Thorium isotopes tracing the iron cycle at the Hawaii Ocean Time-series Station ALOHA
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Abstract
The role of iron as a limiting micronutrient motivates an effort to understand the supply and removal of lithogenic trace metals in the ocean. The long-lived thorium isotopes ($^{232}$Th and $^{230}$Th) in seawater can be used to quantify the input of lithogenic metals attributable to the partial dissolution of aerosol dust. Thus, Th can help in disentangling the Fe cycle by providing an estimate of its ultimate supply and turnover rate. Here we present time-series (1994-2014) data on thorium isotopes and iron concentrations in seawater from the Hawaii Ocean Time-series Station ALOHA. By comparing Th-based dissolved Fe fluxes with measured dissolved Fe inventories, we derive Fe residence times of 6-12 months for the surface ocean. Therefore, Fe inventories in the surface ocean are sensitive to seasonal changes in dust input. Ultrafiltration results further reveal that Th has a much lower colloidal content than Fe does, despite a common source. On this basis, we suggest Fe colloids may be predominantly organic in composition, at least at Station ALOHA. In the deep ocean (>2 km), Fe approaches a solubility limit while Th, surprisingly, is continually leached from lithogenic particles. This distinction has implications for the relevance of Fe ligand availability in the deep ocean, but also suggests Th is not a good tracer for Fe in deep waters. While uncovering divergent behavior of these elements in the water column, this study finds that dissolved Th flux is a suitable proxy for the supply of Fe from dust in the remote surface ocean.
1. Introduction

Determination of the supplies of iron to the ocean is relevant to understanding Earth’s climate and the ocean’s ecology. Ocean storage of carbon dioxide is mediated by iron supply in large areas of the ocean where Fe is a limiting resource, both today (Moore et al., 2013) and during the ice ages (Martínez-García et al., 2014). Additionally, the marine distribution of diazotrophic phytoplankton that modulate the nitrogen cycle may be determined by Fe supply rates (Ward et al., 2013). Atmospheric dust is arguably the major source of Fe to the euphotic zone (Boyd et al., 2010; Conway and John, 2014; Jickells et al., 2005; Tagliabue et al., 2014).

Debate on the sources of marine Fe ensues largely because the techniques to estimate the supply rate of Fe from dust in particular, or Fe residence times in general, are only beginning to be developed.

In this study, we assess the utility of thorium isotopes in seawater to provide rate information on the Fe cycle. By pairing $^{232}$Th, sourced from dust, with radiogenic $^{230}$Th (or $^{234}$Th) that provides a timescale of thorium flux, one can make quantitative estimates of both the total dust flux to the ocean (Deng et al., 2014; Hsieh et al., 2011) and the flux of trace metals released by dust dissolution (Hayes et al., 2013a). Our study site is the Hawaii Ocean Time-series Station ALOHA (22° 45’ N, 158° W) (Church et al., 2013; Karl and Lukas, 1996) in the subtropical North Pacific, where Asian dust is deposited in spring (Boyle et al., 2005; Hyslop et al., 2013; Prospero et al., 2003). Presenting time-series data spanning 20 yrs (1994-2014), we demonstrate that the behaviors of Fe and Th in seawater are consistent with a variable dust source to the surface ocean. Thorium-based fluxes indicate that the residence time of dissolved Fe in the upper 125 m of the water column is less than one year. In the sub-surface ocean (>250
m), the thorium and iron cycles begin to diverge considerably. These divergences reveal new insights into the marine geochemistry of these elements.

2. Background

2.1 Finding the timescale: thorium removal

The rate information on trace metal cycling that we seek is obtained by exploiting the natural radioactive disequilibrium between insoluble $^{230}$Th and its soluble parent $^{234}$U in seawater. The oceanic distribution of $^{234}$U (half-life 245,620 yrs (Cheng et al., 2013)) is homogeneous within a few parts per thousand, as $^{238}$U concentrations vary only with salinity (Owens et al., 2011) and $^{234}$U/$^{238}$U ratios vary by less than 1 per mil (Andersen et al., 2010). Therefore, the decay of $^{234}$U produces $^{230}$Th at a known rate everywhere in the ocean. Due to its particle reactivity, thorium adsorbs onto sinking particulate matter, a process called scavenging, on a timescale of years to decades, much faster than $^{230}$Th decay (half-life 75,584 yrs (Cheng et al., 2013)).

Thus by comparing the amount of $^{230}$Th that remains in seawater to the amount produced by U decay, one can calculate a removal timescale (Eq. 1, Fig. 1), or residence time ($\tau$), of thorium in seawater. Equation 1 is written with radionuclide concentrations in terms of activity (decays per unit time per unit seawater). The denominator contains a $^{230}$Th term to account for $^{230}$Th-decay, which, as mentioned above, can be neglected on the timescale of water column processes (for instance, seawater $^{234}$U activity is 46.6 mBq/kg at salinity 35, while typical $^{230}$Th activities are 1-10 µBq/kg). This approach is analogous to that used with a more commonly used flux tracer, the shorter-lived $^{234}$Th (half-life 24.1 days). By the same principles, using its production rate from parent isotope $^{238}$U, $^{234}$Th inventories can also be used to determine the scavenging rate of Th in seawater (Buesseler et al., 1992; Coale and Bruland, 1985), except that
234\textsuperscript{Th}-decay is much more significant in the water column balance (238\textsuperscript{U} activities being ~40 mBq/kg compared to euphotic zone 234\textsuperscript{Th} activities of 30-40 mBq/kg).

$$\tau_{Th}(z) = \frac{\int_0^z 230\textsuperscript{Th} dz}{\int_0^z (234\textsuperscript{U} - 230\textsuperscript{Th}) \lambda_{230} dz} \quad \text{Eq. 1}$$

To meet the requirements of a steady-state assumption between source and removal terms, we calculate thorium residence times in an integrated sense, from the surface to a particular depth. Thus as one integrates deeper into the water column, the 230\textsuperscript{Th} inventories reflect longer timescales of removal. Residence times calculated in this way also neglect dispersal fluxes by ocean circulation. Lateral gradients in oceanic 230\textsuperscript{Th} concentrations are generally small (Hayes et al., 2015a), while large vertical gradients may make vertical fluxes significant, for instance due to upwelling (Luo et al., 1995).

