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34	The ocean's ability to sequester carbon out of contact with the atmosphere exerts an
35	important control on global climate. The biological pump drives carbon storage in the
36	deep ocean and is thought to function via gravitational settling of organic particles from
37	surface waters. However, the settling flux alone is often insufficient to balance
38	mesopelagic carbon budgets or meet the demands of subsurface biota. Here, we review
39	additional biological and physical mechanisms that inject suspended and sinking
40	particles to depth. Together, these "particle injection pumps" likely sequester as much
41	carbon as the gravitational pump, closing carbon budgets and motivating further
42	investigation of their environmental controls.
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59 Introduction

60 Open ocean waters store (sequester) carbon out of contact with the atmosphere on decadal to 61 millennial timescales, exerting a major control on global climate by regulating atmospheric 62 carbon dioxide partial pressure $(pCO_2)^1$. The magnitude of ocean carbon storage is governed by two well-established mechanisms that maintain a surface-to-deep ocean gradient of 63 dissolved inorganic carbon (DIC) – the biological and the solubility $pumps^{2,3}$. The solubility 64 65 pump delivers cold, dense, DIC-rich waters to depth mostly at high latitudes, whereas the 66 biological pump globally exports particulate organic carbon (POC) from surface waters. POC export is largely attributed to the gravitational settling of a subset of the particle assemblage^{1,4} 67 68 - a process we refer to as the "biological gravitational pump" (BGP). The BGP is the key link between upper ocean photosynthetic carbon fixation, the sustenance 69 of mid-water biota, and carbon storage in the oceans' interior^{4,5}, and is thought to account for 70 ~90% of the vertical DIC gradient, while the solubility pump explains the remainder¹. In the 71 absence of the BGP, models predict atmospheric pCO_2 would be higher by nearly twofold⁶. 72 73 Contemporary and paleoceanographic observations both reveal that carbon sequestration by the BGP is affected by environmental changes in light, temperature, stratification and nutrient 74 availability^{7,8}, and can itself drive dramatic climate shifts such as glacial-interglacial cycles⁸. 75 76 Future climate projections suggest that the functioning of the BGP will be altered by ocean global change^{7,9}, potentially feeding back on anthropogenic climate warming¹⁰. As a 77 consequence, quantification of its functioning requires a reliable baseline of accurate 78 79 measurements.

The underlying principles of the BGP are long established¹¹: organic particles are continually produced and recycled in sunlit surface waters, and a small fraction of these settle into the

82 oceans' interior. The strength of the BGP is often quantified as the rate of particle "export" from the euphotic zone, the surface mixed layer, or across an arbitrary horizon at 100m¹². As 83 84 they sink, particles undergo myriad transformations, which lead to pronounced vertical 85 attenuation of the particle flux that is often described as a power law relationship, referred to as the "Martin Curve"¹³. The efficiency of the BGP is defined here as the time that exported 86 87 carbon is kept sequestered from the atmosphere within the ocean's interior. It is driven by the 88 depth scale of flux attenuation and pathways of ocean circulation that carry remineralized carbon dioxide back to the surface¹⁴. Carbon is sequestered for timescales longer than a year 89 90 by particles that penetrate the permanent pycnocline (beneath the wintertime mixed layer) 91 and up to centuries by those that reach deep water masses (generally >1000m). Together, the 92 strength and efficiency of the BGP determine the total quantity of carbon sequestered 93 biologically in the ocean interior.

94 Recently, analyses of global and regional ocean carbon budgets have identified conspicuous 95 imbalances (i.e., two to three-fold less storage) when BGP export fluxes are compared with those derived from geochemical tracers^{15,16}, highlighting the need to reassess the pathways 96 97 that contribute to carbon storage. Furthermore, rates of site-specific particle export appear to be insufficient to meet the carbon demand of mid-water life (termed mesopelagic biota) by 98 two-to three-fold $^{17-20}$, but in one study can be balanced using community respiration 18 . There 99 100 is considerable debate over the reasons for these carbon deficits, ranging from biases inherent in observational technologies^{17,21} to the potential role of other carbon (dissolved and/or 101 particulate) delivery mechanisms to deep waters^{16,22,23}. Traditionally, the biogeochemical 102 103 functioning of the BGP has been evaluated from quasi one-dimensional (1D) observations of 104 particle flux (Box 1), and extrapolated using Earth System Models (ESMs, parameterised with observations²⁴⁻²⁶) and/or remote-sensing observations²⁶. This approach cannot capture 105

106 more complex mechanisms of carbon export that are highly variable in space and time (Box

107 1), potentially resulting in the reported carbon budget deficits.

