**The relationship between zinc, its isotopes, and the major nutrients in the North-East Pacific**

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**Abstract**

Trace metal micronutrients play key roles in photosynthesis by oceanic phytoplankton. Though they are required in much smaller amounts than the major nutrients (P, N, Si), their bioavailable forms are also present in the seawater solution at much lower levels. Relationships between the dissolved chemistry of the nutrient-type trace metals, their stable isotope variations, as well those of the major nutrients, have highlighted the importance of biological and physical processes in the Southern Ocean in controlling their oceanic biogeochemistry. However, the first-order Southern Ocean processes are overprinted by vertical cycling in other parts of the ocean, particularly those remote from the Southern Ocean, with the North Pacific standing out in particular. Here we present new zinc (Zn) concentration and isotope, as well as major nutrient data, for the NE Pacific and couple these new data with a compilation of published data from across the region, with the objective of better understanding the impact of this important region on oceanic biogeochemical cycles.

The new Zn isotope data for two stations along Line P (P04 and P26) show a large range in 66Zn in the upper ocean (-0.4‰ up to >1‰), associated with a very small isotope fractionation but extreme depletion of the dissolved pool during photic zone biological uptake, and the regeneration of this cellular Zn at very shallow depths (50m). Beneath this, the two profiles approach the 66Zn value of +0.5‰, seen throughout the deep ocean, by about 500m. The minimum 66Zn resulting from regeneration is associated with very high Zn concentrations, particularly at the marginal P04 station where diatoms dominate the phytoplankton ecology. Combining the new data with published Zn and major nutrient concentrations from across the North Pacific, emphasises the role of vertical biological cycling in controlling regional biogeochemistry in the North Pacific, resulting in the partial overprinting of biogeochemical signatures transported out of the Southern Ocean by the ocean circulation. Zinc isotope data document the uptake of this metal into diatoms and the co-regeneration of Zn with phosphate in the upper water column. Silica in contrast is regenerated at greater depth, resulting in a decoupling of the Zn-Si correlation that is set in the Southern Ocean and that dominates the Atlantic. Previous work has suggested that the decoupling of Zn and Si in the North Pacific results from removal of Zn (and other metals) to water column particulate sulphide. In our dataset, and in the compilation of data documenting relationships between Zn and the major nutrients across the North Pacific, this decoupling is clearly due to the different lengthscales of regeneration for organic matter (Zn and P) and diatom opal (Si).

**1. Introduction**

Close to half the carbon fixation, and half the molecular oxygen production, on Earth is due to photosynthesis carried out by oceanic phytoplankton (Field et al. 1998). These organisms have absolute physiological requirements for certain trace metals, which act as essential components of the photosynthetic apparatus, as cofactors in enzymes essential for carbon fixation, and for the acquisition of major nutrients (e.g. Morel et al., 2014). While the cellular requirements of these metals are modest relative to the major algal nutrients (P, N, Si), bioavailable concentrations of metals, and Fe in particular, can be small enough so as to limit the growth and influence the community composition of marine phytoplankton (Boyd et al., 2007; Bruland et al., 2014). Limitation, or co-limitation of cell growth and reproduction is less systemic for other metals, but has sometimes been demonstrated in culture (De La Rocha et al., 2000). It is also the case that cellular quotas relative to oceanic abundances are as high as, or higher than, Fe for certain metals in important organisms – e.g. Zn in diatoms (Twining and Baines, 2013).

The impact of these and other processes on the distributions of oceanic trace metals has been a topic of significant interest for decades (e.g. Bruland et al., 1980), but has received new impetus in recent years. Much more extensive, and higher spatial resolution, trace metal datasets have become available through the GEOTRACES programme ([www.geotraces.org](http://www.geotraces.org)). This data revolution has in turn spurred renewed interest in both new and established observational approaches. For example, though relationships between the dissolved concentrations of trace metals and the major nutrients, and the dominant biological control on these relationships, have long been a focus (e.g. Bruland et al., 1980; Martin et al., 1989; Lohan et al., 2002; Janssen and Cullen, 2015), larger trace metal datasets have made this approach much more powerful. The stable isotope systems of these metals are much more recently developed tools, and researchers are beginning to exploit the timely coincidence of the advances in mass spectrometry and the availability of the clean, large, samples that permit this approach (e.g. Abouchami et al., 2011; Xue et al., 2013; Zhao et al., 2014; Cameron and Vance, 2014; Conway and John, 2014a,b; Conway and John, 2015; Xie et al., 2015).

One outcome of ocean-basin scale sections for trace metals has been the realisation that one-dimensional vertical cycling of trace metal nutrients is only one of the key processes controlling their distributions in a three-dimensional ocean. Thus, for example, Vance et al. (2017) used relationships between dissolved Zn, Si and PO4 to highlight the significance of uptake by Southern Ocean diatoms in creating biogeochemical signals which, when coupled to the physical ocean circulation, appear to control patterns in Zn-Si-PO4 co-variation on a near global scale. This extends to trace metals the concept of Southern Ocean control previously well-established as key to the global decoupling of silica and nitrate (Sarmiento et al., 2004, 2007). Similarly, the Southern Ocean and the water masses sourced there are emerging as key for global patterns in Cd distributions (e.g. Baars et al., 2014; Quay et al., 2015; Xie et al., 2015; Middag et al., 2018). In the case of Cd, specific isotope signals are also generated in the Southern Ocean and advected northwards in mode, intermediate and deep waters, at least as far as the Equator (e.g. Abouchami et al., 2011; Xue et al., 2013; Sieber et al., 2019).

The first order significance of Southern Ocean biogeochemistry in explaining the origins of oceanic Zn-Si-PO4 co-variation, as well as the physics of the ocean circulation in transporting these signals globally (Vance et al., 2017; de Souza et al., 2018), has been confirmed by subsequent studies (Weber et al., 2018; Roshan et al., 2018; Middag et al., 2019). For example, Middag et al. (2019) suggest that no clear signal of *in-situ* vertical cycling processes is seen in the Atlantic. Two other recent large-scale studies have suggested that in the North Pacific the picture is somewhat less clear. Thus, while Weber et al. (2018) conclude that two thirds of the dissolved Zn in the deep North Pacific originates through uptake and regeneration in the Southern Ocean, they also highlight the potential importance of extra-Southern Ocean vertical cycling (both uptake and regeneration, and reversible scavenging) for the origin of the remaining third. Roshan et al. (2018) also suggest that 38±32% of the Zn in the deep North Pacific could originate through either extra-Southern Ocean uptake and regeneration, downward transfer by reversible scavenging, or addition of hydrothermal Zn to the deep ocean. With reference to the subarctic North Pacific in particular, a number of other recent studies (Janssen et al., 2014; Janssen and Cullen, 2015; Kim et al., 2017) have noted a decoupling of Cd and Zn distributions from those of the major nutrients. It has been suggested (Janssen et al., 2014; Janssen and Cullen, 2015) that there may be an impact on the distribution of these metals, particularly at intermediate levels, from water column sulphide precipitation in the low oxygen zones that characterise the NE Pacific.

It is perhaps unsurprising that it is particularly in the subarctic North Pacific that Southern Ocean signatures are overprinted by local basin or sub-basin-scale processes. Along with the North Atlantic, it is the ocean basin most remote from the Southern Ocean. Unlike the North Atlantic, there is no deep water formation in the North Pacific, and the major interior water mass of North Pacific origin, North Pacific Intermediate Water, is only seen in the subtropics (e.g. Yuan and Talley, 1996). The impact of local and basin-scale biological cycling is more likely to be strongly visible in the poorly ventilated waters in and below the subarctic gyres. Here we present new Zn isotope data that have a direct bearing on the issue of water column cycling of Zn. We go further, however, to couple these new data with a compilation of published data from across the NE Pacific, and to compare patterns in the relationships between Zn and the major nutrients to global relationships. The picture that emerges suggests that local export of nutrients significantly overprints signals transported from outside the basin and that, together with the Southern Ocean, the subarctic North Pacific is an important location for export of Zn from the surface to the interior ocean.

**2. Oceanographic Setting**

The focus of this paper is the subarctic northeast Pacific. The locations from which the new data derive (white circles), and those for which we discuss published data (black squares), are shown in Fig. 1.

