resource locator (URL) queries that are made specific to any campaign/station/event by using labels from the campaigns, stations and events registries.

A list of environmental data sets published at PANGAEA can be obtained by combining the following base URL: http://www.pangaea.de/search?q= with a search term. The URL query is made specific to any Tara Oceans campaign, station or event by adding the corresponding label as the search term, see examples below. Already-built URL queries for each campaign, station or event are provided in the three respective registries (Data Citations 6-8).

Campaign-specific environmental data query:
http://www.pangaea.de/search?q=TARA_20110401Z

Station-specific environmental data query:
http://www.pangaea.de/search?q=TARA_100

Event specific environmental data query:
http://www.pangaea.de/search?q=TARA_20110416T1306Z_100_EVENT_CAST

A list of nucleotides data published at ENA can be obtained by combining the following base URL: http://www.ebi.ac.uk/ena/data/search?query= with a search term. The URL query is made specific to any Tara Oceans campaign, station or event by adding the corresponding label as the search term, see examples below. Already-built URL queries for each campaign, station or event are provided in the three respective registries (Data Citations 6-8).

Campaign specific nucleotides data query:
http://www.ebi.ac.uk/ena/data/search?query=TARA_20110401Z

Station specific nucleotides data query:
http://www.ebi.ac.uk/ena/data/search?query=TARA_100

Event specific nucleotides data query:
http://www.ebi.ac.uk/ena/data/search?query=TARA_20110415T1245Z_100_EVENT_PUMP

Technical Validation

Here we provide a first order validation of the Tara Oceans sampling methodology by compiling published values of plankton cell/body size, natural abundance and richness (Table 2). These are compared to the sampling volume and mesh size of the different sampling methods (Figure 5).

Life history traits such as cell/body size and the natural range of abundance determine the general structure and dynamics of food webs and other ecological networks, across multiple scales of organisation (35)(31). Here we characterise the five groups of plankton by their size and abundance in seawater, using values from the literature (Table 2). The range of these characteristics are summarised for each plankton group using coloured areas on the horizontal plane of Figure 5. As already described for a wide range of organisms (36), the literature shows an inverse relationship between plankton size and abundance in the natural environment, so that small viruses (10^2-10^0 µm) generally form the most abundant group
(10^7-10^{11} ind. L^{-1}), whereas the larger metazoans (10^3-10^5 μm) are generally the least abundant group (10^4-10^5 ind. L^{-1}).

Species richness and evenness are used to estimate species diversity, and should therefore be considered when designing sampling strategies and methodologies for biodiversity studies. Here we characterise the five groups of plankton by their species richness in seawater, using values from the literature (Table 2, Refs. 37-68). From these values we made “back of the envelope” calculations of the volume of seawater required to capture 100%, 75% and 50% of species within each group of plankton (Figure 5; coloured areas on the vertical plane). The effectiveness of our sampling strategy can be assessed by comparing these coloured areas with the sampling volume of the various sampling devices used during the Tara Oceans Expedition (horizontal full lines on the vertical plane).

Based on this assessment, it appears that our sampling strategy would capture <50% of total richness for viruses and small size protists (0.8-5 μm). Accordingly, one would need to filter thousands of litres of seawater in order to capture 75% of total richness for these groups. This is both impractical for most field campaigns and dependent on how one defines the currency of richness for these groups, i.e. the concept of species. In all other groups and size fractions, our sampling strategy appears to capture >75% of species richness, and 100% in the case of large size protists and metazoans. It is important to note that data about plankton richness is very scarce in the literature, so that this assessment is only a first approximation. Tara Oceans data will undoubtedly contribute to fill this knowledge gap and improve the sampling design of future ocean biodiversity surveys.

Usage Notes
The Tara Oceans data policy follows the open science principle of open access and early release of raw and validated data sets. All data presented here (Data Citations 6-8) are published under the Creative Commons Attribution 3.0 Unported (CC-by 3.0) and must therefore be cited when used in scientific papers, posters or presentations. As with most scholarly publications, data citations have authors, a title, a year of publication and a digital object identifier (see data citations in the Reference Section). Furthermore, we kindly ask to include the Tara Oceans Consortium in the acknowledgements. When referring to the Tara Oceans Data or to the sampling strategy and methodology of the Tara Oceans Expedition, please cite the present paper.