108 Multiple lines of research have revealed the importance of additional export pathways, 109 physically (e.g. subduction) and/or biologically (e.g. large mesopelagic migrators) -mediated, that inject particles to depth, termed here Particle Injection Pumps (PIPs)^{23,27-30}. These 110 111 mechanisms can potentially export all particle classes to depth, and thus challenge the 112 conventional view of gravitational sinking as the dominant downward pathway for particles 113 into the oceans' interior. The characteristics of PIPs fundamentally change our understanding 114 of biological carbon sequestration: first, PIPs can animate particle transport spatially into 115 three dimensions (3D), in contrast with the BGP where the vertical dimension is predominant 116 (1D); second, global estimates of PIP carbon fluxes are significant relative to those for the $BGP^{27,28}$, and third, these mechanisms cannot be readily quantified using the traditional 117 118 toolbox applied to investigate the BGP (Box 1). Overall, the PIPs will increase the strength 119 of the biological pump beyond estimates based on gravitational flux alone, and can change its 120 efficiency by altering the depth of carbon export.

121 The fate of exported carbon following its delivery to depth has also proven more complex 122 and heterogeneous than previously recognized. Particle flux attenuation is now known to vary 123 systematically in space^{14,31,32} and time³³, suggesting the traditional empirical view¹³ must be 124 replaced by a mechanistic one that considers particle composition and architecture, microbial 125 metabolism, and transformation processes¹⁷.

Together, these developments stand to reshape our understanding of particle transport and remineralisation in the oceans' interior. Here, for open ocean systems we review: the mechanisms, rates, and depths of particle injection by each PIP; the potential for each mechanism to close observed deficits in ocean carbon budgets; and the corresponding

remineralisation depths of exported POC in the deep ocean. We finish by outlining future
research directions needed to synthesize these developments into a new mechanistic, fourdimensional (4D) view of carbon export and sequestration. The review does not detail the
important role of dissolved organic carbon subduction^{22,23}, nor cover the dark microbial
carbon pump³⁴ or chemolithotrophy³⁵ which have been reviewed elsewhere (S-Table 1).

135

136 Particle injection pump mechanisms

137 PIPs differ in their mechanisms, temporal-spatial scales (Fig. 1, Fig. 2a), and/or geographical 138 extent, but have common features: i) they can act on all particles from suspended to sinking 139 (Fig. 1); ii) they typically inject particles below the euphotic zone (i.e., the export depth for the BGP), potentially reaching depths >1000 m^{28-30} depending on the injection mechanism 140 141 (Fig. 1, Fig. 2b); iii) they occur concurrently with the BGP but cannot be measured with techniques developed to quantify gravitational settling 13,32 (Box 1); iv) their dynamic nature 142 (i.e., physical transport^{23,27,28} or patchiness of animal distributions³⁰) means that the interplay 143 144 between their vertical and horizontal vectors and temporal scales varies significantly (Fig. 1). 145 Hence, a 4D sampling framework is required to constrain them (Box 1). The main characteristics of each PIP are elucidated below. 146

Particle export driven by physical subduction includes several processes driving the vertical
transport of near-surface particles that act on different space/time scales: subduction caused
by mixed-layer shallowing (termed the mixed-layer pump^{29,36}); subduction by large-scale
(100-1000 km) circulation (termed the large-scale subduction pump)²³; and subduction by
mesoscale (10-100 km) to submesoscale (1-10 km) frontal circulation (termed the eddysubduction pump^{23,27,28}).

153	Carbon export by the mixed-layer pump is driven by biological accumulation of particles
154	throughout the spring/summer growth season, which are then diluted to the depth of the
155	mixed layer during winter, and left in the oceans' interior during early spring stratification
156	(Box 1). This pump operates on wide-ranging time-scales from days/weeks ³⁷ to seasons ^{29,37} ,
157	predominantly in mid and high latitude regions characterised by strong seasonal variability in
158	mixed-layer depth (Fig. 2a). Although these concepts are long-established ³⁶ , only recently
159	have they been scrutinised in detail using advances in optical profiling (BGC-Argo) floats
160	and satellite particle proxies to track particle accumulation rates in relation to changes in
161	surface mixed-layer depth (Box 1).
162	The large-scale subduction pump is a 3D advective mechanism directed from the seasonal
163	mixed-layer into the oceans' interior, driven by Ekman pumping and horizontal circulation
164	across a sloping mixed-layer ³⁸ . Subduction rates were first estimated for the North Atlantic ³⁹ ,
165	and then globally using data-assimilating models ⁴⁰ . The wide-ranging subduction rates (1-100
166	$m/year)^{39,40}$ are small relative to BGP particle settling rates ^{11,12} , but subduction occurs over
167	large regions of the global ocean boosting the magnitude of carbon delivery to depth.
168	The frontal-associated eddy-subduction pump subducts particle-rich surface waters on
169	timescales of days and across spatial scales of 1-10 km, driven by strong vertical circulation
170	associated with fronts and eddies ^{27,28,41-44} . Gliders are now used to map 3D dynamic eddying
171	flow fields (Box 1), finding evidence for penetration of high particle stocks (co-located POC
172	and chlorophyll indicative of viable phytoplankton) from the spring bloom, conspicuous as
173	distinct filaments at 100-350 m depth at the eddy periphery ²⁸ (Box 1). Mapping revealed the
174	co-location of high POC filaments and negative vorticity to depths near the permanent
175	pycnocline ²⁸ , and the mechanism is supported by high-resolution simulations in which eddy
176	subduction of particles is a recurring feature ⁴⁵⁻⁴⁸ . The strength of the eddy-subduction pump

