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Key Points:
• $^{230}$Th normalization is a robust tool for calculating sedimentary mass fluxes
• $^{230}$Th may be affected by hydrothermal and boundary scavenging in certain discrete regions
• Generally, $^{230}$Th mass fluxes are preferable over age model-based mass accumulation rates

Supporting Information:
• Supporting Information S1
• Table S1

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Abstract: $^{230}$Th normalization is a valuable paleoceanographic tool for reconstructing high-resolution sediment fluxes during the late Pleistocene (last ~500,000 years). As its application has expanded to ever more diverse marine environments, the nuances of $^{230}$Th systematics, with regard to particle type, particle size, lateral advective/diffusive redistribution, and other processes, have emerged. We synthesized over 1000 sedimentary records of $^{230}$Th from across the global ocean at two time slices, the late Holocene (0–5,000 years ago), or 0–5 ka) and the Last Glacial Maximum (18.5–23.5 ka), and investigated the spatial structure of $^{230}$Th normalized mass fluxes. On a global scale, sedimentary mass fluxes were significantly higher during the Last Glacial Maximum (1.79–2.17 g/cm$^2$/kyr, 95% confidence) relative to the Holocene (1.48–1.68 g/cm$^2$/kyr, 95% confidence). We then examined the potential confounding influences of boundary scavenging, nepheloid layers, hydrothermal scavenging, size-dependent sediment fractionation, and carbonate dissolution on the efficacy of $^{230}$Th as a constant flux proxy. Anomalous $^{230}$Th behavior is sometimes observed proximal to hydrothermal ridges and in continental margins where high particle fluxes and steep continental slopes can lead to the combined effects of boundary scavenging and nepheloid.


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interference. Notwithstanding these limitations, we found that $^{230}$Th normalization is a robust tool for determining sediment mass accumulation rates in the majority of pelagic marine settings (>1,000 m water depth).

1. Introduction

Burial fluxes of different components of marine sediment provide insight into a wide variety of surface processes that are central to the Earth system, including marine export productivity, windblown dust deposition on the sea surface, carbon storage as organic matter and calcium carbonate, and hydrothermal activity on the seafloor. The traditional approach to calculating marine burial fluxes relies on determining the average mass accumulation rates based on age model tie points, intervening sediment thickness, and average sediment dry bulk density (e.g., Broecker, 1971). The temporal resolution of this approach is limited by the robustness of the age model, including the number of chronological tie points and their associated errors (e.g., Francois et al., 2004). Furthermore, this approach can easily be biased by sediment redistribution on the seafloor (e.g., Johnson & Johnson, 1970), where lateral sediment transport can exceed the vertical rain of particles from the water column. As a result, constant flux proxies such as $^{230}$Th have been developed to provide more robust estimates of mass accumulation on the seafloor.

Constant flux proxies are geochemical parameters with well-constrained and stable source functions, such as $^{230}$Th (Bacon, 1984; Francois et al., 2004) and $^3$He (Marcantonio et al., 1996; McGee & Mukhopadhyay, 2013; Schlosser & Winckler, 2002; Winckler et al., 2004). $^{230}$Th is produced by the steady decay of uranium dissolved in seawater, after which it is rapidly removed by sinking particles and buried on the seafloor (see section 2) (Bacon, 1984; Francois et al., 1990; Francois et al., 2004; Suman & Bacon, 1989). Because the $^{230}$Th production rate is relatively uniform in space and time, variability in $^{230}$Th concentrations in the sediment can theoretically be attributed to variable dilution by changes in sediment mass flux. Thus, sedimentary $^{230}$Th concentrations can be used to reconstruct changes in sediment mass fluxes over time. This technique, $^{230}$Th normalization, allows both high-resolution sediment mass flux reconstructions independent of age model tie points and isolation of only the vertical component of sedimentation, regardless of the amount of lateral sediment transport.

$^{230}$Th has been used to assess burial fluxes for more than 35 years (Bacon, 1984), with the first comprehensive review of its use, advantages, and limitations published more than a decade ago (Francois et al., 2004). In the intervening 15 years, analysis of $^{230}$Th has become more commonplace, with advances in methodology (e.g., evolving from alpha counting to inductively coupled plasma mass spectrometry [ICP-MS]) resulting in an order of magnitude increase in the amount of data available. At the same time, the GEOTRACES program and associated modeling studies have improved our understanding of $^{230}$Th cycling in the modern ocean. With these changes in mind, and the increasing utilization of sedimentary $^{230}$Th across the global ocean, we have produced an updated compilation that provides an overview of the methodology and current understanding of the $^{230}$Th normalization technique on a global scale.

2. Background: The Marine Geochemistry of $^{230}$Th

In this section, we review the current understanding of $^{230}$Th systematics in the ocean, provide an updated $^{230}$Th production rate, provide revised lithogenic and authigenic correction equations, and present recommendations for best practices in future studies.