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**Author Contributions**

Contributed to writing this paper: Stéphane Pesant, Fabrice Not, Marc Picheral, Stefanie Kandels-Lewis and Noan Le Becot. Contributed to sampling planning during the Tara Oceans Expedition (2009-2013): Gabriel Gorsky, Daniele Iudicone, Eric Karsenti, Stefanie Kandels-
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Competing Financial Interests

The authors declare no competing financial interests.
Figure 1. Sampling devices and working areas on-board SV Tara. Sampling devices and working areas on-board SV Tara are shown from the vessel’s [a] side-view, [b] bird’s-eye-view of the deck, and [c] inside-view. They consist of the [1] Continuous Surface Sampling System [CSSS]; [2] Rosette Vertical Sampling System [RVSS]; [3] wet lab; [4] High Volume Peristaltic pump [HVP-PUMP]; [5] dry lab; [6] oceanography engineers data acquisition and processing area; [7] winch; [8] video imaging area; [9] storage areas at room temperature; [10] storage areas at +4°C and -20°C; [11] MilliQ water system and AC-s system; [12] diving equipment, flowcytobot and ALPHA instruments; and [13] storage boxes. The flow of seawater from the continuous surface sampling system to the dry lab is shown in blue.

Figure 2. Sampling route and stations of the Tara Oceans Expedition. Sampling route of the Tara Oceans Expedition (green track), showing station labels and areas (blue shade) where the annual mean oxygen concentration is <2 ml/L (WOA09), usually corresponding also to high CO₂ concentration and low pH. Detailed information about each station is given in (Data Citation 7).

Figure 3. Sampling route, stations and topical experiments of the Tara Oceans Expedition. Sampling route of the Tara Oceans Expedition (green track), showing stations where plankton were sampled in their environmental context (full red dots) and where only environmental conditions were measured (open red dots). Topical experiments are identified along the sampling route (light blue). Longhurst biogeographical provinces (34) are shown in the background and those sampled during Tara Oceans Expedition are highlighted in blue.

Figure 4. Spatial representation and chronology of sampling events during a 48 hours station. Coloured markers along the route of SV Tara (yellow surface track) correspond to sampling events targeting the surface water layer (red, ), deep chlorophyll maximum layer (green, here at 50 m), and the mesopelagic zone (blue, here at 400 m). At some stations, an Argo drifter (10-m floating anchor and satellite positioning) was used to follow the water mass during sampling (black surface track). White and grey markers correspond to day and night time deployments, respectively, of plankton nets [TYPE-MESH] and rosette [RVSS] casts that covered fixed depth layers of 0-100m, 0-500m or 0-1000m.

Figure 5. Empirical basis for the size-fractionation approach and the choice of sampling devices. The horizontal plane shows the range of body/cell size and natural abundances reported in the literature (Table 2) for viruses (including giant viruses), prokaryotes, protists and metazoans (coloured boxes). The sampling devices used to collect plankton <5 µm in size (i.e. high volume peristaltic pump and rosette with Niskin bottles) and >5 µm in size (i.e. plankton nets) are illustrated as well on the horizontal plane. The vertical plane shows the volume of seawater required to capture 100%, 75% and 50% of species richness reported in the literature (Table 2) for viruses (including giant viruses), prokaryotes, protists and metazoans (shaded boxes). The typical volume of seawater collected by sampling devices are shown in comparison (horizontal thick lines). Also illustrated on the vertical plane: Sieves were used to remove large organisms in the case of plankton nets for protists (5, 20 and 180 µm mesh).
### Table 1. List of sampling devices used during Tara Oceans Expedition (2009-2013)