The northeast Pacific is subdivided by the Subarctic Front (Fig. 1) into the Alaska Gyre to the north and the northern part of the Subtropical Gyre to the south (e.g. Yuan and Talley, 1996). The Subarctic Front not only marks an abrupt change in physical properties, with colder and fresher water to the north, but also an important biogeochemical transition. To the south, surface waters are depleted in major nutrients, while the region to the north is classified as a high-nutrient low-chlorophyll (HNLC) region, with low standing phytoplankton stocks, significant residual major nutrient concentrations (e.g., Si, Fig. 1). Here, artificial enrichment experiments have shown Fe to be limiting to primary productivity (e.g. Martin et al., 1989; see Boyd et al., 2007 for a summary). Major nutrient concentrations decrease again north of 55°N and towards the Alaskan coast.

During late summer, in response to surface warming, the water column is characterized by a thin (~30 m) mixed layer, which shoals towards the coast (e.g. Whitney and Freeland, 1999). Below this a permanent halocline occurs at around 100-150m, at the limit of winter storm and convective mixing. North Pacific Intermediate Water (NPIW; Yuan and Talley, 1996) does not significantly influence the study area and mesopelagic water is very poorly ventilated. Thus, the depth interval 500-2000m is characterised by a low oxygen zone (Janssen and Cullen, 2015), with O2 concentrations ≤ 50 mol kg-1 at all stations shown in Fig. 1 (Martin et al., 1989; Janssen and Cullen, 2015; Conway and John, 2015). Radiocarbon is more depleted in these intermediate waters than any other part of the ocean (e.g. Östlund and Stuiver, 1980). These intermediate waters are sourced from abyssal waters below, and probably to the west, via upwelling and diapycnal transformation (Roemmich and McCallister, 1989). The abyssal waters of Southern Ocean origin flow into the NE Pacific from the south and west (Roemmich and McCallister, 1989), and exhibit higher levels of oxygen and slightly younger radiocarbon ages (Östlund and Stuiver, 1980).

The microbial ecology and phytoplankton dynamics of the subarctic NE Pacific is of particular importance to the discussion of trace metals and their isotopes, as well as their relationships to the major nutrients, later in this paper. Upwelling in the middle of the cyclonic Alaskan gyre (Fig. 1) is marked by domed isotherms (Martin et al., 1989), bringing water rich in major nutrients to the surface. The first measurements of the concentrations of dissolved trace metals (Martin et al., 1989), as well as a number of Fe enrichment experiments (e.g. Martin et al., 1989; see Boyd et al., 2007 for summary), demonstrated the importance of Fe limitation in the open-ocean NE Pacific. Studies of phytoplankton ecology and export production have focused on Line P, and especially Ocean Station Papa (or P26, Fig. 1) in the HNLC region (e.g. Thibault et al., 1999; Boyd and Harrison, 1999). Here, chlorophyll-*a* concentrations (chl-*a*) are usually around 0.2-0.4 g L-1 throughout the mixed layer and year round, with primary production rates generally below 20 g C L-1 day-1. There is little seasonal variation in standing stock, though primary production rates sometimes ascend as high as 30-40 g C L-1 day-1 in later summer (Thibault et al., 1999). In contrast, the marginal station P04 (Fig. 1) shows a very pronounced seasonal cycle in both standing stocks and primary production, similar to the open ocean station in winter, but sometimes exceeding 2 g L-1 and 100 g C L-1 day-1 for chl-*a* and primary production, respectively, in August-September. Station P04 exhibits a sub-surface chlorophyll maximum (SCM, at around 15-30m) in late summer (Thibault et al., 1999; Boyd and Harrison, 1999).

The phytoplankton species composition also varies across Line P, both seasonally and spatially. The dominant cells, year round, found in the photic zone at P26 are *prymnesiophytes* and *pelagophytes*. In contrast, diatoms dominate in August-September at station P04 (Thibault et al., 1999). In the winter the phytoplankton population tends to be more diverse. Thibault et al. (1999) note that sediment traps recovered from 90-100m at station P26 contain more diatoms than photic zone cell populations would suggest, and conclude that sporadic aeolian Fe fertilisation events, leading to episodic diatom blooms, may not have been captured by their phytoplankton census. Boyd and Harrison (1999) present a similar picture, and conclude that diatoms are Fe-stressed in the open ocean NE Pacific but probably not at marginal locations, such as that of station P04.

**3. Sampling and analytical methods**

The new data presented here derive from two stations at either end of Line P (white circles on Fig. 1). The samples were collected during a cruise in August 2011. Samples derive from the open ocean station P26 (Ocean Station Papa, at 50°N 145°W), and the coastal station P04 (48.39°N 126.40°W). Rigorous contamination control procedures were used throughout sampling, with all critical handling steps onboard ship and onshore carried out in either a Class-100 laminar flow hood or a clean room. Samples were collected into acid-cleaned low-density polyethylene bottles. For depths ≤ 40m water was pumped onboard via a trace metal clean Teflon pump and filtered directly through in-line 0.2 μm polyethersulfone membrane filters. Samples from depth ≥ 40m were collected with acid-cleaned 12 litre Teflon-coated Go-Flo bottles attached to a powder-coated trace metal clean rosette. Upon recovery, seawater in the Go-Flo samplers was filtered under gravity through 0.2 μm polyethersulfone membrane filters. Unfiltered samples were analyzed for Si, NO3- and PO43- according to Barwell-Clarke and Whitney (1996). Dissolved O2 data was obtained with a Seabird SBE43 dissolved oxygen sensor calibrated against Winkler titrations. Nutrient and O2 data are courtesy of the Line P Program (Chief Scientist Marie Robert, Institute of Ocean Sciences, Canada). Water samples for chlorophyll analysis were filtered onto Whatman GF/F filters, and extracted with 90% acetone for fluorometric analysis (Strickland and Parsons, 1972).

The methods for isotopic analysis of the NE Pacific samples were identical to those we have described in extensive detail previously (Zhao et al., 2014). Briefly, samples were double-spiked prior to pre-concentration of Zn by co-precipitation with added aluminium hydroxide, column purification and analysis by MC-ICPMS. The blank for this process was 2-3x10-9g (ng) Zn, compared to a processed sample size of 50-500 ng, with the exception of 3 near-surface samples at station P04 (10-30 ng). All samples have had a correction applied for this blank (with a 66Zn value of 0.17±0.18 ‰), and this results in the high analytical uncertainties for the near-surface station P04 samples. The uncertainties reported for most other samples correspond to the analytical reproducibility of ±0.07 ‰ (2SD). Zn concentrations were obtained by isotope dilution. Uncertainties, based on duplicate analysis performed in this laboratory, are <5%.

**4. Results**

The new Zn concentration and isotope data are presented in Table 1, and shown as depth profiles in Fig’s 2 and 3, together with data for the major nutrients and chl-*a* obtained on the same cruise. Zinc concentration data are available for stations P04 and P26 from three previous studies (Martin et al., 1989; Lohan et al., 2002; Janssen and Cullen, 2015), and these data are also compared with those obtained here in Fig’s 2 and 3.

A key feature of station P04 (Fig. 2) that is apparent from all these studies is a particularly rapid increase in dissolved Zn concentrations from the very low values at the surface. In the dataset for both winter and summer 1999, this increase continues to at least 400m, where concentrations reach 14 nM. In 2011 the rapid increase in the upper 50m matches that for the 1999 data, but there is a maximum at 50-70m, after which concentrations decline again to levels close to the shallowest samples measured in August 2012. The prominent Zn concentration maximum is mirrored by a maximum in both Zn/Si (>0.6 mmol mol-1) and Zn/P (up to 6 mmol mol-1). It is important to note that these features of the upper 100m are defined by several data in the 2011 profile and by data from 1999. The maxima in [Zn], Zn/Si, and Zn/P correspond to a pronounced minimum in 66Zn, of -0.2‰. Above this minimum, three data points are heavier, and these occur at depths where chl-*a* is high, though shallower than the chl-*a* maximum. Beneath its minimum, 66Zn increases slowly with depth to the value of around +0.5‰ that characterises much of the deep ocean (e.g. Zhao et al., 2014; Conway and John, 2014b; John et al., 2018; Wang et al., 2019). The station P04 isotope profile generally matches that for the subtropical North Pacific station SAFe extremely well (Conway and John, 2015).