is governed by the vigour and penetration of the vertical circulation, in conjunction with local POC stocks over the frontal area^{27,49}. Eddy subduction rates span 1-100 m d⁻¹ (c.f. 20 to > 100 m d⁻¹ for the BGP^{11,12}) depending on the eddy or frontal structure. Modelling indicates that these subducted particles are remineralised more rapidly (i.e., at relatively shallow depths) relative to gravitationally-sinking particles²⁷.

182 The concept for the 'mesopelagic migrant pump' is based on long-established observations of

diurnal vertical migration⁵⁰ (Box 1). This pump extends the remineralisation scale by

injecting particles to greater depth before decomposition begins 51,52 , as determined by gut

retention time of migrating animals⁵¹⁻⁵³ and the depth of their migration (typically $\sim 400 \text{ m}^{53}$).

186 The injected particles are zooplankton faecal pellets with sinking rates of 10-100's m d^{-1} (ref.

187 51), faster than loosely-packed organic aggregates settling from the surface 11,12 , and will

188 penetrate deeper in the water column before remineralisation. This pump therefore influences

all important facets of the particle flux that govern carbon sequestration – total export rate,

190 depth of peak flux, and flux attenuation depth scale.

191 Diurnal vertical migration results in active subsurface transport and carbon sequestration, and

is usually reported for mesozooplankton and often included in BGP estimates⁵¹. However,

193 vertical migration by larger mesopelagic carnivorous organisms (from greater daytime depths

than mesozooplankton) are not sampled by conventional BGP approaches^{52,54}. Targeted

studies (Box 1) have quantified this pump driven by large mesopelagic migrant carnivores in

the Pacific⁵⁴, and other regions (S-Table 1). The underlying mechanism is upward migration

to graze mesozooplankton⁵⁴ followed by rapid (hours) downward migration⁵³, with

respiration (release of CO_2), exudation, and defecation (release of POC/DOC)^{51,55} often

below the permanent pycnocline⁵⁶, at depths up to 600m (Box 1).

Trawl surveys suggest that ~50% of mesopelagic organisms migrate, ranging regionally between 20-90% depending on temperature, turbidity and oxygen concentrations^{54,56}. The carbon sequestration rate by this pathway is governed by the metabolic transfer efficiency of migrators, and particles are injected at their residence depth, often at the upper boundary of oxygen minimum zones where their respiration intensifies oxygen depletion⁵³.

205 Active transport by vertically-migrating metazoans can also occur on longer timescales (Box

1). For example, in high latitude regions the winter hibernation of copepods (members of the

207 mesozooplankton) at depths between 600-1400m gives rise to a so-called 'seasonal lipid

208 pump^{30'}: during hibernation, they catabolise carbon-rich lipids accumulated during summer in

209 upper layers and thereby shunt carbon (but not nitrogen and phosphorus) below the

210 permanent pycnocline 30 . The strength of the seasonal lipid pump is governed by copepod

abundance, size and temperature, which together control their respiration rate and help

explain the existence of carbon flux hotspots (i.e. patchiness)³⁰.

213 Another vertical export mechanism that operates on seasonal migration timescales is

214 mortality at depth of hibernating zooplankton particularly in high latitude regions 57,58 ,

sequestering carbon to depths >500 m depth. Global extrapolation of seasonal lipid pump

fluxes, along with the over-wintering mortality flux is problematic due to difficulties in

sampling and generalizing across distinct regional mechanisms³⁰ (S-Table 1).