| Referenced in Methods Sub Section | Targeted Plankton Size-fraction | Sampling volume (Litres) per deployment | Device label | Device description |
|----------------------------------|---------------------------------|----------------------------------------|--------------|--------------------|
| [1] Environmental Parameters only | Environmental Parameters only | [BATOS] | Meteo France Voluntary Observing Ships meteorological station equipped with a PTB220 barometer (Vaisala), a PT100 platinium thermometer (Heraeus Sensor), a HMP35 DE hygrometer (Vaisala) for relative humidity, an ultrasonic wind sensor from GILL, and a hull contact temperature sensor PT100 (SPCK PROSENSOR) affixed on the hull of the ship, below the waterline |
| [2] Environmental Parameters only | Environmental Parameters only | [TSG] | Thermostalinograph (SeaBird), mounted on the [CSSS] |
| [2] Environmental Parameters only | Environmental Parameters only | [CSSS] | Continuous Surface Sampling System (5) |
| [2] Environmental Parameters only | Environmental Parameters only | [AC-S] | Spectrophotometer (WETLabs) mounted on the [CSSS], providing hyperspectral particulate absorption and attenuation |
| [2] Environmental Parameters only | Environmental Parameters only | [ALFA] | A 2-wavelength fluorometer mounted on the [CSSS], providing chlorophyll, phycoerythrin and CDOM fluorescence as well as phytoplankton photosynthetic parameters (Fv/Fm protocol) |
| [2] Environmental Parameters only | Environmental Parameters only | [FlowCytobot] | Imaging cytometer for phytoplankton > 30 µm |
| [2] Environmental Parameters only | Environmental Parameters only | [Fluorometer] | Chlorophyll fluorometer (WETLabs) mounted on the [CSSS] |
| [2] Environmental Parameters only | Environmental Parameters only | [FRRF] | Fast Repetition Rate Fluorometry mounted on the [CSSS], assessing photosynthetic efficiency |
| [2] Environmental Parameters only | Environmental Parameters only | [SeaFET] | Ocean pH sensor (Satlantic), mounted on the [CSSS] |
| [2] Environmental Parameters only | Environmental Parameters only | [UltraPath] | Long pathlength spectrophotometer, measuring CDOM absorption |
| [2] [3] Environmental Parameters only | Environmental Parameters only | [PAR] | Photosynthetically Active Radiation sensor (Biospherical Instruments Inc.) |
| [3] Environmental Parameters only | Environmental Parameters only | [sbe9S] | Standalone Sea-Bird CTD (sbe9S) mounted directly on the oceanographic cable. |
| [3] Environmental Parameters only | Environmental Parameters only | [sbe19] | Standalone Sea-Bird CTD (sbe19) mounted directly on the oceanographic cable. |
| [3] [5] Environmental Parameters only | Environmental Parameters only | [RVSS] | Sensors mounted on a Rosette Vertical Sampling System [RVSS], including Sea-Bird components: conductivity-temperature-depth (CTD, sbe9C) and oxygen sensor (sbe43); WETLabs optical sensors: chlorophyll & CDOM fluorometers, 25 cm transmissiometer for (650 nm), and one-wavelength backscatter meter for particles; |
| Referenced in Methods Sub Section | Targeted Plankton Size-fraction | Sampling volume (Litres) per deployment | Device label | Device description |
|---------------------------------|---------------------------------|-------------------------------------|--------------|------------------|
| [3]                             | Environmental Parameters only   | [RADIOMETRE-TSR8]                  | Tethered Spectrophotometer nitate sensor (ISUS). In 2013, a second oxygen sensor (sbe43) and a four frequency acoustic profiler (Aquascat) were added. |
| [3]                             | Environmental Parameters only   | [RADIOMETRE-COPS]                  | Compact Optical Profiling System (Biospherical Instruments Inc.) |
| [3]                             | Environmental Parameters only   | [SECCHI]                           | Secchi disk, diameter (cm)=20 |
| [4] [6a]                        | whole water                    | $10^3$                             | [NISKIN]     | At a few stations, Niskin bottles were mounted individually on the oceanographic cable and triggered with messengers, each collecting 8 L or 12 L. |
| [4] [6a]                        | whole water                    | $10^2$-$10^3$                      | [NISKIN] [RVSS] | 10 Niskin bottles (6 x 8-L and 4 x 12-L) equipped with silicon and Teflon parts were mounted on a Rosette Vertical Sampling System [RVSS]. Each deployment of the [RVSS] could collect 8-12 L from different depths or a total volume of 96 L from one depth. |