Though data for station P26 are also generally consistent with previous studies (Fig. 3), conditions at this location did differ in August 2011 in several important respects. For example, chl-*a* concentrations show a distinct maximum in the photic zone in August 2011, at levels greater than station P04, a feature that is not usual for this location (Thibault et al., 1999; Boyd and Harrison, 1999). Similarly Zn concentrations and Zn/P are somewhat higher right at the surface than found previously. Zinc concentrations, Zn/Si and Zn/P all show minima at the base of the mixed layer and a steep gradient down to about 150m. Zinc concentration and Zn/P continue to increase more slowly beneath this depth, while Zn/Si ratios show a maximum at 150m, decreasing slowly below. Zinc isotopes show a minimum similar to station P04, but at P26 this minimum is more pronounced (-0.4‰) and shallower (25m). As at station P04, 66Zn above this minimum is heavy, though there is only one datum, and beneath it there is the same slow increase to the deep ocean value of +0.5‰. The minimum in 66Zn occurs within the interval of high chl-*a* in the upper 50m. Beneath about 600m, the isotope data again match those for SAFe (Fig. 3), but the P26 profile increases to the value of +0.5‰ that characterises the deep ocean at a much shallower depth – 150-200m versus 600-800m.

**4. Results**

**5. Discussion**

**5.1 Data comparisons along Line P**

Fig. 4 compares data for the two stations studied here, at either end of Line P, with published data for stations in between (Lohan et al., 2002; Janssen and Cullen, 2015). On Fig. 4 the marginal station P04 clearly stands out, being characterized by very prominent excursions in [Zn], Zn/Si and Zn/P in the 50-100m depth range. As noted earlier, our findings here are consistent with the fact that diatoms dominate the phytoplankton at station P04 in summer (Thibault et al., 1999). For example, diatom cells are well-established to have very high Zn/P ratios (e.g. Twining and Baines, 2013) so that the particularly low dissolved ratios in the upper 20m at station P04 are consistent with uptake. The excursion to values as high as 6 mmol mol-1 at 50-100m, just below the mixed layer, is best explained by co-regeneration of Zn and P from this pool at close to the cellular ratio (Twining et al., 2014). The excursion to high Zn/Si at the same level is consistent with the above scenario, given the deeper levels for opal dissolution versus organic matter respiration.

High Zn/Si ratios in the upper subarctic NE Pacific generally were first pointed out by Lohan et al. (2002), who also noted a pattern of decreasing upper-ocean Zn/Si ratio with distance offshore. The depth profiles in Fig. 4 reinforce this pattern, but also emphasise that the enrichment in Zn is particularly prominent in the immediate sub-surface. This is striking at station P04 as noted above, but it also relatively obvious at station P12 (Fig. 4). On the other hand, by stations P20 and P26 further to the west it has largely disappeared. This spatial pattern is also consistent with previous work on the microbial ecology along Line P. Thibault et al. (1999) report that diatoms, though not dominant as at station P04, still constitute a significant proportion of the algal population at station P12. On the other hand, other algal groups (*prymnnesiophytes, pelagophytes*) dominate further offshore. We suggest that it is cycling of Zn via uptake into the organic matter of diatom cells, together with regeneration of this Zn in the immediate sub-surface, that maintains this very shallow sub-surface pattern in the more eastward stations along Line P.

At offshore stations (P16, P20, P26), Zn concentrations increase relatively slowly and monotonically with depth in the upper 100m of the water column. At all offshore stations concentrations then increase much more rapidly beneath 100m, reaching values close to their deep ocean maximum between 200 and 600m. Again there is a spatial pattern, with the deep ocean value reached at 200m at station P26 versus 600m at stations P04 and P12. This rapid change between the photic zone and the deep ocean, also noted earlier for Zn isotopes at P26, occurs at depths that closely correspond to the permanent halocline. At P26, beneath 100m, Zn is virtually conservative with salinity. The east-west pattern, we will show later, is actually radial, and is controlled by upwelling centred on the Alaska Gyre.

**5.2 Upper ocean controls on Zn isotopes in the North Pacific and beyond**

There is currently debate over what controls Zn isotopes in the upper ocean. Intense biological drawdown in the surface Southern Ocean appears not to be associated with a significant isotope effect (e.g. Zhao et al., 2014; Wang et al, 2019; Sieber et al., in review). Thus, surface water here, despite extreme depletion of Zn, is not isotopically different from deep water. As a result, all water masses exported from the Southern Ocean, despite Zn concentrations that differ by two orders of magnitude, are identical in 66Zn at close to +0.5‰. Outside the Southern Ocean, large datasets for Zn isotopes from the North Atlantic and South Pacific (Conway and John, 2014b; John et al., 2018) demonstrate that upper ocean Zn is up to 1.5‰ *lighter* than the deep ocean, not heavier as would be expected from *in-situ* biological drawdown. We suggest that these observations pose two different questions: (1) is biological cycling *within* the upper ocean regime inconsistent with culture studies that suggest uptake of the light isotope and; (2) why is the low latitude upper ocean *as a whole* isotopically light? The data in this paper are directly relevant to question (1) above. We return briefly to question (2) at the end of this section.

Culturing studies (John et al., 2007; John and Conway, 2014; Köbberich and Vance, 2017, 2018; Samanta et al., 2018) have demonstrated that a range of organisms take up the light isotopes of Zn relative to the bulk culture media. John et al. (2007) demonstrated that, at the concentrations relevant to the real ocean, the apparent fractionation is small, corresponding to a difference between the bulk medium and the cells, 66Znmedium-cells, of approximately +0.2 to +0.4‰. As originally noted by John et al. (2007), and reinforced by Köbberich and Vance (2019), this difference is actually consistent with no difference between the isotope composition of free Zn in the culture medium and that taken up, given that most medium Zn is complexed to EDTA and that EDTA-complexed Zn is 0.3 to 0.4‰ heavier than the free Zn species (Markovic et al., 2017). However, Zn in almost the entire global surface ocean (except for the Southern Ocean) is organically-complexed like the Zn in culturing media so that, since cells take up free Zn (Sunda and Huntsman, 1992), the cells should still be enriched in the light isotopes relative to the dissolved pool. Markovic et al. (2017) present experimental findings showing that isotope fractionation between free Zn and the organically-bound complex depends on the thermodynamic stability constant for that complex, which for EDTA is about 6 orders of magnitude greater than those for the molecules that chelate Zn in the ocean (*e.g.*, Bruland, 1989; Ellwood and van den Berg, 2000; Markovic et al., 2017). On the other hand, Bruland (1989) also showed that the *conditional* stability constant for Zn-EDTA complexes in seawater are lower than those for natural organic ligands, due to side reactions between EDTA and Ca and Mg ions that do not occur for the natural ligands (Bruland et al., 1989). Whatever the precise reason, it is clear that surface Zn isotopes barely shift in residual surface seawater north of the upwelling zone in the Southern Ocean, despite the uptake of about 95% of the upwelled Zn pool into diatom cells (Zhao et al., 2014; Wang et al., 2019).

In the upper 50 m or so of the sub-Antarctic zone of the Pacific, Zn is heavier than the sub-surface (Samnta et al., 2017). Moreover, variations in Zn isotopes in the upper 150 m correlate positively with chl-*a*, consistent with uptake of light Zn and its regeneration beneath the chlorophyll maximum. Haptophytes are the main phytoplankton group in this region. In a companion study (Samanta et al., 2018), these same authors demonstrated that cultured *Emiliana huxleyi* cells are enriched in the light isotope relative to bulk media, by about 0.4‰. The authors note the similarity between this apparent isotope fractionation and the isotopic differences in the real ocean across depth and variations in chlorophyll *a*. However, the isotopic spread in the water column does not equate to the fractionation factor for uptake seen in culture. In fact, the relationship between Zn concentrations and 66Zn for the water column at the studied sites would, for a closed-system Rayleigh scenario, imply only a very small fractionation – ≤ 0.1‰ – between bulk seawater Zn and that taken up.