218

219 The potential for double accounting

The export flux from the BGP is mediated by sinking particles, whereas PIPs can provide
additional pathways for all particle classes, from suspended to sinking, to exit the surface
ocean (Fig. 1). Thus, there is potential overlap between particles delivered from the surface

223 ocean to depth via the BGP and by injection from PIPs. Such overlap – termed here as

224	'double-accounting' – may occur where particles associated with the BGP and a PIP are
225	difficult to distinguish and hence could be attributed to more than one pump (Fig. 1). At
226	depth, transformations such as aggregation alter particle characteristics, including size and
227	sinking rate, and hence particles injected by the PIPs can join the sinking flux usually
228	attributed the BGP (Fig. 1). A further factor that introduces overlap between the BGP and
229	PIPs results from the inclusion, for historical reasons ⁵⁹ , of one component of the mesopelagic
230	migration pump (diurnal migration by mesozooplankton) into the 1D sampling framework of
231	the BGP, while other components (e.g. patchier diurnal migration by larger mesopelagic
232	carnivores ⁵) are not. Hence double-accounting can confound our understanding of the
233	relative importance of PIPs to ocean carbon storage.
234	Is it possible to tease apart these areas of overlap? Forty years study of the BGP has
235	uncovered a complex biogeochemical system with multiple drivers and distinguishing
236	characteristics ^{11,60} . This body of research helps to frame the differences and similarities
237	between particles delivered to depth by PIP's and those settling via the BGP. Each PIP is
238	distinct with respect to its combination of injected particle type (suspended cells to faecal
239	pellets of large mesopelagic migrants), the timing and depth of injection (Fig. 2a-b), and
240	associated particle transformations (aggregation/disaggregation) ^{11,12,61} . Additionally, the
241	subsurface "fate" of particles (i.e. where they remineralize), which determines the longevity
242	of carbon sequestration, is driven by the complex interplay between these properties and
243	transformations ^{12,60,61} : Particle composition and architecture set their sinking speed, while
244	myriad processes that are biologically- (microbes/zooplankton) and physically-mediated
245	(fragmentation/ disaggregation) ^{12,62-64} decompose and repackage them over depth (Fig. 1).
246	Therefore, particle fate provides another avenue to distinguish the contributions of PIPs from
247	the BGP.

248	To date, evidence on the subsurface fate of injected particles has been largely indirect ^{27,28,49} .
249	Surveys of eddy-subduction pumps suggest that injected particles may be remineralised at
250	depths <200 m, based on ammonium peaks ⁴⁹ , time-series of biogeochemical gradients ²⁸ , or
251	particle modelling studies ²⁷ . In the NE Atlantic, reported high rates of particle
252	remineralisation (glider-based biogeochemical gradients) must be reconciled with concurrent
253	evidence of coincident, coherent chlorophyll plumes at depths >300 m indicative of
254	subducted viable phytoplankton ²⁸ . This glider-based time-series reveals pronounced
255	patchiness ²⁸ suggesting that inference of the fate of injected particles even from state-of-the-
256	art observations is challenging.
257	Better constraining the contribution of each PIP to mesopelagic carbon budgets will require
258	characterisation of the injected particle assemblage and their transformations during
259	downwards transport ^{12,65-68} . Particle aggregation in PIPs may be driven by
260	convergence/subduction ⁶⁹⁻⁷⁰ and/or differential sinking ^{65,67} , potentially leading to altered
261	modes of subsurface transport (Fig. 1). BGC-Argo profile observations allow quantification
262	of the size, type, seasonal succession, and penetration depths of particles injected by the
263	mixed-layer pump ³⁶ – properties which have the potential to differentiate them from fast-
264	sinking particles (i.e., BGP) whose distinctive 'spiky' bio-optical signature is readily detected
265	using multiple sensors ⁷¹ (S-Figs. 2 and 3). Advances in bio-optics are already making cryptic
266	signatures associated with slow-sinking particles and zooplankton vertical migration less
267	opaque, lessening the possibility of double-accounting. Such double-accounting may be
268	avoided through the identification of unique characteristics of pumps including seasonality
269	(Fig. 2a), distinctive regional features ³⁰ , or multi-variate oceanographic diagnostics ⁷² .
270	

272 Carbon sequestration potential

273 The potential carbon sequestration by each PIP can be quantified as the product of their 274 carbon injection rate and their sequestration timescale, i.e. time until remineralised carbon is 275 returned to the surface (see Supplementary Methods). This timescale is determined both by 276 the injection depth of particles and their eventual fate, i.e. the degree to which they sink or circulate through the ocean before remineralising to CO_2 . In general, deeper particle injection 277 278 and rapid sinking translates to longer carbon sequestration because the "passage time" from 279 the ocean interior to the surface increases with depth (Fig. 2b). Here, we assemble prior 280 estimates of carbon injection rate and depth (S-Table 1), along with new modelling projections (Fig. 2), to estimate carbon sequestration by each PIP and assess their 281 282 significance relative to the BGP.