2.2 Finding the source: lithogenic metal fluxes

The dominant isotope of seawater thorium is primordial and long-lived (half-life 14.1 x 10^9 yrs) 232\textsuperscript{Th}. It is added to the ocean only through the partial dissolution of continental material, which in the context of Station ALOHA we consider to be primarily aerosol dust. Once in the water column, 232\textsuperscript{Th} is assumed to undergo scavenging removal (Fig. 1) at the same rate, i.e. with the same residence time, as 230\textsuperscript{Th} (or 234\textsuperscript{Th}), as scavenging tendencies are characteristic of all isotopes of an element. Assuming a steady state for Th concentrations, with knowledge of the Th residence time derived from 230\textsuperscript{Th}, one can calculate the flux of dust-derived 232\textsuperscript{Th} necessary to support the observed 232\textsuperscript{Th} inventory (Eq. 2). As in calculating thorium residence times, the derived dissolved 232\textsuperscript{Th} flux is reflective of the integrated depth zone, rather than at a particular depth. More details on 232\textsuperscript{Th} flux calculations are reported by Hayes et al. (2013a).

$$232\textsuperscript{Th} \text{ flux}(z) = \frac{\int_0^z 232\textsuperscript{Th} dz}{\tau_{Th}(z)} \quad \text{Eq. 2}$$
In comparison to the relative simplicity of the supply and removal terms in the thorium cycle, seawater iron cycling has many more terms to consider. These include biological uptake, remineralization, redox chemistry, anthropogenic or hydrothermal sources, in addition to supply by dust and removal by scavenging (Fig. 1). Scavenging of Fe also occurs but at a different rate than that of Th. The utility of this element pair is their common source from dust. This is, of course, an idealization of the “simple” thorium cycle. Th will be involved to some extent with (perhaps inadvertent) uptake into and remineralization from organic matter (Barbeau et al., 2001; Hirose and Tanoue, 1994). We have also observed a strong hydrothermal sink for Th in the Atlantic (Hayes et al., 2015a) in addition to an abyssal source of Th from sediments in the North Pacific (Hayes et al., 2013a). Nonetheless, particularly in the remote surface ocean well above the seafloor, dust dissolution and scavenging appear to be the dominant terms for Th cycling.

Thus, in this manuscript we propose using dissolved $^{232}$Th flux as a proxy for the Fe released during dust dissolution. This can be done with knowledge of the Fe/Th ratio in the dust and the relative fractional solubility of the two elements ($S_{Fe/Th}$, Eq. 3).

$$\text{Dust-dissolved Fe flux} = \text{dissolved } ^{232}\text{Th flux} \times (\text{Fe/Th})_{dust} \times S_{Fe/Th} \quad \text{Eq. 3}$$

By weight, the Asian desert dust that undergoes long-range transport over the North Pacific contains $^{232}$Th at 14.3 ± 0.8 ppm, based on fine grained (<8 µm) source materials (McGee, 2009; Serno et al., 2014), and Fe at 3.8 ± 0.4 %, based on a literature compilation by Mahowald et al. (2005). Therefore, we assume the Fe/Th ratio in dust at Station ALOHA of 2660 ± 320 g/g or 11,040 ± 1450 mol/mol. These ratios are close to the average for the upper continental crust of Fe/Th = 3271 g/g = 13,553 mol/mol (Taylor and McLennan, 1995).

The relative fractional solubility of Fe and Th in dust is currently unconstrained. Hayes et al. (2013a) assumed $S_{Fe/Th} = 1$ as a starting point, based solely on the similarly insoluble nature
of these two elements in seawater. While much more work is needed to constrain this parameter, here we continue to assume $S_{\text{Fe/Th}} = 1$, and our observations of the time-series variability in the seawater $\text{Fe}^{232}/\text{Th}$ ratio (section 4.4) support this assumption.

2.3 Iron residence times

We cannot rule out significant marine Fe sources by anthropogenic (e.g., derived from fossil fuel combustion) aerosols, continental margin sediments, or deep-sea hydrothermal vents. We can, however, pursue the notion that if dust were the only Fe source to the water column, the comparison between measured dissolved Fe inventories to the source (dust-dissolved Fe flux) would produce a measure of the turnover rate or residence time of dissolved Fe in seawater (Eq. 4). This residence time again represents the residence time within the integrated water column. Additional sources of Fe, such as combustion aerosols or hydrothermal fluids, would cause the dust-based Fe residence time to be an overestimate. Relevant to iron cycling, this residence time provides a rough timescale over which one can expect Fe concentration to vary as a result of variation in sources, such as springtime Asian dust events (Boyle et al., 2005).

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\text{Dissolved Fe residence time} = \frac{\text{Fe inventory}}{\text{dust-dissolved Fe flux}} \quad \text{Eq. 4}
\]

3. Materials and Methods

3.1 Sample collection during 2012-2014

Samples were collected on several cruises on the R/V Kilo Moana, led by the Center for Microbial Oceanography: Research and Education (C-MORE), to Station ALOHA in July-September 2012 (HOE-DYLAN), May-June 2013 (HOE-PhoR-I), September 2013 (HOE-PhoR-II) and March 2014 (HOE-BOE-I). Depth profiles for $^{230}\text{Th}^{232}\text{Th}$ analysis were collected from
the ship’s Niskin bottle rosette, filtered with a 0.45 µm Acropak cartridge filter, and acidified to pH 1.8 with Savillex-distilled 6 M HCl.

Filtered surface seawater (0.4 µm) was collected for $^{232}$Th (which requires smaller volumes than for $^{230}$Th), as well as for dissolved Fe, using the trace-metal clean MITESS sampler (Bell et al., 2002) at near daily time intervals on the 2012-2013 C-MORE cruises. MITESS collection methods, including “Vane” sampling for Fe depth profiles, on the HOE campaigns are discussed fully by Fitzsimmons et al. (in review). Within 3 hours of collection, the seawater was filtered using 0.4 µm polycarbonate track etch filters (PCTE, Whatman). Particulate samples were immediately frozen, and dissolved filtrates were acidified to pH 2 with trace metal clean HCl. The filters used for filtering MITESS water were analyzed for particulate Fe and $^{232}$Th (representing on average 0.7 liters of seawater).

On HOE-PhoR-II, cross flow filtration was performed to assess colloidal $^{232}$Th/$^{230}$Th using protocols developed to study colloidal Fe (Fitzsimmons and Boyle, 2014a). Seawater was pre-filtered at 0.45 µm and, within 1-2 hours, pumped over a Millipore Pellicon XL filter made of regenerated cellulose with a nominal molecular weight cutoff of 10 kDa, roughly equivalent to an effective pore size of 10 nanometers. Both permeate and retentate fractions were analyzed to determine any loss of Th by adsorption, which turned out to be minimal (88-100% dissolved Th recovery).

3.2 Hawaii Ocean Time-series (HOT) seawater

Seawater samples, typically 0.5 liter size, have been collected during the HOT program for trace metal analysis at MIT periodically since 1997. Most of these samples were collected as unfiltered water using the MITESS sampler (Bell et al., 2002) and subsequently preserved by acidification to pH 2 with HCl. We report data from samples collected via ship-based MITESS
collections as well as MITESS units deployed on a mooring (1997-2000, 2004-2005). In shipboard sampling, a sample bottle filled with high-purity dilute (~0.001 M) HCl is lowered over the side on a clean-wire and opened at depth, allowing ~15-20 minutes for the bottle to be completely flushed with the denser seawater, before bottle closure and sample retrieval. The moored sampling worked similarly except that bottles were filled with stronger acid (1 M HCl) prior to sampling. The moored sampling potentially posed a metal contamination risk due to the effective stronger leaching of the HDPE bottles, increasing the chance for leached Th, for example, to remain in the sample bottle at the time of collection. We report the moored sampler data with a unique symbol in our figures and interpret them with caution. Further sampling details are given by Boyle et al. (2005).