To a large extent the pattern found by Samanta et al. (2017) is also seen here for the NE Pacific data, though the relationship between the location of 66Zn minima and maxima to those of chl-*a* are not as straightforward (Figs. 2, 3). In the case of both stations studied here, the surface-most 66Zn datum is heavy while the sub-surface is light. Though the surface return to heavy Zn isotopes is defined by a single datum for station P26, for station P04 it is better resolved. As discussed in section 5.1 all the evidence for station P04 points to the cycling of Zn in the upper 200m via uptake into and regeneration from diatom cells. Given the uncertainties and scatter in the data, the difference in 66Zn between the mixed layer and the immediate sub-surface is about 0.5-1‰. Again, however, this does not represent the degree to which the diatom cells take up light Zn – in fact, the relationship between Zn abundance and isotopes in the upper 200m at station P04 could imply preferential uptake of the light isotope, as for Samanta et al. (2017), by ≤ 0.1‰.

The finding of slightly light Zn isotopes in the immediate sub-surface relative to the surface is actually ubiquitous, was first made by Bermin et al. (2006), and keeps recurring (e.g. Zhao et al., 2014; Wang et al., 2019). In terms of the North Atlantic, where the most extreme case of an overall light upper ocean can be found, Wang et al. (2019) show that more than half the depth profiles presented in Conway and John (2014b) actually show a pattern whereby the surface-most data are heavier than the immediate sub-surface. Thus, the pattern of Zn isotope variation *within* the upper ocean is not necessarily inconsistent with culture studies

Regarding question (2) from earlier, explanations of the main processes driving the isotope composition of the tiny low latitude upper ocean pool have, naturally, focused on removal processes such as *in-situ* biologically-driven export or scavenging (e.g. Bermin et al., 2006; Zhao et al., 2014; Conway and John, 2014b; John et al., 2018). Since the isotope composition of the upper ocean is light overall, this has led to the suggestion that the removal process favours the heavy isotope (e.g. John and Conway, 2014; Conway and John, 2014b; Weber et al., 2018). However, these assumptions now need to be re-visited. The ultimate origin of low latitude water and the Zn it contains lies in subduction in the sub-Antarctic region. Zinc has largely been stripped from this water by diatom uptake and export. In this framework, the very low upper ocean Zn concentrations are the result of distal biological drawdown in the Southern Ocean – i.e., the upper ocean waters transported northwards from the Southern Ocean are already very low in Zn - the low Zn concentration of the low latitude upper ocean is a preformed signal. Thus, it is not at all clear whether any subsequent process involves removal or *addition*. Given the sensitivity of this very small upper ocean pool of Zn to a perturbation, *addition* of isotopically light Zn is also, actually, a viable option, one that needs to be more fully addressed in future work.

**5.3 Is there significant removal of Zn to sulphide in the mid-depth NE Pacific?**

In an analysis of variations in Zn concentrations and Zn/Si ratios, with depth and along Line P, Janssen and Cullen (2015) showed that Zn and Si concentrations are slightly decoupled in the NE Pacific in comparison with other parts of the world ocean. Specifically, these authors hypothesised the preferential removal of Zn from the water column as solid Zn sulphides form in particularly anoxic microenvironments around remineralising organic material, at depth levels where O2 falls below about 60M, in low oxygen zones. That there is a degree of Zn-Si decoupling in the NE Pacific is beyond doubt. The key data supporting data are plotted in Fig. 5, which shows the current as well as other data from Line P as well as data from other North Pacific sites in Fig. 1 (Martin et al., 1989; Lohan et al., 2002; Janssen and Cullen, 2015; Conway and John, 2015).

The observation that led Janssen and Cullen (2015) to suggest decoupling is the fact that data in the NE Pacific lie off and above the Zn-Si correlation as defined globally (e.g. the black line on Fig. 5A from Vance et al., 2017), so that the correlation has a distinct convex upwards nature. The suggestion in Janssen and Cullen (2015) is that Zn and Si data from the NE Pacific effectively lie on two different straight-line segments – one for the well-oxygenated upper ocean where Zn and Si increase together because of regeneration, and one for the low oxygen zone where Si concentrations continue to increase with depth but Zn concentrations level off due to removal into putative sulphidic particulates.

One issue with this interpretation is that, in fact, the Zn-Si correlation is never completely linear. Vance et al. (2017) emphasised the fact that the major Southern Ocean-sourced upper ocean water mass, Sub-Antarctic Mode Water (SAMW), has its origin in subduction from the mixed layer. This physical feature greatly diminishes the importance of the fact that the regeneration length scales of organic material versus opal are different (e.g. Berelson, 2001; Ragueneau et al., 2002). In simple terms, if diatom uptake and growth can remove Zn and Si together from the surface mixed layer, relative to PO4, the resulting low Zn/P, low Si/P signal will be imprinted on water masses subducted from that layer, whilst Zn- and Si-rich biogenic particles are exported into the deep Southern Ocean, where their decomposition and dissolution will lead to an enrichment in both Zn and Si. This coupled export outweighs the differences between the lengthscales of regeneration of organic matter (with the Zn) and the opal (with the Si), and produces a correlation between these two elements. However, even in the Southern Ocean the Zn-Si relationship is slightly convex upwards (Fig. 5A), and in fact the convex-upward nature of the NE Pacific relationship is simply a more pronounced version of the same thing.

This suggestion receives strong support through comparison of depth profiles of the major nutrients in the NE Pacific with that for excess Zn (Fig. 6). In Fig. 6, excess Zn relative to the global Zn-Si correlation is shown as Zn\* = Zn - Si × Zn/Si(deep), where the Zn/Si(deep) = 0.064 mmol mol-1 (Vance et al., 2017). Fig. 6A shows that there is indeed a reversal in Zn\* at about 300 m, at the same depth level at which O2 concentrations decrease to <100 mol kg-1. However, beneath that depth level Zn remains in excess: Zn\* is far above zero. The Zn\* peak at around 300m is a result of intense regeneration of Zn with PO4 (Fig. 6B) *above* that level, equivalent to the steep lower portion of the Zn-Si correlation, while opal dissolution is much more modest. The reversal in Zn\* at around 300m, equivalent to the break in slope in the Zn-Si correlation (Fig. 5A), is similarly driven by the deeper regeneration of silica from opal (Fig. 6C) accompanied by very little further increase in Zn (and PO4) through regeneration.

Thus, we suggest that the convex-upwards nature of the Zn-Si relationship is, mainly, a simple consequence of three important controls: (1) Zn in phytoplankton, including diatoms, is nearly exclusively located in organic material (e.g. Twining and Baines, 2013); (2) as a result Zn and PO4 are regenerated together, at the Zn/PO4 ratio of the cell, in the upper ocean (Twining et al., 2014); (3) the lengthscale of regeneration of opal is greater. These controls, while always a factor to some degree as shown by the Southern Ocean data in Fig. 5A, become more important in regions of the ocean, like the subarctic NE Pacific, where the combination of high particulate fluxes and limited ventilation of the shallow subsurface allows biogeochemical signals of local nutrient cycling to build up in the water column. Thus, the data that stand out on Fig. 5A are those for station P04, forming their own trend with very high Zn/Si ratio, which we attribute to the dominance of diatoms at this location, their very high Zn contents and the fact that this Zn is regenerated from organic material at very shallow levels. The high Zn/PO4 ratio of diatoms also make these data stand out on the Zn-PO4 correlation (Fig. 5B). Otherwise, the deep NE Pacific data simply continue the deep Zn-PO4 array – there appears to be very little room for removal of Zn relative to PO4 in the oxygen-deficient zone of the NE Pacific. Clearly, on the two plots in Fig. 5, it is silica concentrations in the deep Pacific that stand out: in other words, the cause of the convex-upwards nature of the Zn-Si correlation is not preferential removal of Zn in the sub-surface, rather it is preferential regeneration of Zn in the upper ocean and preferential dissolution of opal in the deep.

**5.4 Regional-scale features of the distributions of major nutrients and metal micronutrients in the NE Pacific**

The previous section highlighted the fact that decoupling of Zn and Si in the NE Pacific arises through differential remineralisation of Zn and Si with depth, with Zn sourced with phosphorus from sinking organic material in the upper ocean while Si is sourced from opal deeper in the water column. We contend that this is simply a manifestation of processes that occur elsewhere, but are more readily visible here because of relatively sluggish circulation, long sub-surface residence times (e.g. Östlund and Stuiver, 1980) and high particulate fluxes in a region of general upwelling (e.g. Honjo et al., 2008). We examine this further, as well as other aspects of the subarctic NE Pacific in particular, via a north-south sequence of stations from SAFe (35°N, Fig. 1; Conway and John, 2015) to T9 (58.5°N; Martin et al., 1989), using depth profiles (Fig. 7), plots against potential density (Fig. 8) and sections at specific depths (Fig. 9).