2.1. Production of $^{230}$Th in the Water Column

$^{230}$Th is produced in seawater by radioactive decay of long-lived $^{234}$U. Because the marine residence time of uranium (~400,000 years; Henderson, 2002) is orders of magnitude longer than the ocean mixing time (~1000 years), $^{230}$Th production is ubiquitous in the water column and occurs at a relatively uniform rate. This production rate ($\beta_{230}$, in units of decays per cubic centimeter per thousand years, dpm/cm$^3$/kyr) can be calculated using the activity of uranium in seawater ($A_{234U}$, which is equivalent to the concentration of $^{234}$U multiplied by the decay constant of $^{234}$U), and the decay constant of $^{230}$Th ($\lambda_{230}$), as demonstrated by Francois et al. (2004). As more precise values of the decay constants are determined, $\beta_{230}$ is progressively refined over time. Uranium concentrations are conservative and
scale with salinity (Chen et al., 1986; Owens et al., 2011), but this relationship is defined in terms of the major uranium isotope, $^{238}\text{U}$. We thus rewrite equation (1a) as equation (1b) by replacing the activity of $^{234}\text{U}$ with the activity of $^{238}\text{U}$ multiplied by the $^{234}\text{U}/^{238}\text{U}$ activity ratio in seawater (1.1468; Andersen et al., 2010). We can then replace the concentration of $^{238}\text{U}$ with the salinity ($S$) relationship of Owens et al. (2011) to obtain equation (1c). Finally, we use the latest half-life for $^{230}\text{Th}$ (75,584 ± 110 years; Cheng et al., 2013) to calculate its decay constant, and we assume a salinity of 35 to determine the mean ocean $\beta_{^{230}}$ (equation (1d)).

$$\beta_{^{230}} = \lambda_{^{230}} A_{^{234}\text{U}}$$  \hspace{1cm} (1a)

$$\beta_{^{230}} = \lambda_{^{230}} A_{^{238\text{U}*}} \left( \frac{A_{^{234\text{U}}}}{A_{^{238\text{U}}}} \right)_{SW}$$  \hspace{1cm} (1b)

$$\beta_{^{230}} = \lambda_{^{230}} \left[ 0.0786 S - 0.315 \right] \left( \frac{A_{^{234\text{U}}}}{A_{^{238\text{U}}}} \right)_{SW}$$  \hspace{1cm} (1c)

$$\beta_{^{230}} = \left( \frac{\ln(2)}{75,584} \right) \left[ 0.0786 S - 0.315 \right] \left[ 1.1468 = 2.562 \pm 0.05 \times 10^{-5} \text{ dpm cm}^{-3} \text{kyr}^{-1} \right]$$  \hspace{1cm} (1d)

Salinity variations affect $\beta_{^{230}}$ at a rate of $0.08266 \times 10^{-5} \text{ dpm/cm}^3 \text{kyr}$ for each change in salinity by 1 (unitless, according to the practical salinity scale of 1978). This rate is only slightly greater than the error associated with $\beta_{^{230}}$, and statistically significant changes to $\beta_{^{230}}$ require relatively extreme changes in salinity. For example, $\beta_{^{230}}$ is about 10% lower in water with $S = 32$ compared to $S = 35$, and $\beta_{^{230}}$ is 10% higher in water with $S = 38$ compared to $S = 35$. Salinity variations within the water column are unlikely to greatly affect the net $^{230}\text{Th}$ production on the time scales of sedimentation, and in general, we recommend using a single $\beta_{^{230}}$ for each record to maintain consistency.

Unlike uranium, which is highly soluble, $^{230}\text{Th}$ is strongly particle reactive and is thus rapidly removed from seawater by sorption onto sinking particles (particle scavenging; Bacon & Anderson, 1982). Dissolved and particulate $^{230}\text{Th}$ concentrations generally increase linearly with water depth. This feature is best explained by reversible scavenging, a process by which $^{230}\text{Th}$ adsorbed onto the surface of sinking particles continuously exchanges with the dissolved $^{230}\text{Th}$ pool as particles settle through the water column (Bacon & Anderson, 1982; Nozaki et al., 1987). As $^{230}\text{Th}$ is highly insoluble, its residence time in seawater does not exceed a few decades (20–40 years; Nozaki et al., 1981).

### 2.2. Analytical Methodology

Analysis of thorium ($^{230}\text{Th}$ and $^{232}\text{Th}$) and uranium ($^{238}\text{U}$, $^{235}\text{U}$, and $^{234}\text{U}$) generally proceeds by aliquoting 100–200 mg of sediment, spiking with $^{229}\text{Th}$ and $^{236}\text{U}$, complete acid digestion, column chromatography to isolate and concentrate the nuclides, and measurement by ICP-MS (e.g., as described in Fleisher & Anderson, 2003). The specific details of this procedure may vary between laboratories, based on, for example, available instrumentation, sample throughput, and required precision. For example, the digestion usually includes a “cocktail” of nitric acid, hydrofluoric acid, and perchloric acid (e.g., Jacobel et al., 2017b), but some studies replace perchloric acid with hydrochloric acid and hydrogen peroxide (e.g., Skonieczny et al., 2019) and others omit any chlorinated acid altogether (e.g., Palchan & Torfstein, 2019). Some digestions are also assisted by pressurized microwave systems (e.g., Thöle et al., 2019).