We also make use of literature seawater $^{232}$Th/$^{230}$Th data, collected at Station ALOHA in September 1994 (HOT-57), reported by Roy-Barman et al. (1996).

### 3.3 Thorium and iron analyses

Dissolved $^{230}$Th concentrations at Station ALOHA are as low as $10^{-18}$ moles per kilogram seawater ($10^{-18}$ mol $^{230}$Th $= 0.1746 \mu$Bq). Therefore, for measurement by inductively-coupled plasma mass spectrometry (ICP-MS), 4-5 liter water samples are required. Thorium concentrations were determined by isotope dilution by spiking with $^{229}$Th (not present in natural seawater). Sample preparation (pre-concentration, acid digestion, and chromatographic purification) was performed using published methods (Anderson et al., 2012; Auro et al., 2012). A portion of the $^{230}$Th samples were prepared and analyzed at the Lamont-Doherty Earth Observatory (L-DEO), using an Element XR single-collector ICP-MS. The remaining $^{230}$Th samples were prepared at the Massachusetts Institute of Technology (MIT) and analyzed using a
Neptune Plus multi-collector ICP-MS at Brown University. Th-232 was also analyzed in samples prepared for $^{230}$Th.

Analysis of $^{232}$Th, at $10^{-15}$ mol (i.e. femtomoles) per kg seawater, required smaller samples (200-800 mL) and was measured on archive HOT and HOE samples for which sample volume did not allow $^{230}$Th determination. While not as prone to contamination as some other trace elements, clean lab techniques were required to produce blanks that were consistent and low enough to allow detection of the relatively small sample size of ~20-40 femtomoles $^{232}$Th. Therefore, modifications of the cited procedures for Th analysis (Anderson et al., 2012; Auro et al., 2012) were made. Instead of co-precipitation with added Fe, pre-concentration of $^{232}$Th was achieved using magnesium hydroxide co-precipitation, such as that described for Pb by Reuer et al. (2003). Thorium was purified using a smaller amount (100 µl rather than 1 ml) of anion-exchange resin (AG1-X8) on columns fashioned from Teflon shrink-tubing. Samples were loaded onto AG1-X8 resin in 8 M HNO$_3$ and Th was eluted with 6 M HCl (instead of 12 M HCl, to reduce acid blank), following Edwards et al. (1987). Blank determinations were made on 125 mL aliquots of acidified seawater samples whose $^{232}$Th content had been determined during previous $^{230}$Th analysis. The mean procedural blank ($n = 6$) was $3.5 \pm 1.6$ fmol $^{232}$Th, resulting in a detection limit of 4.8 fmol $^{232}$Th. Samples for seawater $^{232}$Th were prepared and analyzed at MIT, using a Micromass IsoProbe multi-collector ICP-MS with detection by a Daly-style ion counter.

In this study, we refer to measured trace metal concentrations as dissolved (filtered at 0.4 or 0.45 µm), particulate (>0.4 µm), or total (acidified unfiltered water). The “total” concentrations in this sense are sometimes referred to as “total dissolvable”, allowing for the possibility that some forms of Th are not mobilized into solution by acidification to pH 2 or
collected with co-precipitation. Since our goal in interpreting seawater $^{230}$Th concentrations is to determine scavenging rates based on uranium decay, we made small (0-10%) corrections for the dissolved $^{230}$Th released from dust (or lithogenic material in general). This correction is based on measured dissolved $^{232}$Th and a lithogenic $^{230}$Th/$^{232}$Th mole ratio of $4 \times 10^{-6}$ (Roy-Barman et al., 2009). The corrected dissolved $^{230}$Th values are denoted as “xs”.

Particulate $^{232}$Th and particulate Fe, were analyzed at Florida State University by total digestion of the filter samples and subsequent analysis by ICP-MS, using slightly modified versions of published protocols (Ho et al., 2011; Morton et al., 2013; Upadhyay et al., 2009). In brief, samples were microwaved (CEM MARS Xpress) for 40 minutes at 180°C with HNO$_3$ and H$_2$O$_2$ (to digest the organic and less refractory biogenic and authigenic components) and HF (to digest the more refractory lithogenic components). The detection limit (based on 3 standard deviations of the digested acid blanks) for particulate $^{232}$Th was 8 fmol/L (n=19) and the particulate Fe detection limit was 0.2 nmol/L (n=21). Dissolved Fe was measured by isotope dilution after pre-concentration onto nitrilotriacetate resin on the Micromass IsoProbe ICP-MS at MIT (Lee et al., 2011). Further details on Fe analyses are discussed by Fitzsimmons et al. (in review).

Data presented in this study can be accessed in the Supplemental Material online.

4. Results and Discussion

4.1 $^{230}$Th-$^{232}$Th depth profiles to 1.5 km

We focus first on the 2012-2013 thorium isotope depth profiles in the upper 1.5 km of the water column for a sense of the type of data used to calculate thorium fluxes (Fig. 2). High resolution depth profiles were analyzed in late July 2012, early June 2013 and late September 2013. The mixed layer depths during these sampling casts (based on 0.125 kg/m$^3$ density change)
were 54, 33 and 53 m, respectively, and below 100 m these profiles displayed little
distinguishing hydrography (Figs. 2C, 2D, 2E).

For dissolved $^{232}$Th (Fig. 2A), there were increased concentrations near the surface,
minimum concentrations at the depth of maximum chlorophyll concentration (the DCM, ~120-
140 m), and a relatively constant local concentration maximum at 500-600 m depth. At
intermediate depths (900-1200 m), each profile exhibited smooth variations in concentration but
concentrations at the different sampling dates varied by up to 30%.

The surface $^{232}$Th maxima are consistent with aerosol dust as the major source of $^{232}$-Th to
Station ALOHA, as recognized by Roy-Barman et al. (1996). An interesting feature of these
high-depth resolution measurements is that the surface (5 m) $^{232}$-Th concentration was lower than
that in the core of the mixed layer (25 m depth) at these three sampling times. This is perhaps
related to small-scale scavenging and export dynamics, or particle cycling in general.

The coincidence of the subsurface chlorophyll maximum and the minimum in $^{232}$-Th is
apparently a universal feature for lithogenic trace elements such as Al, Ti and Fe (Dammshäuser
et al., 2013; Fitzsimmons and Boyle, 2014b; Ohnemus and Lam, 2015). This was also true for
dissolved and particulate Fe at Station ALOHA during this study (Fitzsimmons et al., in review).
Increased particle aggregation efficiency, such as through the formation of fecal pellets, may
more efficiently scavenge dissolved $^{232}$-Th from this depth.