Many features of the depth profiles, going from north to south, coincide with those noted by Janssen and Cullen (2015) proceeding west along Line P. Thus, both westward along Line P (Fig. 4) and northwards from SAFe as far as station T8 (55.5°N) on Fig. 7, the Zn concentration data shift away from continuously increasing profiles with depth (station P04 versus P26, station SAFe versus T8) toward profiles with two distinct sections: first, rapidly increasing Zn concentrations down to 400m or so, followed by rather flat profiles versus depth. The Zn/P depth profiles mirror those of Zn concentrations almost perfectly, again emphasising the similarity in the depths of the remineralisation of these two elements. And, again, the contrast with Si is emphasised by the depth profile for Zn/Si, with a shoaling and sharpening of the Zn/Si peak (Fig. 7) proceeding northwards. It is important that the final station in the north-south sequence considered here – station T9 at 59.5°N, north of the centre of the Alaska gyre – shows a reversal in all these trends, so that its depth profiles are similar to those for station T6, at 45°N. Thus, the biogeochemical pattern of variation in the NE Pacific is, in fact, radially centred on the Alaska gyre, and is almost certainly related to upwelling. Indeed, the variation with depth at stations in the centre of the gyre look like classical advection-dominated profiles. Moreover, the variations in the shapes of depth profiles across line P (Fig. 4) are also seen in the north-south sequence (Fig. 7), but these differences largely disappear in the potential density plots (Fig. 8). The differences that remain between stations in Fig. 8 are also intriguing. For example, the two stations in the centre of the gyre (P26 and T8) show an enrichment in Zn relative to all other stations at low densities (~25.7–26.5 kg/m3) that correspond to the very shallowest subsurface water column (~50–150 m) at these stations. This feature – also visible in Zn/P and Zn/Si in Fig. 8 – is, again, extremely suggestive of a net source of Zn to sub-mixed layer waters, caused by enhanced regeneration of Zn within the upwelling Alaskan gyre. In this context, it is worth noting that Kim et al. (2017) invoke remineralisation of Zn-rich biogenic particles as a potential cause of the shallow Zn-enrichment observed in the western and central subarctic North Pacific.

These diagrams also show again that, in the context of variations in nutrient concentrations with depth, it is silica that stands out as different in its behaviour versus depth in the NE Pacific. Thus, for stations in the Alaska Gyre at 50-60°N, *both phosphate and the metals* Freach values within 10% of their maxima, at about 300-400m (Fig. 9). This is emphasised for Zn by the Zn/P ratio depth profile in Fig. 7. Clearly Zn, Cd, are remineralised with phosphate in the upper 400m. Silica, alone, shows a significant increase between 300 and 1000m, and a smaller increase between 1000 and 1500m (Fig. 9). The remineralisation of opal in the deep ocean is reflected in the fact that Si concentrations, in contrast to phosphate and the metals, almost double between 300-400m and 1500-2000m. The Zn/Si ratio havles at all stations, from its peak at 200-400m.

**6. Conclusions**

In this paper we have presented new Zn concentration and isotope data for two stations in the NE Pacific (Line P), and have compared the new data with Zn abundance patterns, and their relationship to the major nutrients, across the NE Pacific. The new Zn isotope data are consistent with a very small (<0.1‰) isotope fractionation during uptake in the photic zone, with preferential uptake of the light isotope, that is consistent with both Southern Ocean data (Zhao et al., 2014; Wang et al., 2019; Sieber et al., in review) and culture experiments (John et al., 2007; Samanta et al., 2018; Köbberich and Vance, 2019). The isotopically-light cellular Zn is regenerated at very shallow levels, resulting in high dissolved Zn concentrations and an isotope minimum at about 50-100m. Beneath this regeneration maximum 66Zn approaches the deep ocean average as seen elsewhere. These patterns are particularly clear at the marginal station (P04) where previous work (Thibault et al., 1999) have documented the dominance of diatoms in the phytoplankton ecology.

Combining the new data with published Zn and major nutrient concentrations from across the North Pacific, confirms the role of vertical biological cycling in controlling regional biogeochemistry in the North Pacific. Even in the North Pacific (c.f. Weber et al., 2018), however, and to an even greater extent in the Atlantic (Middag et al., 2019), these vertical processes are superimposed on the dominant global pattern that originates through advection northwards of Southern Ocean water masses (Vance et al., 2017; de Souza et al., 2018). In the North Pacific, the vertical cycling processes as well as the clear co-regeneration of Zn and phosphate from organic matter in the upper ocean and the deeper regeneration of Si, partially decouples Zn and Si covariation. Our interpretation greatly de-emphasises the role of low oxygen zones in driving this decoupling. Though we not that we cannot discount this latter process completely, our analysis suggests little need to invoke additional upper-ocean Zn removal other than by cellular uptake, either by removal into sulphides or any other abiotic process.

**Acknowledgements**

We thank the Captain and crew of the CCGS J.P. Tully and Chief Scientist and head of the Line P Time Series program Marie Robert for supporting this research. We are grateful to Dr. Christina Schallenberg for help with trace metal clean sampling and the Department of Fisheries and Ocean’s Canada for pigment and nutrient analyses on the August 2011 cruise. This research was supported by ETH Zürich and the Swiss National Science Foundation (SNF) through grant 200020-16590 (to DV), and by the European Union through a Marie Sklodowska-Curie Fellowship (to GFdS). MCL’s trace metal research is supported by NERC grant NE/N001125/1. JTC was supported by a Natural Sciences and Engineering Research Council (NSERC) Discovery Grant. We thank two anonymous reviewers for their comments, and editor Laura Robinson for very efficient editorial handling.

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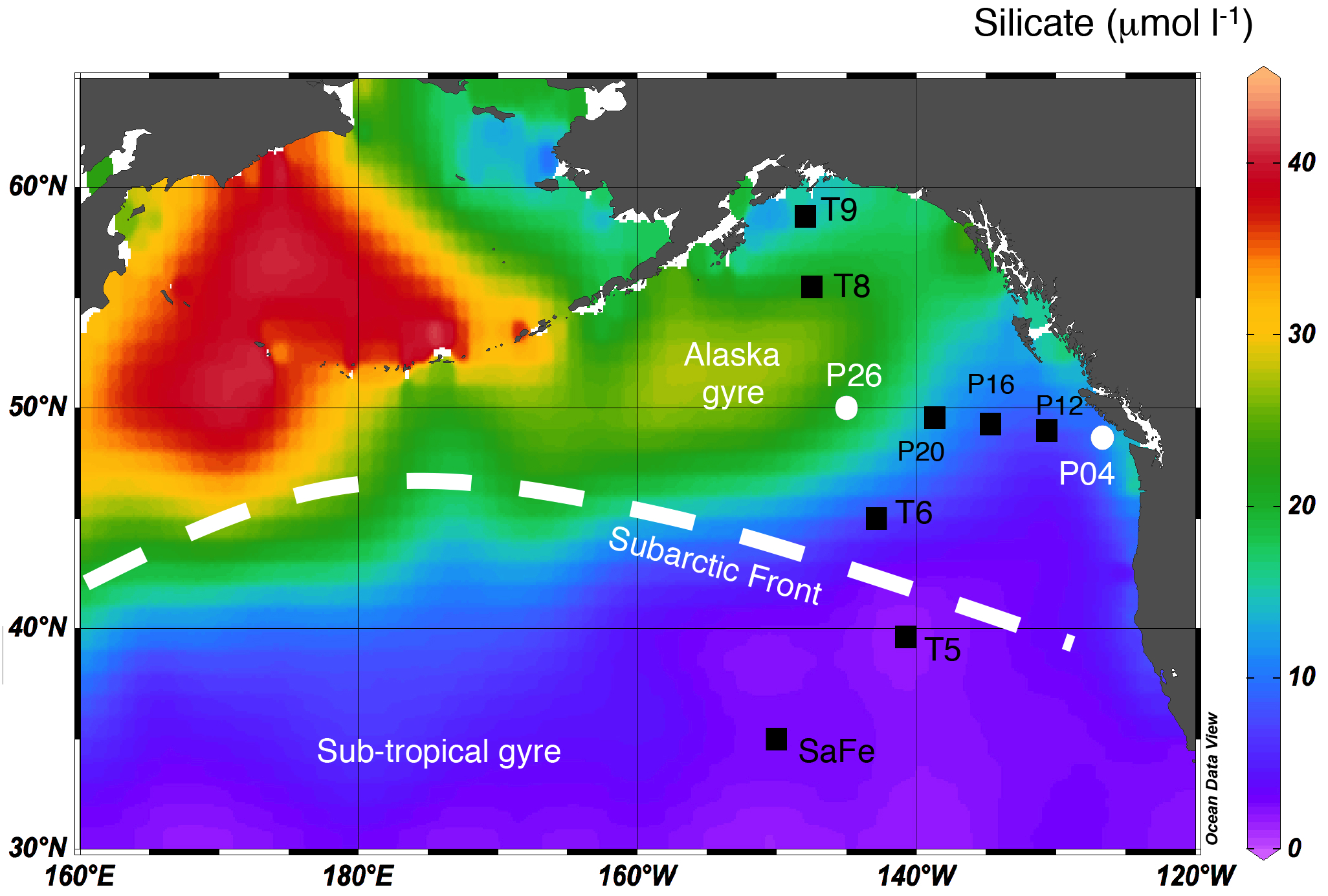
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**Table 1:** Zn concentration, isotopic and major nutrient data for the NE Pacific.