| [6a]                            | whole water                    | $10^4$                             | [BUCKET]     | 10-L plastic bucket attached to a line |
| [6a]                            | whole water                    | $10^2$-$10^3$                      | [HVP-PUMP]   | High Volume Peristaltic Pump (Tech-pump A40) |
| [6a]                            | whole water                    | $10^2$-$10^3$                      | [HVP-PUMP]+ [sbe9S] | High Volume Peristaltic Pump (Tech-pump A40), equipped with a CTD [sbe9S] |
| [6a]                            | whole water                    | $10^2$-$10^3$                      | [HVP-PUMP]+ [ECOTriplet] | High Volume Peristaltic Pump (Tech-pump A40), equipped with an [ECOTriplet], three-optical-sensor (WETLabs), measuring pressure, temperature, CDOM fluorescence and chlorophyll fluorescence |
| [6a]                            | whole water $<5 \mu m$         | $10^2$-$10^3$                      | [SIEVE-200]+ [SIEVE-20]+ [SIEVE-5] | Serial filtration through conical, nylon sieves SIEVE-200: type=nylon, shape=conical, mesh size (µm)=200, diameter (cm)=90 SIEVE-20: type=nylon, shape=conical, mesh size (µm)=20, diameter (cm)=90 SIEVE-5: type=nylon, shape=conical, mesh size (µm)=5, diameter (cm)=60 |
| [6a]                            | whole water $<20 \mu m$        | $10^2$-$10^3$                      | [SIEVE-200]+ [SIEVE-20] | SIEVE-200: type=nylon, shape=conical, mesh size (µm)=200, diameter (cm)=90 SIEVE-20: type=nylon, shape=conical, mesh size (µm)=20, diameter (cm)=90 |
| [6a] [6b]                       | 5-20 µm                        | $10^2$-$10^3$                      | [NET-SINGLE-5]+ [SIEVE-20] | Plankton net, type=single, #nets=1, mesh size (µm)=5, mouth opening (m^2)=0.196349, length (m)=2.5 SIEVE-20: type=metal, shape=pan, mesh size (µm)=20, diameter (cm)=30 |
| Referenced in Methods Sub Section | Targeted Plankton Size-fraction | Sampling volume (Litres) per deployment | Device label | Device description |
|-----------------------------------|----------------------------------|----------------------------------------|--------------|-------------------|
| [6a] [6b] 20-180 µm              | $10^4$-$10^5$                    | [NET-DOUBLE-20]+ [SIEVE-180]           | Plankton net, type=Double, #nets=2, mesh size (µm)=20, mouth opening (m^2)=0.192442, length (m)=2.5 |
| [6a] [6b] 180-2000 µm           | $10^4$-$10^5$                    | [NET-BONGO-180]+ [SIEVE-2000]          | Plankton net, type=Bongo, #nets=2, mesh size (µm)=180, mouth opening (m^2)=0.258319, length (m)=3 |
| [6b] >50 µm                     | $10^4$-$10^5$                    | [NET-WPII-50]                           | Plankton net, type=WPII, mesh size (µm)=50, mouth opening (m^2)=0.248505, length (m)=2.75 |
| [6b] >200 µm                    | $10^4$-$10^5$                    | [NET-WPII-200]                          | Plankton net, type=WPII, #nets=1, mesh size (µm)=200, mouth opening (m^2)=0.248505, length (m)=2.75 |
| [6b] >300 µm                    | $10^5$-$10^6$                    | [NET-BONGO-300]                         | Plankton net, type=Bongo, #nets=2, mesh size (µm)=300, mouth opening (m^2)=0.258319, length (m)=3 |
| [6b] >300 µm                    | $10^6$                           | [NET-MULTI-300]                         | Multiple opening-closing plankton net, type=Multinet (Hydrobios), #nets=5, mesh size (µm)=300, mouth opening (m^2)=n/a, length (m)=2.5 |
| [6b] >680 µm                    | $10^6$                           | [NET-REGENET-680]                       | Plankton net, type=Regent, #nets=1, mesh size (µm)=680, mouth opening (m^2)=0.754296, length (m)=4 |
| [6b] >500 µm                    | $10^4$-$10^5$                    | [NET-MANTA-ST-500]                      | Neuston net deployed in 2011-2012, type=Manta suitcase Trawl (ALGALITA), #nets=1, mesh size (µm)=500, mouth width (m)=0.61, length (m)=2.5 |
| [6b] >500 µm                    | $10^5$-$10^6$                    | [NET-MANTA-OI-500]                      | Neuston net deployed in 2013, type=Manta (OCEAN INSTRUMENTS), #nets=1, mesh size (µm)=500, mouth width (m)=0.70, length (m)=2.5 |
Table 2. Cell/body size, abundance and species richness reported in the literature for the four groups of plankton in coastal and oceanic environments.