Scavenged $^{232}$Th may be partially released through remineralization of particles from the
near-surface upon sinking to mesopelagic depths (300-500 m). Thus remineralization may be
responsible for some of the subsurface $^{232}$-Th maxima at 400-600 m depth. In support of this
view, this depth range coincides with a rapid increase in phosphate concentration and apparent
oxygen utilization, as inferred from HOT climatology
On the other hand, the attenuation of particulate organic carbon at Station ALOHA is most intense at shallower depths, between 100 and 200 m (Bishop and Wood, 2008).

The dominant basalts of the Hawaiian Islands (tholeiitic) are low in Th content, 0.8 ± 0.4 ppm, according to available data in PetDB (www.earthchem.org/petdb) (Lehnert et al., 2000). Nonetheless, with our seawater observations, we cannot fully rule out lateral input of Th from the Hawaiian Islands. For instance, dissolved Mn concentrations reach a maximum near 800 m depth at Station ALOHA (Boyle et al., 2005) that may reflect a coastal source of metals.

The variability in $^{232}$Th concentration at intermediate depths (900-1400 m) could be due to the effect of hydrothermal activity at the nearby Loihi seamount. The iron oxide particles associated with hydrothermal plumes strongly scavenge Th, and depleted deep-sea Th concentrations have been observed up to 1400 km away from a vent site in the Atlantic (Hayes et al., 2015a). While intermediate water $^{233}$Th variability suggests the influence of hydrothermal scavenging here, the effect is apparently too weak to perturb the near-linear $^{230}$Th profiles (Fig. 2). Time-variability in the influence of the Loihi hydrothermal system on trace metals at ALOHA is discussed more fully by Fitzsimmons et al. (in review).

The $^{230}$Th profiles also displayed interesting temporal variations. The theory of reversible scavenging contends that a steady-state is achieved between thorium adsorption and desorption on uniform particles that settle at a constant rate (Bacon and Anderson, 1982). Under these assumptions, one expects $^{230}$Th concentrations to increase linearly with depth with a boundary condition of zero concentration at the surface. While the observed depth profiles are essentially linear (Fig. 2B), it appears that mixing at the surface homogenizes $^{230}$Th concentrations to some depth. Interestingly, the layer of relatively homogeneous $^{230}$Th extends deeper than the density-
defined mixed layer (30-50 m), down to the deep chlorophyll maximum (Fig. 2; see also report by Barone et al. (2015)). This phenomenon is worthy of future time-series study. Potentially a remnant of deep winter mixed layers (> 100 m), the homogeneous surface $^{230}$Th layer could also represent some combination of vertical mixing and enhanced scavenging related to export of organic matter from the euphotic zone.

Another significant observation is that while the surface $^{230}$Th concentrations from June and September 2013 were nearly identical (1.2 μBq/kg), the surface $^{230}$Th concentrations from July 2012 were about a factor of 2 lower (0.6 μBq/kg). This implies a relatively rapid change in scavenging and/or export production. Future time-series studies are warranted to further assess the short-term (daily-monthly) variability in euphotic zone $^{230}$Th concentrations and how closely these changes can be correlated with organic matter export. In the next section, we assess what changes in the removal timescale are implied by these results.

### 4.2 Surface thorium residence times

Residence times of dissolved $^{230}$Th as described in section 2.1 using the 2012-2014 results are presented in Figure 3. In this assessment, we integrate production due to $^{234}$U decay (based on salinity) and the measured $^{230}$Th inventory to 150 m depth. This allows comparison to Th residence times calculated on the basis of $^{234}$Th: $^{238}$U disequilibrium established by previous work at Station ALOHA, during April 1999-March 2000 (Benitez-Nelson et al., 2001) and June-July 2008 (Buesseler et al., 2009). The $^{234}$Th results differ slightly from the approach used here for dissolved $^{230}$Th since the $^{234}$Th fluxes are calculated using unfiltered seawater. Since adsorbed $^{230}$Th concentrations are on the order of ~10-20% of total $^{230}$Th (Roy-Barman et al., 1996), residence times based on total $^{234}$Th can be expected to be up to 10-20% greater than those based on the dissolved phase only. The low percentage of adsorbed $^{230}$Th appears to hold
generally for the remote ocean (Hayes et al., 2015b), but it could be higher in coastal or marginal seas with high terrigenous or riverine input (e.g., (Andersson et al., 1995)).

We also assessed the influence of vertical mixing on surface $^{230}$Th inventories, which could influence the derived residence time. Assuming a vertical mixing coefficient ($K_v$) of $10^{-5}$ m$^2$/s (Charette et al., 2013), using a linear regression of the $^{230}$Th depth profiles (from 150 to 300 m), we can calculate the $^{230}$Th added to the upper 150 m by vertical mixing as $K_v \times d\text{Th}/dz$. The results indicate that this vertical mixing term is 3-5% of the integrated production due to $^{234}$U decay in the upper 150 m. Therefore we can assume that vertical mixing does not significantly affect the $^{230}$Th residence time estimates at Station ALOHA.

Nearly all of the thorium residence time estimates fall in the range of 1 to 3 years with no evident seasonal cycle (Fig. 3). In the HOT climatology, organic carbon export at 150 m is highest in May-August. While export seasonality is relatively weak in this oligotrophic, subtropical location (Church et al., 2013), long-term monitoring has revealed episodic export events related to diatom blooms and symbiotic cyanobacteria, typically in late July and early August (Karl et al., 2012).

The concept of “residence time” used here is based on a steady-state assumption for sources and sinks. Therefore with a residence time of ~2 years, one would not expect significant variation in the removal timescale over a period of months. However, the range in observed Th residence times for Station ALOHA indicates that this steady-state assumption is not quite correct. More precisely, the steady-state for scavenging removal of Th appears to hold within a factor of 2-3. The range in removal timescales observed based on $^{230}$Th is similar to that based on $^{234}$Th. Thus, it seems the rate of thorium scavenging can change dynamically at Station ALOHA.
possibly related to export pulses, but the data are consistent with a long-term average thorium residence time of 2 ± 1 years in the upper 150 m.

4.3 Surface $^{232}$Th concentrations

With relatively good control on the removal timescale of thorium, we turn to observed variability in surface $^{232}$Th concentrations. Barring significant fluxes due to lateral circulation, this variability represents the balance between removal by scavenging and input by dust. Smaller volume requirements for analysis (<1 liter) allowed us to investigate $^{232}$Th from daily, monthly and decadal timescales.

Collected during a series of cruises in summer 2012 (HOE-DYLAN), daily-scale samples of 250 mL were analyzed for dissolved and particulate $^{232}$Th. Sample size required combining the samples from 2-4 days for dissolved $^{232}$Th, contributing to some temporal smoothing. Dissolved concentrations ranged from 45-90 fmol/kg (Fig. 4C). Particulate $^{232}$Th, although measured at a higher, daily resolution, had a higher range of variability, from 10-290 fmol/kg. Of the total seawater $^{232}$Th (dissolved + particulate) during HOE-DYLAN, on average 42% was in the particulate phase (range 26-66%). This fraction particulate is higher than that for $^{230}$Th (~15%, Roy-Barman et al., 1996) since particulate $^{232}$Th represents both adsorbed Th and structural Th in mineral dust.