|  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| **Depth** |  | **[Zn]** | **66Zn1** | **2** |  | **Salinity** | **[PO4]** | **[Si]** | **[NO3]** |
| **m** |  | **nmol kg-1** | **‰** |  |  |  | **M** | **M** | **M** |
|  |  |  |  |  |  |  |  |  |  |
|  | ***P04, NE Pacific, 48.39°N, 126.40°W*** | | | | | | | | |
| 10 |  | 0.09 | 0.43 | 0.11 |  | 32.183 | 0.28 | 2.5 | 0 |
| 25 |  | 0.08 | 1.17 | 0.30 |  | 32.253 | 0.31 | 3.2 | 0 |
| 40 |  | 0.96 | 0.31 | 0.07 |  | 32.412 |  |  |  |
| 50 |  | 3.95 | -0.22 | 0.07 |  | 32.518 | 0.68 | 6.4 | 3.7 |
| 75 |  | 4.07 | -0.06 | 0.07 |  | 32.507 | 0.97 | 8.6 | 9.4 |
| 100 |  | 3.20 | 0.08 | 0.07 |  | 32.647 | 1.45 | 20 | 17.4 |
| 150 |  | 3.49 | 0.22 | 0.07 |  | 32.657 | 2.01 | 34.9 | 28.0 |
| 200 |  | 5.05 | 0.28 | 0.07 |  | 33.079 | 2.16 | 43.5 | 30.9 |
| 250 |  | 6.17 | 0.21 | 0.07 |  | 33.033 |  |  |  |
| 300 |  | 7.36 | 0.34 | 0.07 |  | 33.757 | 2.57 | 60.6 | 36.6 |
| 600 |  | 10.1 | 0.49 | 0.07 |  | 33.768 | 3.16 | 96.1 | 43.7 |
| 800 |  | 10.4 | 0.45 | 0.07 |  | 33.888 |  |  |  |
| 1000 |  | 9.11 | 0.52 | 0.07 |  | 33.891 | 3.28 | 127 | 45.1 |
| 1200 |  | 10.3 | 0.52 | 0.07 |  | 33.925 | 3.24 | 136 | 44.3 |
|  |  |  |  |  |  |  |  |  |  |
|  | ***P26, NE Pacific, 50°N, 145°W*** | | | | | | | | |
| 10 |  | 0.65 | 0.25 | 0.13 |  | 32.519 | 1.10 | 16.0 | 11.6 |
| 25 |  | 0.88 | -0.38 | 0.13 |  | 32.517 | 1.11 | 15.9 | 11.7 |
| 40 |  | 1.01 | 0.17 | 0.07 |  | 32.512 | 1.34 | 11.0 | 13.6 |
| 50 |  | 1.09 | 0.29 | 0.07 |  | 32.519 | 1.49 | 27.0 | 17.2 |
| 100 |  | 1.95 | 0.42 | 0.07 |  | 32.648 | 1.69 | 32.5 | 21.1 |
| 150 |  | 6.03 | 0.40 | 0.07 |  | 32.750 | 2.42 | 61.0 | 34.4 |
| 200 |  | 7.24 | 0.50 | 0.07 |  | 32.733 | 2.67 | 73.0 | 38.6 |
| 300 |  | 7.88 | 0.58 | 0.07 |  | 32.811 | 2.89 | 87.8 | 42.5 |
| 600 |  | 8.68 | 0.57 | 0.07 |  | 32.797 | 3.11 | 119 | 45.0 |
| 1000 |  | 9.75 | 0.49 | 0.07 |  | 33.680 | 3.18 | 147 | 45.8 |
| 1400 |  | 9.97 | 0.56 | 0.07 |  | 33.675 | 3.18 | 160 | 45.0 |
| 2000 |  | 9.83 | 0.49 | 0.07 |  | 33.773 | 3.01 | 171 | 43.8 |
|  |  |  |  |  |  |  |  |  |  |

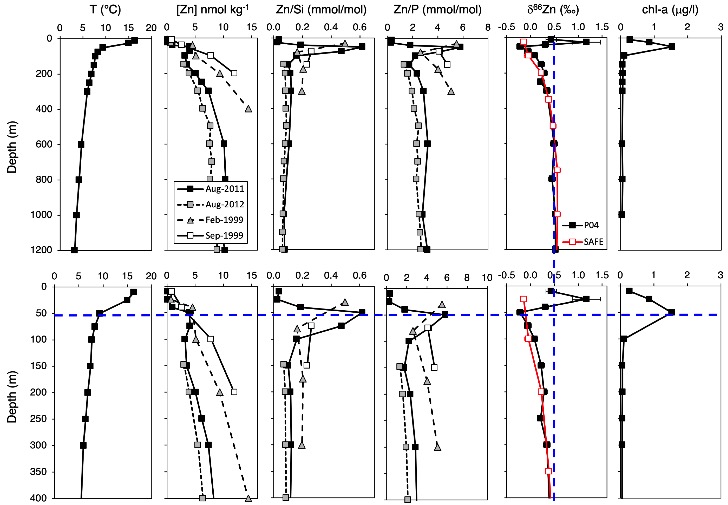
166Znreported as per mil (‰)deviations of the 66Zn/64Zn ratio from the Lyons JMC standard = [(66Zn/64Znsample/66Zn/64ZnJMC-Lyons)-1]x1000. Uncertainties quoted are the larger of the two estimates of uncertainty: external reproducibility (see Methods) versus internal error.

**Vance et al. – Figure 1**

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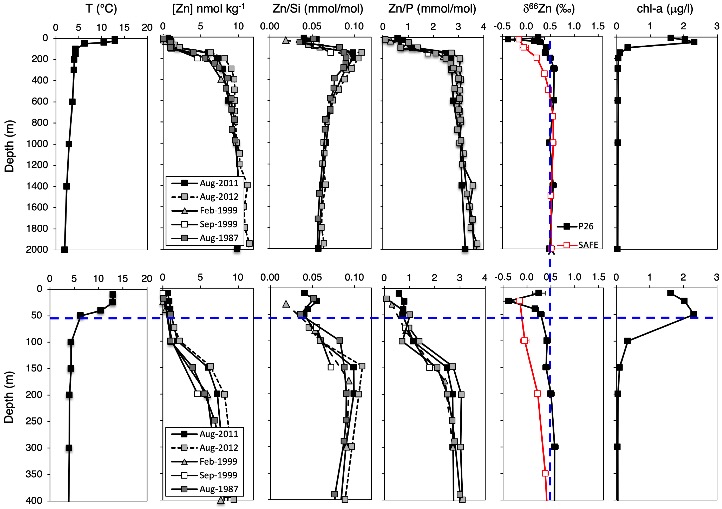
**Figure 1:** Map of the northeast Pacific, showing surface annually-averaged dissolved silicate concentrations (World Ocean Atlas; Boyer et al., 2013), with the locations of the two stations for which new Zn concentration and isotope data are presented here (white circles, stations P04 and P26). Other stations for which published data are discussed here are shown as black squares (SaFe: Conway and John, 2015; VERTEX stations T5-T9; Martin et al., 1989). Previously published data for Zn and major nutrient concentrations are also available for stations P04, P26 and other line P stations (Lohan et al., 2002; Janssen and Cullen, 2015) and are also discussed here. The approximate position of the northern boundary of the Subarctic Front (white dashed line) is taken from Yuan and Talley (1996).