283 Some targeted studies provide concurrent estimates of carbon injection by individual PIPs and the BGP^{27,28}, whereas others^{30,54,57,58} facilitate comparison of regional-scale PIP fluxes 284 285 with independent estimates of the BGP. Both approaches reveal that PIPs each have the potential to contribute significant rates of POC export. The reported upper bounds of global 286 PIP estimates summed together is 8.7 Pg C yr⁻¹, which is comparable to the BGP export flux 287 (Table S1). This comprises 1.1-2.1 Pg C yr⁻¹ for the large-scale/mesoscale physical pumps 288 (also includes DOC^{22,23}), and 0.25-1.0, 0.9-3.6 and (-0.09) to 2.0 Pg C yr⁻¹ from the lipid 289 290 seasonal, mesopelagic migration, and eddy-subduction pumps, respectively (Fig. 2c). Thus, 291 their cumulative contribution may be as much as $\sim 40\%$ of total particle export (i.e., 292 BGP+PIPs) suggesting considerable potential to resolve the imbalances reported for mesopelagic carbon demand¹⁷, between nutrient and carbon export budgets¹⁵, and to lessen 293 294 the variability between model estimates of global carbon sequestration (S-Table 1).

295 We estimated the sequestration timescales for each PIP based on the "passage time" from the injection depth to the surface in an observationally-constrained ocean circulation model¹⁴. 296 297 Particles injected at the depth of the wintertime mixed-layer by the large-scale physical 298 pumps (mixed-layer and subduction) result in sequestration for 25-100 years, assuming 299 subduction occurs before re-entrainment next winter. In turn, deeper injection by the eddy 300 subduction pump (up to 450 m), mesopelagic migration pump (up to 600 m), and seasonal 301 lipid pump (up to 1400 m) translates to sequestration timescales up to 150, 250, and 500 302 years respectively (Fig. 2b). These timescales will increase if it is assumed that sinking rather 303 than suspended particles are injected, which remineralise deeper than the injection horizon 304 (see Supplementary Methods). 305 Given the wide-ranging estimates of carbon injection rate (Fig. 2c) and depth (Fig. 2b) for 306 each PIP, oceanic carbon sequestration by these mechanisms cannot be estimated with 307 precision (Fig. 2d). However, choosing central values from the reported ranges of each 308 property allows a first order comparison between PIPs and the BGP. The mesopelagic 309 migration pump emerges as the most significant PIP, potentially storing $\sim 60\%$ as much carbon as the BGP in the ocean interior if large, sinking particles (i.e. faecal pellets) are 310 311 injected. The C storage potential of the seasonal lipid, eddy-subduction and large subduction 312 pumps are ~20%, 10% and 5% of the BGP respectively, assuming each injects suspended 313 particles. The latter small net value is due to offsetting of subduction by strong obduction (upward transport of water parcels) in the equatorial oceans³⁹. Based on these central values 314 315 (Fig. 2d), it is likely that the reservoir of respired carbon in the ocean interior contributed by 316 the PIPs approaches that contributed by the BGP, and may therefore help to close globalscale mesopelagic carbon budgets^{15,16}. 317

318

Tracer constraints on the fate of exported carbon

320 Oceanic carbon sequestration by the BGP and wide-ranging biophysical mechanisms that 321 inject biogenic particles to depth depends critically on the fate of exported carbon (Fig. 2). 322 However, at present tracing the remineralisation of particles (regardless of their export 323 pathway) as they settle and circulate through the global ocean remains a logistical challenge, 324 due to the difficulties of deep-water particle sampling. Recently, new methods have used 3D 325 ocean data assimilation models to leverage geochemical "remineralisation tracers" including 326 oxygen and nutrients. These tracers integrate particle remineralisation signatures over long 327 timescales, and their global distributions are characterised by orders of magnitude more observations than are available for particles^{16,31,73}. Two distinct approaches have been applied. 328 329 The first diagnoses remineralisation rates directly from phosphate accumulation along 330 transport pathways in a circulation model, and reconstructs particulate flux profiles required to explain the global distribution of remineralised phosphate³¹. The second assimilates 331 geochemical and satellite data into mechanistic biogeochemical models to optimise key 332 333 particle flux parameters, yielding mechanistic insights while leveraging the observations less directly⁷³. 334

335 Both approaches have yielded similar results and provide evidence for regional variability in 336 particle flux attenuation, with the flux attenuating slowly at high latitudes and quickly in subtropical gyres, while the tropics lie between these two extremes (Fig. 3a). These 337 338 simulations reveal that carbon exported from high latitude and tropical surface waters is 339 sequestered longer in the oceans' interior than carbon exported in the oligotrophic gyres 340 (Figure 3b), with important implications for feedbacks between the particle export and global 341 climate. Atmospheric pCO_2 is likely more sensitive to past changes in high latitude export than previously recognised⁸, and the future expansion of subtropical habitats⁹ may result in 342 less efficient (although not currently quantifiable) carbon sequestration in a warming world. 343

Regional variations in particle flux attenuation have largely been interpreted in terms of the balance between decomposition and sinking rates³². A likely explanation for the diagnosed latitudinal pattern is the temperature-dependent metabolism of heterotrophs responsible for particle decomposition^{32,73}, although variations in particle size and/or ballast are valid alternatives⁷³. There may also be a secondary effect of oxygen, with decomposition slowing in anoxic zones^{73,74}, and even hypoxic waters due to anaerobic microenvironment formation in particles⁷⁵.