Prior to the 1990s, in the early development of the proxy, Th and U nuclides were analyzed by alpha spectrometry, a slow process of counting individual nuclide decays that required an additional electroplating step in sample preparation (e.g., Anderson & Fleer, 1982). Today, most measurements are conducted via multicollector ICP-MS. Some studies use single-collector ICP-MS to increase throughput (e.g., Costa & McManus, 2017; Pichat et al., 2004), primarily at the expense of precision on the low-abundance $^{234}\text{U}$. The majority of studies report uncertainties based on the reproducibility of sediment standards (e.g., Costa & McManus, 2017; Palchan & Torfstein, 2019; Thöle et al., 2019), although the specific standards vary from laboratory to laboratory.
2.3. $^{230}$Th in Marine Sediments

The total $^{230}$Th measured in sediment includes not just $^{230}$Th scavenged from the water column (or excess $^{230}$Th, $^{230}$Th$_{ex}$ hereafter) but also lithogenic and authigenic nonexcess components. The $^{230}$Th$_{ex}$ is calculated by subtracting the contributions of the lithogenic and authigenic $^{230}$Th activities as follows (Henderson & Anderson, 2003):

$$A_{^{230}Th_{ex}} = A_{^{230}Th_{total}} - A_{^{230}Th_{lith}} - A_{^{230}Th_{auth}} \quad (2)$$

Lithogenic $^{230}$Th ($^{230}$Th$_{lith}$) is derived from the incorporation of continental material, hereafter referred to as lithogenic material, into marine sediments. Nonexcess $^{230}$Th is also derived from the in situ decay of authigenic U, which is precipitated under reducing sedimentary conditions. While this $^{230}$Th is not authigenic sensu stricto, we refer to it as authigenic $^{230}$Th ($^{230}$Th$_{auth}$) for simplicity. These two additional sources of $^{230}$Th must be quantified and subtracted, following the procedures detailed below.

2.3.1. Lithogenic Correction

Depending on its location, a core site can receive substantial lithogenic input from rivers, aeolian dust, and/or iceberg discharge. Lithogenic material contains lattice-bound $^{232}$Th (10.7 ppm for upper continental crust on average; Taylor & McLennan, 1995), while biogenic material (e.g., calcium carbonate and opal) is virtually devoid of this isotope. Thus, the lithogenic $^{230}$Th activity can be determined as follows:

$$A_{^{230}Th_{lith}} = \frac{A_{^{238}U}}{A_{^{232}Th}} \cdot A_{^{232}Th}_{lith} \quad (3)$$

where ($A_{^{238}U}/A_{^{232}Th})_{lith}$ is the lithogenic ratio of $^{238}$U to $^{232}$Th in activity units. This correction relies on three assumptions: (i) the lithogenic fraction of the sediment is at secular equilibrium for $^{238}$U and $^{230}$Th, (ii) the ($A_{^{238}U}/A_{^{232}Th})_{lith}$ is known, and (iii) all measured $^{232}$Th is lattice bound rather than adsorbed. Generally, studies calculating $^{230}$Th$_{ex}$ use prescribed ($A_{^{238}U}/A_{^{232}Th})_{lith}$ based on the recommendations summarized by Henderson and Anderson (2003): Atlantic (0.6 ± 0.1), Pacific (0.7 ± 0.1), and Southern (0.4 ± 0.1) Oceans. However, the ($A_{^{238}U}/A_{^{232}Th})_{lith}$ values that have been employed within each basin vary substantially among publications (supporting information, Figure S1), rendering data comparison, compilation, and modeling difficult.

While the bulk silicate Earth ($A_{^{238}U}/A_{^{232}Th})_{lith}$ is ~0.74 (Allegre et al., 1986), U and Th can be fractionated in continental materials by igneous processes, chemical weathering, transport, and sedimentation. In particular, U dissolves much more easily in oxygenated water than Th, so that, for example, deeply weathered continental rocks are expected to be depleted in U relative to Th. This mobilization of U contributes to the highly variable ($A_{^{238}U}/A_{^{232}Th})_{lith}$ observed in sedimentary rocks (0.15 to 155; Adams & Weaver, 1958) compared to fresh, unweathered igneous rocks (0.4 to 1.6; Bourdon & Sims, 2003). It can thus be difficult to predict the relevant ($A_{^{238}U}/A_{^{232}Th})_{lith}$ for deep-sea sediments, which may integrate material from multiple geological sources with highly variable ($A_{^{238}U}/A_{^{232}Th})_{lith}$. Several studies have highlighted that the most appropriate ($A_{^{238}U}/A_{^{232}Th})_{lith}$ value may diverge substantially from the recommended value for a given ocean basin (Costa & McManus, 2017; Missiaen et al., 2018; Pichat et al., 2004; Walter et al., 1997). For instance, input from young volcanic provinces and/or inland regions with high runoff may locally deviate the ($A_{^{238}U}/A_{^{232}Th})_{lith}$ value from that of the basin average (Pichat et al., 2004). At the same time, the observed variability in deep-sea sediments is markedly lower than the variability reported for potential parent rock material (Missiaen et al., 2018), suggesting that the integrative nature of marine deep-sea sediment mixes individual lithogenic signals toward a more homogeneous ($A_{^{238}U}/A_{^{232}Th})_{lith}$ range.