**Vance et al. – Figure 2**

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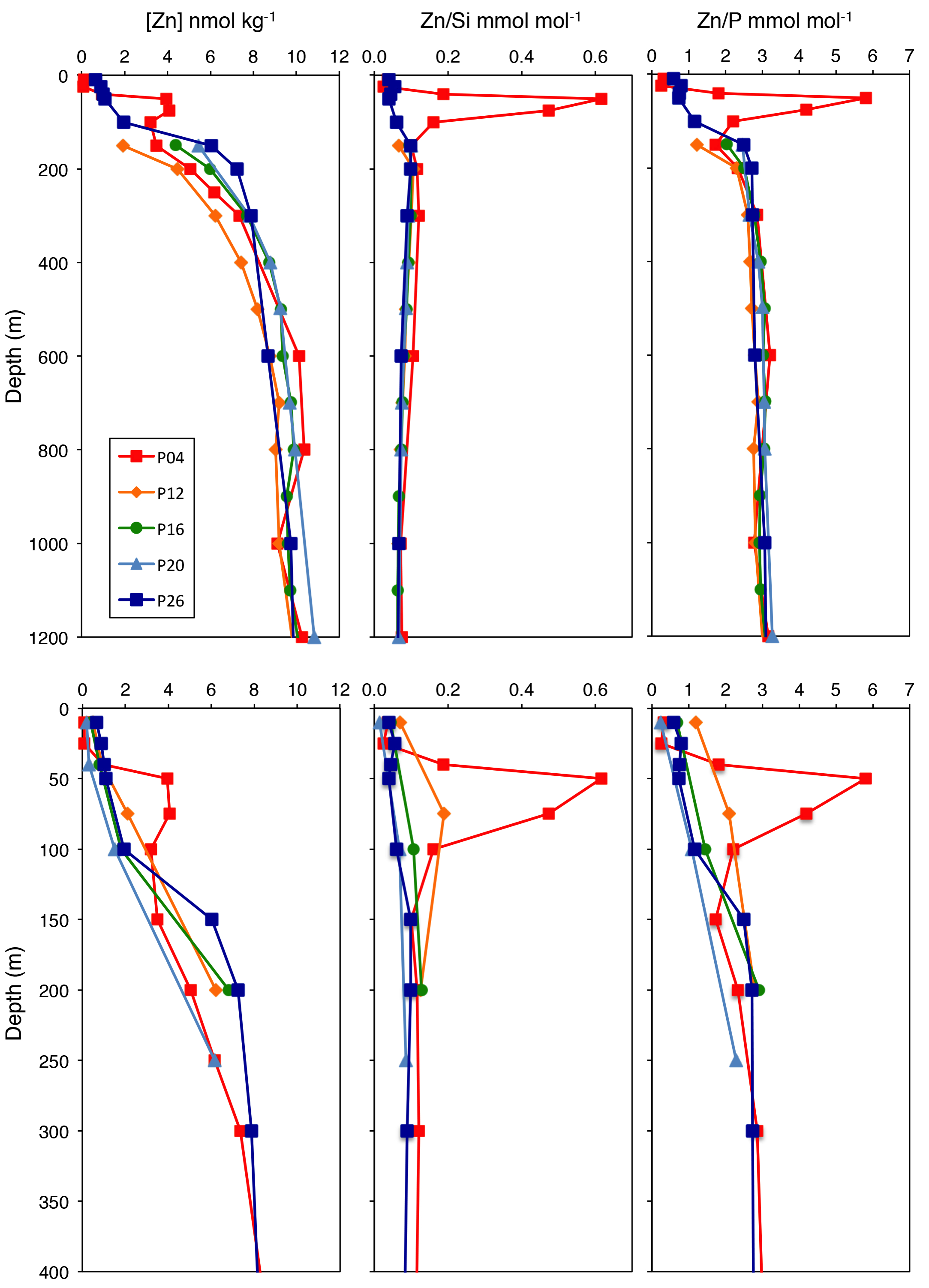
**Figure 2:** Depth profiles of temperature, Zn concentration, Zn/Si, Zn/P, 66Zn and fluorescence-based chl-*a* for station P04 for August 2011 (black filled squares in all panels). The full profiles are shown at the top and a more detailed view of the upper 400m at the bottom. The vertical blue line in the isotope plots marks a value of +0.5‰, typical of the deep Pacific and close to the average deep ocean value for other regions of the ocean (Zhao et al., 2014; Conway and John, 2014b; Conway and John, 2015; Samanta et al., 2017; John et al., 2018). The horizontal blue line marks the level of the chlorophyll maximum as derived from calibrated fluorescence measurements. The Zn, Zn/Si and Zn/P profiles show data from this study (August 2011) as well as data from previously published studies (Aug 2012: Janssen and Cullen, 2015; Feb. and Aug. 1999: Lohan et al., 2002). The isotope profiles also show data for the SAFe station (Conway and John, 2015).

**Vance et al. – Figure 3**

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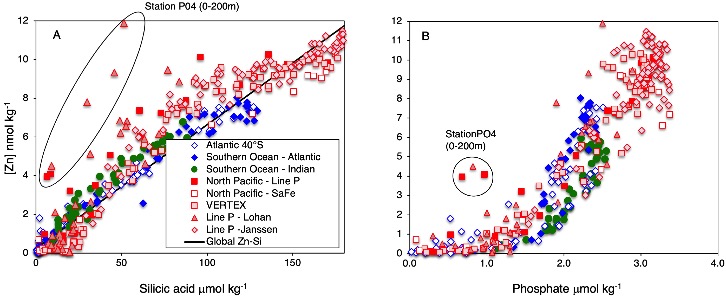
**Figure 3:** Depth profiles of temperature, Zn concentration, Zn/Si, Zn/P, 66Zn and chl-*a* for station P26 for August 2011 (black filled squares in all panels). The full profiles are shown at the top and a more detailed view of the upper 400m at the bottom. The vertical blue line in the isotope plots marks a value of +0.5‰, typical of the deep Pacific and close to the average deep ocean value for other regions of the ocean (Zhao et al., 2014; Conway and John, 2014b; Conway and John, 2015; Samanta et al., 2017; John et al., 2018). The horizontal blue line marks the level of the chlorophyll maximum as derived from calibrated fluorescence measurements. The Zn, Zn/Si and Zn/P profiles show data from this study (August 2011) as well as data from previously published studies (Aug 2012: Janssen and Cullen, 2015; Feb. and Aug. 1999: Lohan et al., 2002; Aug. 1987: Martin et al., 1989). The isotope profiles show data for the SAFe station (Conway and John, 2015).

**Vance et al. – Figure 4**

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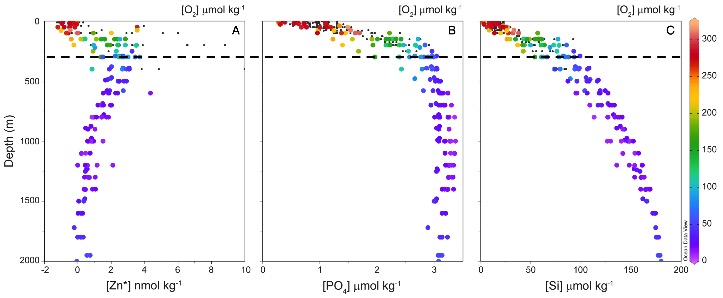
**Figure 4:** Depth profiles of Zn concentration, Zn/Si and Zn/P, comparing data from this study for stations P04 and P26 for August 2011 with published data for other Line P stations. The locations of all stations plotted are shown on Fig. 1. The full profiles are shown at the top and a more detailed view of the upper 400m at the bottom. For the top panels the data obtained here are compared with those for August 2012 in Janssen and Cullen (2015). No data are reported in the latter paper for the upper 150m so that the expanded profiles at the bottom use the data of Lohan et al. (2002) for September 1999.