To some degree, model-derived particle flux profiles may also reflect the relative magnitude 351 352 of different export pathways (PIPs and BGP), which vary in the injection depth and nature of 353 particles they supply, since geochemical tracers integrate the effects of all export mechanisms. 354 Deep injection by PIPs would result in slower flux attenuation over depth, whereas injection 355 of suspended particles that remineralise shallower in the water column would be diagnosed as 356 rapid flux attenuation. Predicting future changes in ocean carbon sequestration will require a 357 better understanding of the contribution of injection versus remineralisation processes to 358 sequestration efficiency (Fig. 3b), given the different environmental sensitivity of these 359 processes.

360 The need for prediction motivates development of new techniques to distinguish particle flux 361 associated with the BGP and each PIP. Particle stoichiometry (i.e., C:N:P) may be central to 362 identifying particular mechanisms that decouple their export. For example, diagnosing 363 oxygen consumption between 500-1500 m (depth of zooplankton hibernation) without concomitant nutrient accumulation would point to carbon export by the seasonal lipid pump³⁰. 364 365 Alternatively, diagnosing seasonal cycles of nutrient accumulation and oxygen consumption 366 rates would help distinguish remineralisation of particles exported by physical pumps versus 367 particle settling, which should exhibit distinct seasonality (Fig. 2a). This approach may soon

368 be possible given the burgeoning spatial/temporal resolution of tracer data provided by BGC-

369 Argo floats (S-Figure 1), and emerging float sensor technology (S-Table 2).

370

387

371 Extrapolation – towards a 4D view of particle export

372 Improving the accuracy of the initial estimates of the magnitude of carbon sequestration 373 presented in Figure 2d requires the development of a 4D picture of particle flux and storage 374 in the oceans' interior. It is clear from our synthesis of PIP mechanisms that multiple scales, 375 from sub-mesoscale to basin, must be accommodated if PIPs are to be assembled, first 376 spatially and then temporally, into a complete 4D picture. Again, lessons on how to approach 377 such upscaling can be gleaned from BGP research which imprinted both spatial and seasonal signatures (satellite remote-sensing/modelling)²⁶ onto short-term (days-weeks) observations 378 379 taken at specific sites (Box 1). The timescales and lifetimes of features such as submesoscale 380 eddies/fronts or seasonal mesopelagic export signatures (Fig. 2a) must be characterized to 381 define the temporal footprint of each PIP and move towards a 4D viewpoint. This framework 382 must be linked to the seasonality of pelagic particle production to assess if there is distinctive 383 period for the subduction of significant stocks of these upper ocean particles (Fig. 2a). For 384 example, it is well-established that submesoscale dynamics are strongly seasonal, with stronger and deeper penetration during winter than summer⁷⁶. 385 386 Some published approaches towards extrapolating PIP's globally, and to climatological time

scales, are outlined in S-Table 1. The identification of the specific drivers of each PIP

- 388 mechanism should help improve modelling and hence extrapolation. We advocate the utility
- 389 of explicitly incorporating the different PIP mechanisms into predictive, mechanistic models
- 390 as a means to extrapolate PIPs into 4D. In the case of the extrapolation of the submesoscale
- 391 eddy subduction PIP, increasing the model grid resolution to incorporate these features is

392	necessary and is now achievable in regional configurations ^{77,78} . In contrast, other physically-
393	mediated PIPs such as the large-scale subduction and mixed-layer pumps are already
394	represented in global models, and so their extrapolation requires the development of
395	diagnostics to enable the simulated POC/DOC distributions to be better evaluated against
396	observations ²³ . At present, the biologically-mediated PIPs are not incorporated into state-of-
397	the-art biogeochemical models ^{9,14,31,77,78} . While simulating animal behaviour at the global
398	scale remains a grand challenge in ocean modelling, simple parameterisations have been
399	developed to predict the geochemical effect of the mesopelagic migrant pump ⁶ , which might
400	be further expanded to incorporate hibernation and therefore the seasonal lipid pump. It is
401	only very recently that diel vertical migration has been incorporated for the first time in a
402	global ocean general circulation model and used to estimate the associated flux of carbon at
403	the global scale (see Aumont et al. in S-Table 1). Although promising, this approach remains
404	challenging because it is based on a computationally-intensive, end-to-end ecosystem model
405	in which all trophic levels from phytoplankton to top predators interact.