The decadal time-series observations (1994-2014) of total $^{232}$Th (Fig. 4A) exhibit a range in concentration (~50-300 fmol/kg) that is consistent with the higher frequency observations of particulate Th in 2012-2013. Since most of the data fall within the range of 50-150 fmol/kg, we are not fully confident in the five observations of elevated concentrations (150-300 fmol/kg) observed in 1994, 1998, and 1999 samples. In particular, the 1998 and 1999 samples were collected using the moored MITESS units, as discussed in section 3.2. Mooring-collected water
at times had higher Th concentrations than contemporaneous ship-based sampling (Fig. 4) and
thus the possibility of contamination during sampling, sample storage, or sample analysis cannot
be fully discounted. In fact, the variable 1994 results from Roy-Barman et al. (1996) came from
multiple samples collected on the same Niskin bottle cast. Spatial variability, related to
mesoscale eddies, is another potential source of rapid changes in surface $^{232}$Th concentration.
Conservatively excluding the elevated observations >150 fmol/kg, no significant temporal term
trend can be derived.

When all observations are placed on a monthly axis (Fig. 4B), there is little indication of
elevated surface $^{232}$Th concentrations during the spring (Mar-Jun) season of Asian dust transport
over the North Pacific. It appears that dissolved $^{232}$Th may be relatively constant throughout the
year, consistent with the Th residence times of ~2 yrs derived in section 4.2. Unfortunately, few
observations of dissolved $^{232}$Th have been yet made during the spring season when dust input can
increase by 2 orders of magnitude (Hyslop et al., 2013). Of course, these data are sparse, but they
do provide a baseline of variability against which future trace metal observations can be
measured.

4.4 Fe/Th ratio behavior in surface water and in colloidal content

Before applying the $^{232}$Th flux technique, comparison of the time-series behavior of Fe
(Fitzsimmons et al., in review) and $^{232}$Th is informative in terms of relative solubility and relative
removal rates (Fig. 5). This is possible because both elements have been analyzed on the same
samples from HOE-DYLAN, HOE-PhoR and many of the HOT archive samples.

In the context of daily, monthly and decadal variability, it appears that the ratio of total
and particulate Fe/$^{232}$Th tends to be at or above the dust-ratio of 11,040 mol/mol, while dissolved
Fe/$^{232}$Th is at or below the dust-ratio (Fig. 5A & 5B). These observations are consistent with
input at the dust Fe/$^{232}$Th ratio and a strong sink from biological uptake for Fe. Thus, the
dissolved phase is left depleted in Fe relative to $^{232}$Th, while the particulate phase becomes
enriched in biogenic Fe. The total Fe/Th ratio often exceeds the dust ratio as well, possibly
because biogenic particulate Fe may be efficiently recycled and thus may reside in the surface
longer than particulate Th. Interestingly, the samples from moored MITESS units (open symbols
in Fig. 5) had Fe/Th ratios quite close to the near-crustal ratio of Asian dust. This fact does not
necessarily exclude the possibility for Th contamination in these samples but does suggest any
potential metal contamination was of near-crustal composition.

The partitioning between dissolved and total/particulate Fe/$^{232}$Th centers on the dust ratio
(Fig. 5C). We interpret this to mean that the relative fractional solubility of Fe and $^{232}$Th ($S_{Fe/Th}$)
is close to 1. An alternate interpretation would be that $^{232}$Th is more efficiently leached from
dust, leaving the particulate phase enriched in Fe/$^{232}$Th and the dissolved phase depleted in
Fe/$^{232}$Th. However, given the known ability for phytoplankton to efficiently utilize Fe from dust
sources (e.g., (Rubin et al., 2011)), the assumption of $S_{Fe/Th} = 1$ during dissolution followed by
rapid biological uptake of Fe seems more likely. It is difficult to assign a quantitative
uncertainty to the relative solubility with the existing data. The measured seawater Fe/$^{232}$Th
ratios ($n = 30$) are on average within $54 \pm 51\%$ (1 sigma) of the Asian dust ratio. Therefore, the
relative solubility is likely close to 1 with less than 50% uncertainty, but 50% could be used as a
conservative uncertainty estimate ($S_{Fe/Th} = 1 \pm 0.5$). Consideration of the size-partitioning of Fe
and Th within the dissolved phase provides another constraint on the pathways these elements
take after being released by dust. This investigation was also used as an opportunity to
determine whether $^{232}$Th and $^{230}$Th have coherent speciation, as assumed for the $^{232}$Th flux
method. Figure 6 presents these results based on measurements of ultra-filtered seawater from
HOE-PhoR-II in September 2013. We define colloidal Th as dissolved (< 0.45 µm) minus soluble (< 10 kDa).

Of the measured dissolved Th, 8-25% was found in the colloidal phase (0.45 µm-10 kDa ≈ 0.01 µm). The total Th recovery during ultrafiltration was nearly complete (88-100%), implying that this 8-25% of dissolved Th was indeed colloidal in size, not an artifact of Th sorption/loss to the ultrafiltration system. Furthermore, at least at 15 m, 130 m (DCM), and 1000 m, the colloidal percentage for $^{232}$Th and $^{230}$Th agreed within the uncertainty of the measurements. This result implies coherent speciation of these thorium isotopes despite very different sources, and it supports the use of $^{230}$Th as a tracer for $^{232}$Th removal. This coherent speciation result agrees with previous measurements of the $^{232}$Th/$^{230}$Th ratio of filtered (< 0.2 µm) and ultrafiltered (< 1 kDa) solutions from the Mediterranean Sea (Roy-Barman et al., 2002).

The role of colloids in Th scavenging has much history and deserves a few words of context. Early models of scavenging inferred that Th likely goes through a colloidal intermediate before being scavenged by larger, sinking particles (Honeyman et al., 1988; Honeyman and Santschi, 1989). Subsequent attempts at measuring colloidal Th focused largely on $^{234}$Th (see review by (Guo and Santschi, 2007)), in part due to its use in quantifying organic matter export. A generalization might be made that outside of the coastal ocean, colloidal $^{234}$Th was found to be a relatively small (<15%) proportion of the total dissolved (e.g., (Guo et al., 1997; Huh and Prahl, 1995; Moran and Buesseler, 1992)), which is also consistent with our $^{230}$Th and $^{232}$Th results. Recent observations from the North Atlantic (Hayes et al., 2015b), however, observed scavenging characteristics consistent with a strong role for Th colloids as predicted by the “colloidal pumping hypothesis” of Honeyman and Santschi (1989), even at open-ocean particle
concentrations of < 10 µg/kg seawater. Further observations on the geographic distribution of
colloidal Th are clearly warranted.