Early approaches to refining ($A_{^{238}U}/A_{^{232}Th})_{lith}$ estimates either measured bulk sediment ratios in predominantly lithogenic sediment (Veiga-Pires & Hillaire-Marcel, 1999) or applied a range of ($A_{^{238}U}/A_{^{232}Th})_{lith}$ based on a compilation of possible lithogenic sources (Pichat et al., 2004). Later studies argued that the minimum measured bulk sediment ($A_{^{238}U}/A_{^{232}Th})_{lith}$ over the studied time series would be the closest estimate to the actual ($A_{^{238}U}/A_{^{232}Th})_{lith}$ (Böhm et al., 2015; Costa & McManus, 2017;
Lippold et al., 2009; Mulitza et al., 2017). Another approach has been to use \(A_{234U}/A_{238U}\) to identify sediment with no authigenic contribution, within which the bulk sediment \(A_{238U}/A_{232Th}\) would be a more accurate estimate for the local lithogenic value (Bourne et al., 2012). This approach assumes a seawater \(A_{234U}/A_{238U}\) of 1.1468 (Andersen et al., 2010) and a lithogenic \(A_{234U}/A_{238U}\) of 1 (i.e., secular equilibrium). However, \(A_{234U}/A_{238U}\) ratios below secular equilibrium are known to occur frequently, particularly in slowly accumulating deep-sea sediments (e.g., DePaolo et al., 2012; Ku, 1965), and so this approach should be used with caution. Finally, sequential sediment leaching has been applied to isolate the authigenic fraction of sediment, and it has demonstrated substantial variability (0.4 to 0.7) in \(A_{238U}/A_{232Th}\) within a single sediment core in the Atlantic (Missiaen et al., 2018). This range exceeds the uncertainty that is usually associated with \(A_{238U}/A_{232Th}\) in the literature (±0.1) and presents a challenge to the treatment of \(A_{238U}/A_{232Th}\) as a constant through time.

Refining \(A_{238U}/A_{232Th}\) is important because of the propagating effects on \(230Th_{xs}\) calculations, particularly in sediment with a high proportion of lithogenic material (Burckel et al., 2016; Guihou et al., 2018; Lippold et al., 2012). An accurate evaluation of the \(A_{238U}/A_{232Th}\) value can be key to properly reconstructing the amplitude and timing of \(230Th_{xs}\) changes, especially for shallow coastal sediment cores, which receive significant lithogenic inputs (>30% of the total sediments). Future studies should aim to (i) develop a simpler method to evaluate the temporal \(A_{238U}/A_{232Th}\) variations from routine measurements and (ii) further investigate the importance of adsorbed versus lattice-bound \(232Th\) for lithogenic corrections when using the \(230Th\) normalization technique. Given the available tools, we recommend the following treatment for future \(230Th_{xs}\) records: (i) evaluate the detrital contribution to the sediment (% lithogenic) using the measured bulk \(232Th\) activities; (ii) assess the sensitivity/robustness of the \(230Th_{xs}\) record to changes in the \(A_{238U}/A_{232Th}\) value; (iii) include appropriate uncertainties on \(A_{238U}/A_{232Th}\) of, for example, at least 10% at 2σ; and (iv) propagate the uncertainties and potential temporal variability in \(A_{238U}/A_{232Th}\) into the calculations for \(230Th_{xs}\).

### 2.3.2. Authigenic Correction

After subtraction of lithogenic \(230Th\), the residual \(230Th\) concentration must be corrected for authigenic \(230Th\), which may have accumulated in the sediment due to the decay of nonlithogenic uranium, commonly known as authigenic uranium (\(U_{auth}\)). The precipitation of \(U_{auth}\) in marine sediments is the primary sink of U from the ocean (Klinkhammer & Palmer, 1991; McManus et al., 2005) and occurs when soluble U (VI) is reduced to U (IV) (Anderson, 1982). This transformation is thought to be microbially mediated (Francis et al., 1994; Ganesh et al., 1997; Lovley et al., 1991; Sani et al., 2004) and occurs in reducing porewaters where oxygen is limited by a combination of low bottom water oxygen and/or a high organic carbon rain rate (Finneran et al., 2002; McManus et al., 2005). As the reduction and precipitation of U begins in porewaters, it creates a concentration gradient between high-U seawater and low-U porewater that transfers U from seawater to sediment as long as reducing conditions are maintained (Anderson et al., 1989; Barnes & Cochran, 1990). In some sedimentary environments, typically those characterized by well-oxygenated bottom water and low organic productivity, no \(U_{auth}\) is found and the magnitude of the authigenic correction will be negligible. In other environments, particularly those where porewater redox conditions are variable and the conditions for \(U_{auth}\) precipitation are periodically or continuously sustained, uncertainties arising from the \(U_{auth}\) correction can be substantial.