407 Transforming our view of ocean carbon export

408 Our synthesis of physically- and biologically-mediated PIPs reveals that they are directly 409 transporting significant stocks of biogenic particles to depth, of a cumulative magnitude that 410 may be equivalent to the carbon storage of the BGP. The potential of PIPs to make a major 411 contribution to the ocean carbon budget must now be explored in more detail, commencing 412 with those PIPs that are most likely to contribute to carbon sequestration. Synthesising 413 estimates of particle export, injection depth, and circulation timescales reveals that the 414 mesopelagic migrant pump has the greatest potential to contribute to carbon sequestration, 415 followed by the seasonal lipid pump and the various physical pumps (Fig. 2d). In the case of

the seasonal lipid pump, its geographical realm of influence is already established³⁰, whereas
less is known about the regional contributions of the mesopelagic migrant pump⁵.

418 For all PIPs, the most pressing research issue – needed to address double-accounting issues 419 and improve estimates of carbon sequestration - is to better understand the mechanisms of particle transformations^{17,65-68} (Fig. 1) within a 4D framework. Specifically, the fate of 420 421 exported particles between their injection depth and the permanent pycnocline remains poorly 422 constrained. A first step will be improved particle characterisation, in particular the ability to 423 distinguish zooplankton from other particle types, and to aggregate Particle Size Distribution 424 (PSD) profiles through the development and application of new sensors (S-Table 2). Future development of acoustic and imaging technologies⁷⁹ must be deployed on a range of 425 426 platforms from ships (i.e., calibration) to an array of long-lived (i.e., years), geographically-427 diverse BGC-Argo floats. These developments towards improving particle characterisation 428 will reduce the likelihood of double-accounting. Moreover, the alignment of BGC-Argo 429 deployments (Box 1) with the characteristic space and time scales of PIPs will enable better 430 quantification of the role of patchiness in driving observed local/regional hotspots in biological PIPs^{30,54,56}. In time, following the development and testing of a Coastal-Argo 431 432 platform, they can also be deployed to coastal and shelf seas to explore the role of PIPs in 433 these regions (S-Table 2).

The way forward in refining estimates of the contribution of PIPs in closing the ocean carbon budget¹⁵⁻¹⁷ also requires leveraging advancements in ocean biogeochemical modelling. Models are valuable testbeds to probe the sensitivity of carbon storage mechanisms, and guide future observations. For example, model sensitivity analyses point to the pivotal role of PSD in determining the fate of exported carbon^{31,73}, but the processes that set the PSD of exported particles and its evolution over depth remain crudely parameterized. Developing robust models of particle transformations between multiple size classes, and incorporating them into general circulation models, will allow us to trace the fate of particles injected by
different PIPS and dissect their contribution to carbon sequestration, while avoiding doubleaccounting issues.

Inverse methods that can assimilate PSD fields from new BGC-Argo technologies⁸⁰ will
allow models to "learn" from the data, further refining them to best reflect the real ocean.
Furthermore, downscaling of physical models is essential to simulate the locations of PIP
injections in support of observational programmes such as high resolution altimetry⁸¹, and the
integration of detailed particle transformations into submesoscale models⁸².

To transform the comprehension of particle export from one- to three- and eventually four-449 dimensions, machine learning approaches⁸³ will need to be employed, which can be trained to 450 451 predict unknown variables such as particle flux from better sampled variables. Approaches like artificial neural networks⁸⁴, will enable and enhance the upscaling of local/regional 452 datasets needed to provide more robust extrapolation^{85,86} to depth, regionally, and annually of 453 454 each PIP. This upscaling is essential to refine estimates of the contribution of each PIP to 455 carbon sequestration. BGC-Argo datasets will also eventually be combined with new satellite products such as hyperspectrally-resolved ocean colour observations of biology processes⁸⁷ 456

457 and submesoscale characterisation of sea level using high-resolution altimetry 81 .

Satellite and water-column remote-sensing, along with targeted process studies, will yield
expansive datasets that can be assimilated into regional and global models of ever increasing

460 realism and resolution. Together, these approaches will lead towards a robust, four-

dimensional view of carbon sequestration by the ocean's multi-faceted bio-physical particlepumps.