Our paired observations of Th and Fe size partitioning nonetheless provide additional
information on their physicochemical speciation in a comparative sense. Dissolved Fe has a
much higher colloidal content at ALOHA than Th (Fig. 6). Above the DCM, dissolved Fe can be
>50% colloidal. In the deeper water column, to 1.5 km depth, colloidal Fe is relatively constant
at 40% (with the exception of one sample < 10% colloidal at 650 m). Since Fe and \(^{232}\)Th are
apparently solubilized from dust with equal fractional solubility, this difference in size-speciation
is most likely also due to the selective uptake or complexation of Fe by organic substrates.
Ligands, in the form of macromolecular organic molecules or organic colloidal particles, most
likely complex Fe released from dust quite rapidly in the upper water column (Bressac and
Guieu, 2013; Mendez et al., 2010). We hypothesize that organic Fe-binding ligands are
predominantly responsible for converting such a large percentage of dissolved Fe to colloidal
size. The inorganic speciation of Th in seawater is largely hydroxo- complexes (Santschi et al.,
2006). Other similarly hydrolyzable metals such as Al and Ti do not have significant colloidal
components (Dammshäuser and Croot, 2012).

Similar to Al and Ti, the abundance of colloidal ligands (>10 kDa) with an affinity to
complex Th must also be small compared to the source of dissolved Th from dust. This finding
does not necessarily contradict previous evidence for significant organic complexation of Th in
seawater (Santschi et al., 2006). It does require, however, that any significant Th complexation is
done by small (<10 nm), low-molecular weight organic molecules, at least in the subtropical
North Pacific.
Greater uptake of Fe into the colloidal phase is another piece of evidence that suggests that dissolved Fe is cycled more rapidly than Th in the upper water column. The innovation of the $^{232}$Th flux method is our ability to be quantitative about the rates of Fe removal, which are presented in the next section.

4.5 Iron residence times

Using the 2012-2013 $^{230}$Th profile data, we extend our calculations for Th residence time down to 1.5 km water depth in Fig. 7A. The average residence time for the depth zone between the surface and the DCM is 1-2 years. The Th residence times increase nearly linearly with integration depth to 14 years for the average residence time between the surface and 1.5 km. We do not plot integrated values shallower than the DCM (~120 m) on the assumption that steady-state Th scavenging may not apply under the conditions of stronger mixing and organic matter export within the euphotic zone. Dividing the integrated dissolved $^{232}$Th inventories by these residence times gives our estimate of dissolved $^{232}$Th flux, as a function of integration depth, in Fig. 7B.

In June and Sept. 2013, the dissolved $^{232}$Th flux increased with integration depth and began to level-off around 500 m. This pattern reflects that, at these times, the inventory of dissolved $^{232}$Th increased with integration depth slightly more quickly than the increase in Th residence time with depth. Interestingly, in July 2012, the dissolved $^{232}$Th flux decreased with integration depth, reflecting that the Th residence time increased more quickly than the dissolved $^{232}$Th inventory, essentially because the mixed layer $^{230}$Th concentrations were exceptionally low at this time. Estimated $^{232}$Th fluxes are clearly quite sensitive to short-term variability in scavenging rates. We suggest further time-series analysis along with modelling efforts that contain circulation and realistic particle fluxes to determine more quantitatively the sensitivities
involved in calculating dissolved $^{232}\text{Th}$ fluxes during moderate changes in scavenging rates and
dust input.

The three flux profiles converge around 1000 m depth. This is encouraging that over
longer integration times, 10-15 years in this case, we estimate consistent lithogenic metal fluxes
at multiple time points. Using Eq. 1, the dissolved $^{232}\text{Th}$ fluxes are simply converted to dust-
dissolved Fe fluxes, using $S_{\text{Fe/Th}} = 1$ and $(\text{Fe/Th})_{\text{dust}} = 11,040 \text{ mol/mol}$, shown in the second x-
axis in Fig. 7B. The depth profiles of dissolved Fe concentrations from the same sampling
campaigns are shown in Fig. 7C (Fitzsimmons et al., in review). Finally, using Eq. 2, by
integrating Fe inventories and dividing by the dust-dissolved Fe fluxes, we estimate the residence
time of dissolved Fe, as a function of integrated depth in Fig. 7D.

In the upper 250 m, the residence time of dissolved Fe is 6 months to 1 year, again
assuming that the sole source of Fe to the surface ocean at Station ALOHA is aerosol dust
deposition. This range agrees well with the 6 month residence time estimated previously at
Station ALOHA (Boyle et al., 2005), and with other estimates of surface ocean dissolved Fe
residence times from the Atlantic based on measured Fe concentrations and assumptions about
soluble aerosol deposition (Bergquist and Boyle, 2006; Jickells, 1999; Ussher et al., 2013). With
such fast turnover times, dissolved Fe concentrations in surface waters can be expected to vary
on monthly to yearly timescales with changes in the seasonal input of dust from Asia. This is in
fact exactly what was observed over the HOT and HOE time-series (Fitzsimmons et al., in
review).

Available aerosol data suggest that Asian dust transport over the North Pacific had no
significant long-term trend from 1981 to 2000 (Prospero et al., 2003) and perhaps a 6% decline
over the past 10 years (Hyslop et al., 2013). Because of a nearly immediate impact on surface
water Fe concentrations and the associated ecological consequences, it is important to monitor future changes in Fe sources. Sources such as Asian desert dust in our changing climate may vary independently of other Fe sources such as combustion aerosols.

As one integrates further from 250 m to 1500 m, while the dissolved Fe fluxes change only moderately, the dissolved Fe residence times increase quickly to about 10 years at 1500 m depth. This is due to the large increase in Fe concentrations at these depths due to remineralization of Fe from sinking organic material and some portion of Fe accumulated and transported to ALOHA laterally via deep ocean circulation. There is potentially additional input of Fe at ~1 km depth due to Loihi hydrothermal activity. Additional lateral sources would cause our dust-based dissolved Fe residence time to be an overestimate, implying even faster timescales of Fe removal. On the other hand, as discussed in the next section, the 10 year Fe residence time at 1500 m could as well be an underestimate, if the geochemical cycles of Th and Fe become decoupled at greater depths where dust dissolution is no longer the most significant source of dissolved Fe.

4.6 Fe and Th decoupling in the deep ocean

Our focus on the upper water column stems from our motivation to understand trace metal cycling due to aerosol deposition and export production. We can extend our analysis of Fe and Th into the deep ocean (4.5 km water depth at Station ALOHA) to learn about the geochemistry of these elements over decadal-to-centennial timescales. In Figure 8, we compiled available deep profiles from Station ALOHA for dissolved Fe (Boyle et al., 2005; Fitzsimmons et al., in review; Morton, 2010) and dissolved $^{232}$Th and $^{230}$Th (this study; Roy-Barman et al. (1996)).
Variability in dissolved Fe at 1-1.5 km is clearly apparent, likely due to hydrothermal inputs. Below 1.5 km depth, Fe, $^{232}$Th, and $^{230}$Th display relatively constant profile shapes, at least during the sparse sampling dates. From 2 km depth to the bottom, dissolved Fe is nearly constant or slightly decreases with depth to about 0.4 nmol/kg, while dissolved $^{232}$Th actually increases with depth from 50 to 180 fmol/kg below 3000 m. This divergence in profile shape already suggests a decoupling of the behavior of these elements in the deep ocean.