Assuming that the lithogenic end-member is known for a site (see section 2.3.1), \(U_{auth}\) activity \(A_{238U}^{auth}\) can be quantified as follows:

\[
A_{238U}^{auth} = A_{238U}^{total} - \left( \frac{A_{238U}}{A_{232Th}} \right)_{lith} \cdot A_{232Th}
\]

This \(U_{auth}\) then decays to \(230Th_{auth}\) since the time of deposition \(t\), as in equation (5a) and as described in Francois et al. (2004). However, because this process does not occur at secular equilibrium, the ingrowth rate itself will vary as a function of time. To account for this disequilibrium ingrowth, we incorporate the \(230Th\)
age equation, as used for dating corals and speleothems (Edwards et al., 2003), into the $^{230}\text{Th}_{\text{auth}}$ calculation (equation (5b); Henderson & Anderson, 2003).

$$A_{^{230}\text{Th}_{\text{auth}}} = A_{^{238}\text{U}}^\text{auth} \left( 1 - e^{-\lambda_{230}^* t} \right)$$  \hspace{1cm} (5a)

$$A_{^{230}\text{Th}_{\text{auth}}} = A_{^{238}\text{U}}^\text{auth} \left( 1 - e^{-\lambda_{230}^* t} \right) + \frac{\lambda_{230}^*}{\lambda_{234}^* - \lambda_{230}^*} \left(e^{-\lambda_{234}^* t} - e^{-\lambda_{230}^* t}\right) \left( \frac{A_{^{234}\text{U}}}{A_{^{238}\text{U}}}^{\text{SW}} - 1 \right)$$  \hspace{1cm} (5b)

The divergence between equations (5a) and (5b) increases with higher lithogenic corrections (section 2.3.1) and lower $U_{\text{auth}}$ activity. For example, the $^{230}\text{Th}_{\text{auth}}$ activity would be about 0.8% lower for equation (5a) than for equation (5b) for a theoretical 400 ka sediment with bulk $^{238}\text{U}$ of 9 dpm/g, bulk $^{232}\text{Th}$ of 1 dpm/g, and $(A_{^{238}\text{U}}/A_{^{232}\text{Th}})_{\text{ lith }}$ of 0.6. If instead bulk $^{238}\text{U}$ were 3 dpm/g, holding all other variables constant, the difference in $^{230}\text{Th}_{\text{auth}}$ activity between the two equations would increase to about 2.6% lower for equation (5a) than for equation (5b). Equation (5a) always underestimates $^{230}\text{Th}_{\text{auth}}$, relative to equation (5b).

The use of equation (5b) involves two assumptions: The first one is that the age of the sediment and the age of the $U_{\text{auth}}$ deposition are contemporaneous. This assumption is almost certainly an oversimplification as $U_{\text{auth}}$ is deposited at the porewater redox front beneath the sediment-water interface, making the age of the $U_{\text{auth}}$ inherently younger than the sediment in which it is measured. However, given the long half-life of $U$ and considering average marine sedimentation rates of a few centimeters per thousand years, the age offset between the sediment and $U_{\text{auth}}$ is typically negligible. A second, potentially more critical, assumption is that all of the $U_{\text{auth}}$ that contributed to the production of $^{230}\text{Th}$ is still present in the sediment. If post depositional burn down (i.e., diagenetic remobilization) removed a substantial fraction of $U_{\text{auth}}$ after the time of initial deposition, the magnitude of the correction for ingrown $^{230}\text{Th}$ may be too small (Jacobel et al., 2017a).

Loss of $U_{\text{auth}}$ is primarily problematic in cores with low sedimentation rates (<2 cm/kyr; Costa et al., 2018; Mangini et al., 2001). The number of records in which it has been clearly demonstrated as problematic is thus far small and restricted to regions that have experienced large changes in bottom water oxygen and/or organic matter fluxes (Hayes et al., 2014; Jacobel et al., 2017a).

### 2.4. $^{230}\text{Th}$ Normalization

After scavenging from the water column and deposition on the seafloor, the excess $^{230}\text{Th}$ activity in the sediment decreases with a half-life of 75.584 kyr (Cheng et al., 2013). In order to calculate the $^{230}\text{Th}_{\text{eq}}$ activity at the time of initial sediment deposition, $^{230}\text{Th}_{\text{eq}}^0$, this decay must be accounted for by using independent chronological constraints (such as oxygen isotope stratigraphy or radiocarbon dates) and the classic radio-decay equation:

$$A_{^{230}\text{Th}_{\text{eq}}^0} = A_{^{230}\text{Th}_{\text{eq}}} e^{-\lambda_{230}^* t}$$  \hspace{1cm} (6)

The long half-life of $^{230}\text{Th}$ allows utilization of the proxy as far back as 500,000 years, although the errors expand in increasingly older sediments. These errors are largely due to uncertainties in the lithogenic and authigenic corrections. A big advantage of $^{230}\text{Th}$ normalization over fluxes derived using stratigraphic age control points is that $^{230}\text{Th}$-normalized fluxes are relatively insensitive to errors in the age model (<1% error in flux for a 1 kyr error in age). Altogether, propagated uncertainties on $^{230}\text{Th}_{\text{eq}}^0$ activities are typically less than 5% for the last 30 kyr, and often less than 2%, including analytical uncertainties, authigenic corrections, lithogenic corrections, and decay corrections.