**Vance et al. – Figure 5**

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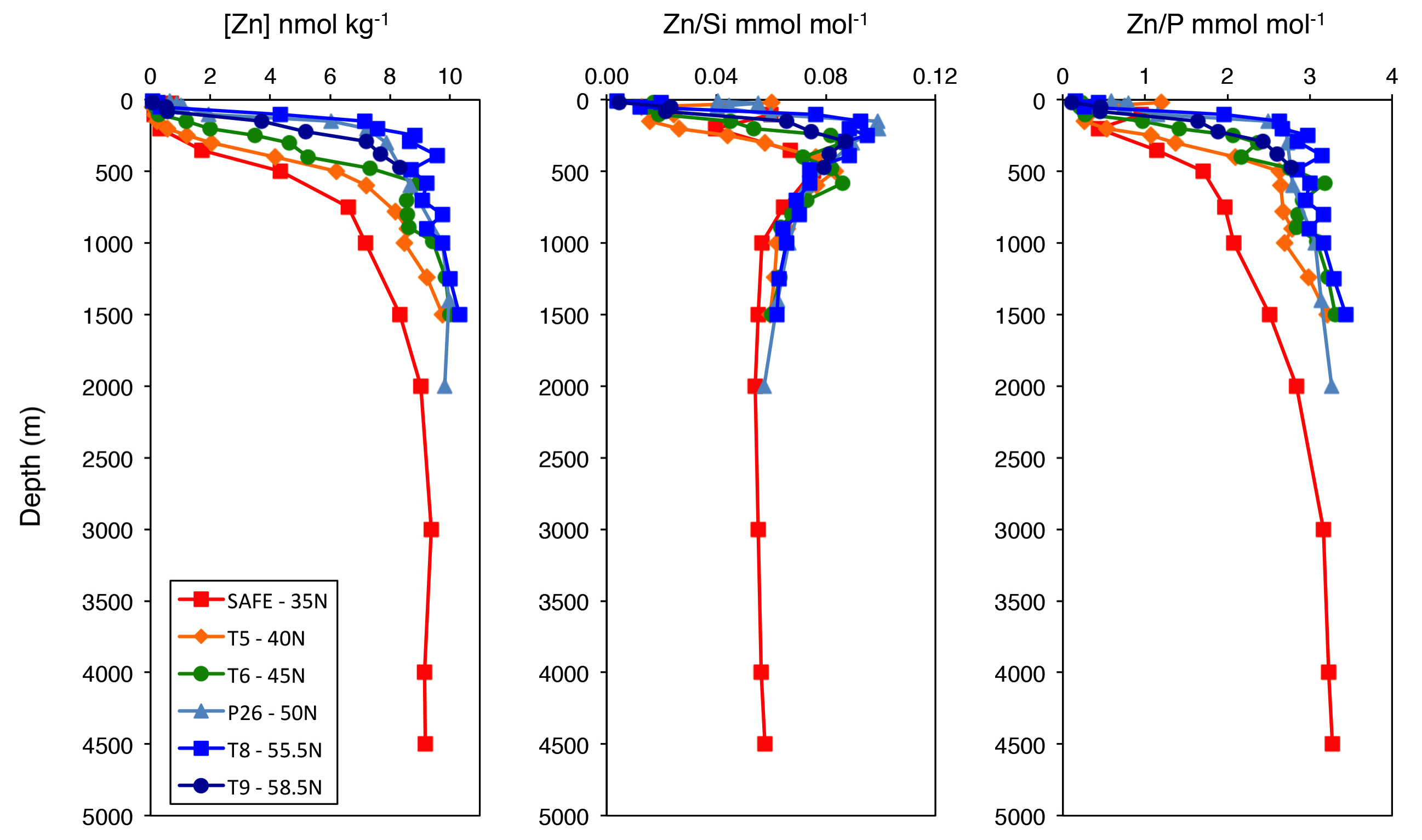
**Figure 5:** Zn versus silica (A) and Zn versus phosphate (B) for water column samples from the NE Pacific (this study: North Pacific – Line P; other Line P data from Lohan et al. (2002) and Janssen and Cullen (2015); SAFe (Conway and John, 2015); VERTEX (Martin et al, 1989)) compared to data for the Southern Ocean (Atlantic data from Zhao et al., 2014; Indian data from Wang et al., 2019) and the South Atlantic (Wyatt et al., 2014). Data for specific samples at Station P04, from this study and from Lohan et al. (2002) are highlighted. The line labelled Global Zn-Si is from Vance et al. (2017).

**Vance et al. – Figure 6**

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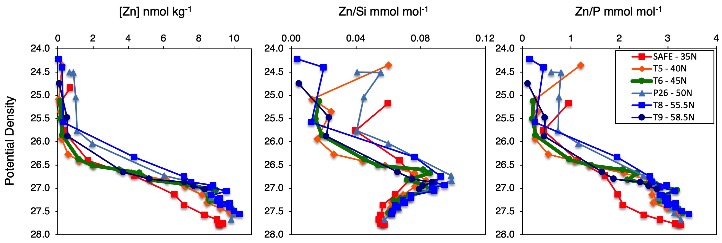
**Figure 6:** Depth profiles of Zn\* (A, see text for definition), PO4 (B) and Si (C) for the NE Pacific. Colour of the data points denotes dissolved O2 concentrations – scale on right – where available (otherwise data shown as black dots). The horizontal dashed line shows the depth level of the Zn\* high. This peak at around 300m is due to: (1) co-regeneration of Zn and PO4 from phytoplankton organic matter above and; (2) continued opal dissolution below (C) but minimal further regeneration of Zn and PO4 (B, Figs. 2-4).

**Vance et al. – Figure 7**

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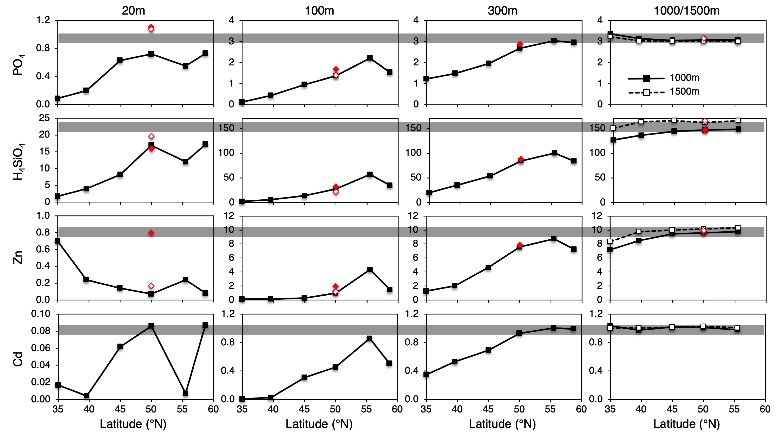
**Figure 7:** Depth profiles of Zn concentration, Zn/Si and Zn/P, showing data along a latitudinal transect through the North Pacific from SAFe (35°N; Conway and John, 2015), from VERTEX stations (Martin et al., 1989) and from this study (P26).

**Vance et al. – Figure 8**

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**Figure 8:** Zn concentration, Zn/Si and Zn/P, versus potential density, showing the same stations and data from the same sources as in the depth profiles in Fig. 7.

**Vance et al. – Figure 9**

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**Figure 9:** The same data as in Figures 7 and 8, with latitudinal trends shown for specific depths. Note the change in scales between the plots for 20m versus the plots for the deeper samples. The squares use data from Conway and John (2015) for SAFe and data from the VERTEX stations for all other latitudes (Martin et al. 1989). Data for P26 from this study (filled red diamonds) and from Lohan et al. (2002) illustrate the range of variation at this station, when available. The data for Zn concentration at P26 again highlight the different nature of the August 2011 dataset, as pointed out in the text for chlorophyll a and other parameters. The grey bands, showing the concentration at 1000m for all elements in all panels, highlight the fact that concentrations of all species except Si reach values very close (within 10%) to their maxima at about 300-400m. Silica concentrations, by contrast, increase by a further 70-90% at depths beneath 300-400m.