The deep ocean appears to contain an additional source for $^{232}$Th. This source is potentially related to resuspension of diagenetically-altered sediments at the seafloor (Hayes et al., 2013a; Okubo et al., 2012). The bottom-increase in $^{232}$Th begins nearly 2 km above the seafloor, much higher than typical benthic vertical mixed layers (50-100 m) (Richards, 1990). This phenomenon, as observed with km-scale nepheloid layers (McCave, 1986), suggests that the $^{232}$Th at abyssal depths of Station ALOHA is being mixed in laterally from locations where isopycnals impinge on surrounding bathymetry.

Also related to bottom sediment resuspension, the July 2012 profile of $^{230}$Th displays a negative concentration anomaly, or deficit of $^{230}$Th, with respect to the linear profile near the seafloor (Fig. 8C). This bottom $^{230}$Th deficit is indicative of enhanced bottom scavenging as observed in many parts of the deep North Pacific (Hayes et al., 2013b; Okubo et al., 2012). It is non-intuitive that a bottom layer where the scavenging removal of Th is enhanced compared to the overlaying water column would also be a strong source of $^{232}$Th. The resuspension of bottom sediments may produce such a strong release of $^{232}$Th that this source more than compensates for enhanced scavenging. Another contributing factor may be that the resuspended thorium could have a much higher $^{232}$Th/$^{230}$Th ratio than the water column due to age-decay of $^{230}$Th in the sediments.
Dissolved Fe, on the other hand, appears unaffected by bottom processes, displaying only a slight decrease in concentration with depth (Fig. 8A). The slight decrease with depth may be related to scavenging of Fe as deep water masses age (Bruland et al., 1994). If we extend our integrated residence time approach to the deep Fe profile at Station ALOHA (Fig. 9), we derive a whole ocean residence for dissolved Fe of only 30 years. This is significantly shorter than the 100-300 year estimates of the ocean residence time for dissolved Fe based on deepwater scavenging (Bergquist and Boyle, 2006; Bruland et al., 1994). This discrepancy must arise because the deep ocean source of $^{232}$Th does not add dissolved Fe to the water column at a crustal ratio, unlike what occurs during near-surface dust dissolution. Thus, the $^{232}$Th flux method for Fe residence times probably should not be extended to the deep ocean.

The question remains: how is an element like Th, a trace component of continental material, added to the deep ocean without a simultaneous release of a major crustal element like Fe? The answer is likely related to solubility.

Dissolved Fe in the deep central North Pacific at ~0.5 nmol/kg has been found to be at near solubility equilibrium with Fe(III) hydroxide (Kitayama et al., 2009; Kuma et al., 2003). These studies determine Fe(III) solubility by adding gamma-emitter $^{59}$Fe(III) to filtered seawater, allowing the solutions to come to solubility equilibrium with Fe(III) hydroxide over several weeks, subsequently filtering the seawater and then counting the $^{59}$Fe gamma-activity on the final filtrate. The observed ~0.5 nmol/kg solubility is elevated over Fe solubility in inorganic seawater because of the presence of organic ligands (Liu and Millero, 2002). Thus, since the deep Pacific is in a near saturation state, dissolved Fe can no longer be expected to increase, even in the presence of increasing Th concentrations.
A problem with this argument is that electrochemically-determined Fe ligand concentrations at Station ALOHA are up to 2 nmol/kg, well in excess of dissolved Fe concentrations (Rue and Bruland, 1995), as found in most of the world ocean (Gledhill and Buck, 2012). However, it may not be kinetically appropriate to compare Fe ligand concentrations directly with seawater solubility. In either estimation, deepwater dissolved Fe is at least close to (within the same order of magnitude) our best estimates of Fe solubility.

While much less is known about Th solubility in seawater, our large underestimate of Fe residence time in the deep ocean implies that the deep North Pacific, with Th at ~180 fmol/kg, is below Th solubility equilibrium. Near seawater pH and ionic strength, the solubility of Th(IV) hydroxide may be as high as 0.5-1 nmol/kg, compared to 1 fmol/kg for crystalline ThO\(_2\), due to the amorphous nature of Th(OH)\(_4\) solids (Neck et al., 2003). Also, electrochemical methods suggest organic Th ligands exist at nanomolar concentrations (Hirose, 2004). Despite our finding of low colloidal Th content, organically-bound Th could of course be present at Station ALOHA if the complexes are smaller than ~10 nm. We advocate direct measurements of Th solubility in seawater, perhaps using radio-tracer additions with similar protocols as developed for Fe (Kuma et al., 1996; Schlosser and Croot, 2008), to confirm that Th exists in the deep ocean at much less than its equilibrium solubility. This would explain the fact that dissolved Th concentrations continue to grow from lithogenic sources in the deep North Pacific, where Fe concentrations become fixed by a solubility limit.

5. Conclusions

Using time-series data from the North Pacific, this study finds variability in surface Fe and \(^{232}\)Th concentrations consistent with a source from Asian dust. The dust source likely has a relative Fe/Th fractional solubility close to 1. The application of \(^{230}\)Th scavenging rates to \(^{232}\)Th
inventories allows the accurate evaluation of the flux of dissolved metals from dust in the remote surface ocean. The source flux of dissolved Fe, derived from $^{230}$Th-based timescales, suggests that dissolved Fe in the upper 250 m is turning over in 1 year or less. A compelling implication of this result is that Fe delivery to phytoplankton can be expected to vary with seasonal-to-interannual changes in dust delivery from Asia. Continued monitoring of Fe-dependent biological processes, such as nitrogen fixation, are crucial to anticipate the consequences of changing land-use and/or industrial processes that may significantly affect eolian sources of Fe to the North Pacific.

In addition, by comparing Fe and Th size-partitioning, we find evidence that colloidal Fe may be of predominantly organic composition in the subtropical North Pacific. We also hypothesize that iron reaches a solubility limit in the deep sea (>2 km) while Th does not. While less controversial for Th, this result questions the relevance of “excess” Fe ligands in the deep sea. Overall, however, the kinetic box model approach to tracing dust-derived elements (Fig. 1) appears well-suited in the upper water column (~250 m).

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Figure Captions
**Figure 1.** Tracing the Fe cycle with the behavior of the long-lived thorium isotopes. Thorium-230 has a well-known source from the radioactive decay of its parent $^{234}$U. This allows a quantitative estimate of Th removal due to scavenging on to particles. This removal rate can be used to estimate the steady-state source of $^{235}$Th from the partial dissolution of aerosol dust, assuming dust dissolution and scavenging dominate the Th cycle which may be most relevant in the remote surface ocean. While Fe has many more terms in its biogeochemical cycling, its ultimate source from dust dissolution can be predicted using known $^{232}$Th fluxes and the relative solubility of Fe and Th. Assuming Fe is derived only from dust, one can then estimate a maximum Fe residence time or minimum turnover rate.