The $^{230}\text{Th}$-normalized mass flux (or preserved rain rate) of sediment (g/cm$^2$/kyr) may then be calculated (Bacon, 1984; Suman & Bacon, 1989):

$$\text{Mass Flux} = \frac{\beta_{^{230}\text{Th}} z}{A_{^{230}\text{Th}_{\text{eq}}}}$$  \hspace{1cm} (7)

where $\beta_{^{230}}$ is the production rate (as in section 2.1), $z$ is the water depth in centimeters, and the term $\beta_{^{230}} z$ is equivalent to the integrated $^{230}\text{Th}$ production ($P$) in the overlying water column. At relatively shallow sites (<1,200 m water depth), it may be important to consider glacial-interglacial changes in sea level (e.g., Grant...
et al., 2014), but this adjustment is generally only necessary when the change in sea level (~120 m, on average, during the Last Glacial Maximum [LGM]) comprises 10% or more of the modern water column depth (e.g., in the Bahamas, Slowey & Curry, 1991; Williams et al., 2016; and in the Red Sea, Palchan & Torfstein, 2019). At deeper sites, the compensatory increase in salinity driven by reduced ocean volume at sea level low stands (e.g., Adkins et al., 2002) largely negates any change in P driven by changes in water column depth (z) by increasing the production rate (P_{230Th}; see section 2.1), and the effects essentially cancel at the mean depth of the ocean (McManus et al., 1998). For example, at Bermuda Rise, modern P at 4,584 m water depth and S = 34.885 is 11.70 dpm/cm²kyr. During the LGM, assuming 120 m lower water depth (4,464 m) and S = 35.84 (Adkins et al., 2002), glacial P was 11.75 dpm/cm²kyr, a negligible difference of only 0.4%.

The $^{230}$Th-normalized flux of any sedimentary component j (e.g., calcium carbonate) can be determined from the fraction of j in the bulk sediments (f_j) by

$$\text{Mass Flux}(j) = f_j \cdot \text{Mass Flux}$$

(8)

2.5. Calculating focusing factors (Ψ)

If the accumulation rate of scavenged $^{230}$Th differs substantially from its inferred production rate (P) in the overlying water column, then the deposited sediment is likely to have been affected by lateral addition/removal (focusing/winnowing) of $^{230}$Th and the associated sediment by bottom currents or downslope redistribution, from a local to larger spatial scale. The degree of sediment focusing (Ψ) can be calculated by comparing the inventory of $^{230}$Th in a dated sediment horizon with the inferred production of $^{230}$Th in the overlying water column over the same time interval (Suman & Bacon, 1989):

$$\Psi = \frac{\rho \int_{z_1}^{z_2} A_{230Th_{bax}}(z) dz}{P(\Delta t)} \approx \frac{\rho \int_{z_1}^{z_2} A_{230Th_{bax}}(z) dz}{P(\Delta t)}$$

(9)

where $\rho$ is the sediment dry bulk density (in g/cm³), $\Delta t$ is the time elapsed (in kyr), and $\Delta z$ is the sediment accumulation (in cm). Where available, sediment density in this compilation is obtained from previously published sources, generally derived from calibrated gamma ray attenuation as determined from a core scanning multisensor track system. Where unspecified, sediment density is arbitrarily set to 0.75 g/cm³, which at least limits the contributed uncertainty to a systematic bias. If the amount of $^{230}$Th buried in the sediment is equal to the amount produced in the water column, then Ψ = 1. Otherwise, Ψ will vary with the addition (focusing, Ψ > 1) or loss (winnowing, Ψ < 1) of sedimentary material.

2.6. Data Compilation

Over 50 years’ (1966–2019) worth of data have been compiled to create the global thorium database ($n = 1,167$) presented here (Adkins et al., 2006; Anderson et al., 2006, Anderson et al., 2009, Anderson et al., 2014, Anderson et al., 2019; Bausch, 2018; Böhm et al., 2015; Bohrman, 2013; Borole, 1993; Bradtmiller et al., 2006, 2007, 2009; Broecker, 2008; Broecker et al., 1993; Brunelle et al., 2007, 2010; Causse & Hillaire-Marcel, 1989; Chase et al., 2003, 2014; Chong et al., 2016; Costa, McManus, & Anderson, 2017; Costa & McManus, 2017; Crusius et al., 2004; Dekov, 1994; Denis et al., 2009; Dezileau et al., 2000, 2004; Durand et al., 2017; Fagel et al., 2002; Francois et al., 1990, Francois et al., 1993; Frank, Eisenhauer, Bonn, et al., 1995, Frank, Eisenhauer, Kubik, et al., 1995, 1996; Fukuda et al., 2013; Galbraith et al., 2007; Geibert et al., 2005, 2009; Gherardi et al., 2009; Hoffmann et al., 2018; Jaccard et al., 2009, 2013; Jacobel et al., 2017a; Jonkers et al., 2015; Kienast et al., 2007; Ku & Broecker, 1966; Kumar et al., 1995; Lam et al., 2013; Lamy et al., 2014; Lao et al., 1992; Lippold et al., 2009, Lippold et al., 2011, Lippold et al., 2012, Lippold et al., 2016; Loubere et al., 2004; Loveley et al., 2017; Lund et al., 2019; Mangini & Dominik, 1978; Marcantoni et al., 1996, Marcantonio et al., 2001, Marcantonio et al., 2014; Martinez-Garcia et al., 2009; McGee et al., 2007, 2010, McGee & Mukhopadhyay, 2013; McManus et al., 1998, 2004; Meier, 2015; Middleton et al., 2020; Missiaen et al., 2018; Mohamed et al., 1996; Mollenhauer et al., 2011; Moran et al., 2005; Mulitza et al., 2008, Mulitza et al., 2017; Muller et al., 2012; Nave et al., 2007; Negre et al., 2010; Neumann & Geibert, 2003; Ng et al., 2018; Not & Hillaire-Marcel Claude, 2010; Nuttini, 2014; Nuttini & Hillaire-Marcel, 2015; Paetsch, 1991; Palchan & Torfstein, 2019; Pichat et al., 2004, Pichat et al., 2014;
Table 1
Summary of Quality Control Criteria and the Subsequent Quality Levels of the Records Within the Database