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Figure 1 Interplay between particle characteristics, mode of export (BGP or PIP), delivery depth and larger scale ocean circulation for a range of pumps. In the upper panel, the box (top left) represents mixed-layer particle types, which either form large sinking particles (i.e., within the BGP, such as faecal pellets, marine snow) or are injected to depth (i.e., PIPs, suspended/ slow-settling heterogeneous particles and cells (i.e., including healthy, slow-sinking phytoplankton⁸⁸)). The vertical yellow arrow signifies the BGP; black lines physically mediated PIPs; and purple lines biologically mediated PIPs. The delivery rates of particles to subsurface strata (in m d⁻¹, ? denotes not known) are presented for each pump. Patchiness in the distribution of vertically-migrating animals (top right) plays a role in driving three-dimensional particle delivery to depth^{89,89}, and is denoted by different fish or copepod stocks in the upper ocean. The box (middle left) presents different particle transformations central to the BGP¹², but whose role is not known so far for PIPs. They include microbial solubilisation, aggregation (marine snow denoted by aggregation 1; heterogeneous faecally-dominated aggregates (aggregation II) and/or dissaggregation¹⁸ to form/break down heterogeneous particles (hatched brown symbols). In the lower panel, depths in parentheses are the reported delivery depths, with the BGP (and some PIPs) exporting some particles to the sea floor. Blue curved arrows represent transport of subsurface material along downward-sloping isopycnals (white dashed lines). Major unknowns include whether physical transport by PIPs can cause particle aggregation (signified by ? in the middle panel below subduction pump, and also applicable for the mixed-layer pump) and hence alter their mode of injection towards gravitational settling (i.e., the BGP). Other unknowns include the potential ballasting role of small mineral particles such as aerosol dust for PIPs.



Figure 2 Carbon export and storage by PIPs compared to the BGP. a) Idealised seasonality of the PIPS for regions that exhibit strong seasonality, where a Spring Bloom dominates C export by the BGP (dark blue line). Coloured bars indicate season of peak C export by the PIPs (acronyms defined in legend). Note the large-scale physical pump should be strongest when mixed layers are deepest, but is likely operative all year (dashed line). b) Sequestration efficiency of the PIPs. Black lines represent the global-mean sequestration timescale for carbon injected at a given depth, defined as the time for remineralised carbon to circulate back to the ocean surface, computed in a data-constrained circulation model (see S-Methods). Solid line assumes that particles are suspended, so remineralisation occurs at the injection depth, whereas dashed line assumes that particles are sinking and remineralise over depth (see Methods). Colored bars show injection depth range of the BGP and PIPs. The efficiency of each pump is defined as the sequestration time from its injection depth. c) Strength of the pump mechanisms, defined as their rate of carbon export or injection (see Table S1). "All PIPs" refers to the sum of the five individual PIPs d) Ocean carbon storage by each pump, defined as the product of the strength (\mathbf{c}) and efficiency (\mathbf{b}). Two scenarios are shown for each PIP, using the sequestration time for suspended (circles) and sinking (square) particles, whereas the BGP is assumed to export only sinking particles. For the sum of PIPs, we present a "most likely" scenario, in which the migrant pump injects sinking particles (faecal pellets), and all other PIPs inject suspended particles (triangle).





Box 1 Approaches used to investigate downward particle export, from the BGP to PIPs.

The BGP is quantified in a biologically-patchy upper ocean (green filaments) using ship-based surface sampling (particle production) and subsurface particle interception by sediment traps, most recently neutrally-buoyant traps downstream of particle source regions (orange instruments, a). This coupled surface-subsurface sampling strategy is logistically-complex, temporally- and spatially-restricted (i.e., represented here by a "statistical funnel"^{91,92}, see a), and hence provides a '1D' view of particle export that is extrapolated to the basin scale using satellite observations and/or modelling. This '1D' viewpoint cannot measure the PIPs presented in b) to e), and is contrasted in a) with the 4D view^{29,93} obtained by an ensemble of BGC-Argo floats (white instruments).

(b) the mixed-layer pump, in which particles are detrained when the pycnocline (blue dashed line) shallows, can been addressed regionally through backscattering (a proxy for POC) profiles measured by BGC-Argo floats³⁶, or globally using satellite surface-ocean backscattering and Argo/BGC-Argo (density/backscattering) vertical profiles³². c) the seasonal lipid pump is quantified using surveys of overwintering copepods at depths below the permanent pycnocline and subsequent scaling of their lipid-enriched biomass in carbon content³³. d) the eddy subduction pump can been quantified using gliders (pink instruments) and subsequent modelling³¹, BGC-Argo floats (bio-optics/oxygen/physics)⁹³ or surveys based on multiple POC profiles in conjunction with coupled models (regional circulation/particle dynamics)^{30,49}. e) quantification of the mesopelagic migrant pump (active diel transport of carbon by mid-water biota, denoted by moon and sun symbols) requires mid-water trawl surveys along with metabolic modelling^{54,55}. Some multidisciplinary studies^{30,31,49} have combined these approaches to cross-compare export flux from the BGP (green arrows (d)) and the eddy subduction pump (purple arrows represent subsurface particle maxima recorded at the eddy periphery³¹). Note, the large-scale subduction pump²³ is not presented here.