| Group of organism | Environment | Cell/body size | Abundance | Species richness |
|-------------------|-------------|----------------|-----------|-----------------|
|                   |             | Dimension: $L^1$ | Dimension: $L^3$ | Dimension: $L^3$ |
|                   |             | Unit: micrometre | Unit: individuals per Litre | Unit: species per Litre |
|                   | Coastal     | Oceanic         | Minimum   | Maximum         | Minimum   | Maximum         | Reference |
| Viruses           | X           | X               | $2 \times 10^{-2}$ | $2 \times 10^{-1}$ | $1 \times 10^9$ | $2 \times 10^{11}$ | (37)      |
| Viruses           | X           | X               | $2 \times 10^{-2}$ | $2 \times 10^{-1}$ | $1 \times 10^{10}$ |                     | (38)      |
| Viruses           | X           |                 | $2 \times 10^{-2}$ | $2 \times 10^{-1}$ | $6 \times 10^7$ | $2,6 \times 10^8$ | (39)      |
| Viruses           | X           |                 | $2 \times 10^{-4}$ | $2 \times 10^{-3}$ | $1,2 \times 10^7$ | $3,5 \times 10^8$ | (39)      |
| Viruses           | X           | X               |                     |                 | $1 \times 10^5$ | $1 \times 10^6$ | (40)      |
| Prokaryotes       | X           |                 |                     | $3 \times 10^9$ |                     | (41)      |
| Prokaryotes       | X           | X               | <                     | $1 \times 10^6$ | $4 \times 10^9$ |                     | (42)      |
| Prokaryotes       | X           |                 | <                     | $2 \times 10^6$ | $1 \times 10^9$ | $5 \times 10^9$ | (43)      |
| Prokaryotes       | X           |                 | <                     | $2 \times 10^6$ | $5 \times 10^8$ |                     | (43)      |
| Prokaryotes       | X           |                 | <                     | $2 \times 10^9$ | $2 \times 10^8$ | $9 \times 10^9$ | (44)      |
| Prokaryotes       | X           |                 | <                     | $2 \times 10^9$ | $2 \times 10^8$ | $6 \times 10^8$ | (44)      |
| Prokaryotes       | X           |                 | <                     | $2 \times 10^9$ | $5 \times 10^8$ | $7,5 \times 10^9$ | (45)      |
| Prokaryotes       | X           |                 | <                     | $2 \times 10^9$ | $2,25 \times 10^8$ | $3 \times 10^8$ | (46)      |
| Prokaryotes       | X           |                 | <                     | $2 \times 10^9$ | $2 \times 10^8$ | $8 \times 10^8$ | (46)      |
| Prokaryotes       | X           |                 |                     | $1 \times 10^9$ |                     | (47)      |
| Prokaryotes       | X           |                 |                     | $1 \times 10^7$ |                     | (48)      |
| Prokaryotes       | X           |                 |                     | $2 \times 10^4$ |                     | (49)      |
| Protists          | X           |                 |                     | $2 \times 10^4$ |                     | (50)      |
| Protists          | X           |                 |                     | $3 \times 10^4$ |                     | (41)      |
| Protists          | X           |                 | $8 \times 10^{-1}$ | $2 \times 10^7$ | $1,83 \times 10^9$ |                     | (51)      |
| Protists          | X           |                 | $6 \times 10^{-1}$ | $1 \times 10^7$ | $1 \times 10^9$ | $4 \times 10^9$ | (52)      |
| Protists          | X           |                 | $2 \times 10^{-1}$ | $2 \times 10^7$ | $1 \times 10^9$ |                     | (53)      |
| Protists          | X           |                 | $8 \times 10^{-1}$ | $2 \times 10^9$ | $5 \times 10^6$ | $3 \times 10^9$ | (54)      |
| Protists   | X    | X    | 2x10^{-1} | 3x10^5 | 1x10^6 | 2x10^7 |   | (55) |
|-----------|------|------|-----------|--------|--------|--------|---|------|
| Protists  | X    |     | 8x10^0    | 1x10^3 | 1x10^7 | 3x10^8 |   | (56) |
| Protists  |      | <    | 2x10^2    | 1.08x10^5 |     |        |   | (57) |
| Protists  | X    | <    | 2x10^2    | 2.6x10^5 |     |        |   | (57) |
| Protists  |      | 1x10^1 | 1x10^2 | 1x10^7 |     |        |   | (58) |
| Protists  | X    | 5x10^3 | 2x10^4 | 1x10^{10} |     |        |   | (58) |
| Protists  | X    | 1.8x10^4 | 5x10^5 | 2x10^4 | 5x10^5 |     |   | (59) |
| Protists  |      |      | 3x10^3 |     |        |        |   | (50) |
| Protists  |      |      | 1.4x10^6 | 1.6x10^7 |     |        |   | (60) |
| Protists  | 1x10^0 | 1x10^0 |     |        |   |        |   | (61) |
| Protists  |      |      | 1x10^6 | >     |        |        |   | (62) |
| Metazoa   | X    |      | 2x10^{-2} | 8x10^1 |     |        |   | (41) |
| Metazoa, copepods | X | 1x10^2 | > | few | 2.5x10^2 |     |   | (63) |
| Metazoa, copepods | 2x10^2 | > |     | 3x10^0 | 4x10^2 |     |   | (64) |
| Metazoa, copepods | X | 5x10^2 | > | few | 8x10^1 |     |   | (65) |
| Metazoa, copepods | X | 1x10^2 | > | few | 1.5x10^2 |     |   | (65) |
| Metazoa   | X    | 1x10^4 | > | 5x10^{-5} | 7x10^{-4} |     |   | (66) |
| Metazoa   | X    | X    | 2x10^3 | > | 6x10^{-4} | 5x10^{-9} |     |   | (67) |
| Metazoa   |      |      | 7x10^3 |     |        |        |   | (68) |
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