<table>
<thead>
<tr>
<th>Criterion</th>
<th># Passing cores</th>
<th>% Database</th>
</tr>
</thead>
<tbody>
<tr>
<td>(1) Were raw concentrations (^{230}\text{Th},^{232}\text{Th}, \text{and}^{238}\text{U}) or (^{230}\text{Th}_{\text{exc}}) provided?</td>
<td>1,142</td>
<td>97.9</td>
</tr>
<tr>
<td>(2) Were errors provided for (^{230}\text{Th},^{232}\text{Th}, \text{and}^{238}\text{U}) provided?</td>
<td>778</td>
<td>66.7</td>
</tr>
<tr>
<td>(3) Is chronology specified by either (\delta^{18}\text{O}) or (^{14}\text{C})?</td>
<td>368</td>
<td>31.5</td>
</tr>
</tbody>
</table>

Quality level (number of criteria passed by each record) | # Passing | % |
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>3 = Optimal</td>
<td>263</td>
<td>22.4</td>
</tr>
<tr>
<td>2 = Good</td>
<td>605</td>
<td>51.8</td>
</tr>
<tr>
<td>1 = Fair</td>
<td>279</td>
<td>23.9</td>
</tr>
<tr>
<td>0 = Poor</td>
<td>14</td>
<td>1.2</td>
</tr>
<tr>
<td>Excluded</td>
<td>6</td>
<td>0.5</td>
</tr>
</tbody>
</table>

Plain, 2004; Pourmand et al., 2004, Pourmand et al., 2007; Purcell, 2019; Roberts et al., 2014; Robinson et al., 2008; Rowland et al., 2017; Ruhlmann et al., 1996; Sarin et al., 1979; Saukel, 2011; Scholten et al., 1990, 1994, 2005, Scholten et al., 2008; Serno et al., 2014, 2015; Shiau et al., 2012; Shimmield et al., 1986; Shimmield & Mowbray, 1991; Shimmield & Price, 1988; Singh et al., 2011; Skonieczny et al., 2019, Studer et al., 2015; Sukumaran, 1994; Thiagarajan & McManus, 2019; Thöle et al., 2019; Thomas et al., 2007; Thomson et al., 1993, Thomson et al., 1995, Thomson et al., 1999; Vallieres, 1997; Veeh et al., 1999, 2000; Veiga-Pires & Hillaire-Marcel, 1999; Voigt et al., 2017; Waelbroeck et al., 2018; Walter et al., 1997; Wengler et al., 2019; Williams et al., 2016; Winckler et al., 2008; Yang & Elderfield, 1990; Yang et al., 1995; Yu, 1994; Zhou & McManus, 2020).

Ideally, data were contributed as primary \(^{230}\text{Th},^{232}\text{Th}, \text{and}^{238}\text{U}\) activities so that \(^{230}\text{Th}_{\text{na}}\), mass fluxes, and focusing factors could all be recalculated using consistent formulas and constants (e.g., \(^{230}\text{Th}\) half-life and \(^{230}\text{Th}\) production rate). In an effort toward inclusivity, we have included these records in favor of greater spatial coverage at the expense of some small degree of inconsistency. Variability in constants has been relatively small, with \(^{230}\text{Th}\) half-life changing by less than 10% (80,000 years, Hyde, 1946; vs. 75,587 years, Cheng et al., 2013) and the production rate changing by less than 5% (2.67 \(\times\) \(10^{-5}\) dpm/cm\(^2\) kyr, Francois et al., 2004; vs. 2.562 \(\times\) \(10^{-5}\) dpm/cm\(^2\) kyr, calculated in section 2.1). As we focus on two relatively young time periods, the combined effect of these inconsistencies should yield only minor deviations between the reported values and those that would have been determined using the updated constants applied here.