**Figure 2.** Depth profiles from the Hawaii Ocean Time-series Station ALOHA from sampling campaigns in 2012-2013. In July 2012 and June 2013, profiles for dissolved $^{232}$Th (A) and $^{230}$Th (B) were collected in two casts (shallow to 250 m and deep to 1500 m) on different days. Relative uncertainty in isotope concentrations was 1-5% and thus errors bars would be close to the symbol size. The hydrographic profiles (C-F) are shown from the shallow cast only.

**Figure 3.** Thorium residence times, or turnover rates, calculated for the upper 150 m at Station ALOHA on a monthly axis combining data from 1999 to 2014. These times are calculated by comparing integrated Th inventories to integrated production by uranium decay. The $^{234}$Th-based results are reported by Buesseler et al. (2009) and Benitez-Nelson et al. (2001). Note the $^{230}$Th-based results from March 2014 are not based on profiles but on single samples from 25 m, assuming uniform concentrations in the upper 150 as seen in the 2012-2013 profiles (Fig. 2).

**Figure 4.** Station ALOHA time-series data from the surface ocean (0-10 meters depth) on dissolved (filtered at 0.45 or 0.4 µm), total (unfiltered) and particulate (digested 0.4 µm filter) $^{232}$Th in full time-series (1994-2014) (A), monthly climatology (1991-2014) (B) and during a daily resolution period in July-Sept. 2012 (C). Note change in scale of y-axes at 160 fmol/kg. Results from 1994 were reported by Roy-Barman et al. (1996). Open circles represent samples collected using a mooring rather than ship-based sampling (Sec. 4.1). Relative uncertainty in dissolved, total and particulate $^{232}$Th concentrations was 1-10%.

**Figure 5.** Station ALOHA time-series data from the surface ocean (0-10 meters depth) (A), monthly climatology (B) and a daily resolution period in July-Sept. 2012 (C) of the dissolved (filtered at 0.45 or 0.4 µm), total (unfiltered) and particulate (digested 0.4 µm filter) Fe/$^{232}$Th ratio. Note change in scale of y-axes at 25,000 mol/mol. The dotted lines represent the Fe/$^{232}$Th ratio of Asian dust of 10,800 ± 1,200 mol/mol (1σ). Note in (C), four samples with particulate Fe/$^{232}$Th ratios greater than 40,000 are not shown. Open circles represent samples collected using a mooring rather than ship-based sampling (Sec. 4.1).

**Figure 6.** Depth profiles of the percentage of dissolved metals (<0.45 µm for Th or <0.4 µm for Fe) that are in the colloidal size fraction (roughly 10-400 nm) from Station ALOHA in late September 2013. Colloidal content is estimated by subtracting the metal concentration in 0.4 µm filtered seawater (dissolved) from that passing through a 10 kDa membrane filter by cross-flow filtration (soluble). Colloidal fractions of $^{232}$Th and $^{230}$Th agree within uncertainties, while Fe colloidal content is 2-3 times larger.
**Figure 7.** Application of dissolved $^{232}$Th fluxes to predict the residence time of dissolved Fe in seawater at Station ALOHA during 2012-2013. Dissolved Th residence times (A) are calculated as a function of integration depth using radioactive disequilibrium between $^{234}$U and $^{230}$Th. Integrated values shallower than the deep chlorophyll maximum (~120 m) are not included on the assumption that steady-state Th scavenging may not apply within the euphotic zone. The integrated $^{232}$Th inventories divided by these residence times produces an estimate of the dissolved $^{232}$Th flux (B) due to dust dissolution. Assuming equal fractional solubilities dissolution and a near crustal composition for Asian dust, the flux of dissolved Fe from dust can be predicted using the second x-axis in (B). The integration of dissolved Fe inventories based on concentration profiles shown in (C) (Fitzsimmons et al., in review), produces our estimate of dissolved Fe residence time in (D, note change in scale of x-axis at 1.2 yrs).

**Figure 8.** Full ocean depth profiles from Station ALOHA for dissolved Fe (A), $^{232}$Th (B) and $^{230}$Th (C) using data from this study (July 2012) and compiled from the literature. Iron data from April 2001 and July 2002 were reported by Boyle et al. (2005) and from June 2002 by Morton (2010). Dissolved Th data from 1994 were reported by Roy-Barman et al. (1996). Note in (C) the dotted grey line is the linear regression of $^{230}$Th data between 1 and 3.5 km, which when extended to the seafloor demonstrates that the bottom two samples are less than expected from reversible scavenging and imply enhanced scavenging (assuming no other processes affect supply and removal of $^{230}$Th here).

**Figure 9.** Application of dissolved $^{232}$Th fluxes to predict Fe residence times for the full depth ocean at Station ALOHA. Here data from July 2012 are used to calculate $^{232}$Th fluxes (A). The depth profiles of Fe concentrations presented in Fig. 8 were averaged to calculate the dissolved Fe residence times as a function of integration depth (B). The 30 year ocean residence for dissolved Fe is significantly lower than the century-scale residence times derived by other approaches, suggesting that $^{232}$Th flux may not be an accurate proxy for Fe sources in the deep ocean.

References


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Figure 1

234U \rightarrow \text{Radiogenic Source} \quad \text{230Th} \quad \text{232Th} \quad \text{Lithogenic Source} \quad \text{Dust} \quad \text{Sinking Particles} \quad \text{Scavenging Removal} \quad \text{Fe} \quad \text{(much more complicated)} \quad \text{Biological Uptake} \quad \text{Respiration/Regeneration} \quad \text{Redox}
Th residence times: 150m, Station ALOHA

- This study (Th-230)
- Buesseler et al. 2009 (Th-234)
- Benitez-Nelson et al. 2001 (Th-234)
Figure 4
Figure 6  % Colloidal metals (of dissolved)
Figure 7

A: Diss. Th residence time [yrs]

B: Diss. $^{232}$Th Flux [nmol/m²/yr]

C: Dissolved Fe [nmol/kg]

D: Diss. Fe Residence Time [yrs]
Figure 8

A) Dissolved Fe [nmol/kg]

B) Dissolved $^{232}$Th [fmol/kg]

C) Dissolved $^{230}$Th$_{xs}$ [μBq/kg]
Figure 9

A

Diss. $^{232}$Th Flux [nmol/m$^2$/yr]

Integration Depth [km]

Dust-diss. Fe Flux [µmol/m$^2$/yr]

B

Diss. Fe Residence Time [yrs]

Integration Depth [km]

?
Supplemental Data

Click here to download Electronic Annex: Hayes ALOHA Th Fe Supplemental Data for GCA.xlsx