All data are presented using the age models in the original publications. Generally, the ages are derived from radiocarbon or \(\delta^{18}\text{O}\) stratigraphy, but some data (particularly core tops) have no or only basic age information based on assumed constant sedimentation rates. As the associated uncertainties do not permit precision at millennial time scales, we focus only on the late Holocene (LH) and the LGM and do not consider deglacial events such as Heinrich Stadial 1, for which more stringent age constraints would be required. We conducted sensitivity tests to determine the optimal time frame for the Holocene (0–3, 0–5, and 0–10 ka) and the LGM (19–23, 18–24, and 18.5–23.5 ka) (supporting information Figure S2). The majority of records show only minimal deviation among the different time windows; the main effect of reducing the time window is to limit the number of cores included.

For example, defining the Holocene as 0–3 ka results in 825 cores (71% of the database), whereas defining the Holocene as 0–5 ka yields 982 cores (84% of the database) and as 0–10 ka includes 1,068 cores (92% of the database). Although by definition the Holocene spans 0–11.7 ka, we focus here on the late Holocene (0–5 ka) in order to (i) avoid intra-Holocene climatic variations, (ii) minimize potential incorporation of deglacial values due to age model uncertainties, and (iii) better align the duration (5 kyr) integrated for both time slices.

For the LGM time slice, we tested three different chronozones based on the recommendations from Mix et al. (2001). The effect on the database for the three different time windows is minimal: 297 cores (25% of the database) for 18–24 ka versus 266 cores for 19–23 ka (23% of the database). We select the intermediate option (18.5–23.5 ka, 281 cores, 24% of the database) as the best compromise between sustaining adequate spatial coverage and limiting the potential incorporation of data from the bounding Heinrich events due to age model uncertainties. Henceforth, we define the late Holocene time slice as 0–5 ka and the LGM time slice as 18.5–23.5 ka.

Finally, all data have been screened for quality control, where records were passed when they positively met the criteria described below (Table 1). In addition to raw radionuclide concentrations and age model constraints, we considered whether stated uncertainties were available, the associated magnitude of those errors, and the specified lithogenic corrections. A total of six cores (0.5% of the database) were excluded.
from our analysis because the data failed to pass our criteria. Lithogenic corrections (section 2.3.1) were applied using the \((A_{238U}/A_{232Th})_{th}\) reported in the original publication (supporting information Figure S1), generally ranging between 0.4 and 0.7, but three cores were excluded due to high reported \((A_{238U}/A_{232Th})_{th}\) (greater than 0.8). An additional three core tops without age control were excluded because the resulting calculated fluxes were anomalously high (e.g., by an order of magnitude) compared to neighboring cores with better age constraints.

Overall quality levels were computed by summing each record’s scores on the individual criteria. A record is optimal if it is based on a chronology that is constrained by \(\delta^{18}O\) or \(^{14}C\) and it provides both the raw nuclide concentrations and the associated errors. About one quarter of the records in the database achieved this highest quality level. The large majority of the records in the database are good, passing two of the three criteria, while the remaining quarter are of fair or poor quality. Restricting the database by quality level primarily reduces the spatial coverage, with little impact on the overall data patterns observed (supporting information Figure S3). Time slice data (LH and LGM) and quality screening for all sites are provided in supporting information Table S1. Maps of raw \(^{230}Th_{xs}\) are provided in supporting information Figure S4.

3. \(^{230}Th\) Global Database Results

3.1. Atlantic Ocean

Holocene mass fluxes in the Atlantic (Figures 1a and S5 and Table 2) are highest in the northwestern basin, particularly Baffin Bay and the Labrador Sea, where fluxes reach values as high as ~12 g/cm\(^2\)kyr. In the Nordic Seas, mass fluxes range from <1 to 2.3 g/cm\(^2\)kyr and are generally lower than fluxes in the central northern basin to the south of Iceland, which range from 1 to 5 g/cm\(^2\)kyr. Most equatorial Atlantic sites show mass fluxes between 1 and 2 g/cm\(^2\)kyr, except near the mouth of the Amazon River. Mass fluxes are lower in the South Atlantic than in the North, almost all below 2 g/cm\(^2\)kyr.

During the LGM (Figure 1b), mass fluxes are high (5–20 g/cm\(^2\)kyr) in Baffin Bay, the Labrador Sea, and the western North Atlantic subtropical gyre, all sites that likely received glaciogenic sediment from the Laurentide Ice Sheet. Mass fluxes at sites off western Europe fall between 3 and 5 g/cm\(^2\)kyr, while subtropical sites near the Mid-Atlantic Ridge have the lowest fluxes (1–2 g/cm\(^2\)kyr) in the North Atlantic. Nordic Seas mass fluxes in the LGM range between 2 and 4 g/cm\(^2\)kyr, up to double the Holocene fluxes at these locations. Much of the North Atlantic basin thus shows higher LGM mass flux relative to the Holocene (Figure 1c). South Atlantic LGM fluxes are lower than those in the northern basin: Almost all fall below 2 g/cm\(^2\)kyr, with a few exceptions near the equator or the Southern Ocean. LGM/Holocene mass flux ratios in the South Atlantic are mostly less than or equal to 1, except for a handful of sites showing a doubling to tripling of mass fluxes during the LGM off southern Brazil and in the southern Cape Basin. There is no significant trend in LGM/Holocene mass flux ratios with core site water depth.

Holocene focusing factors tend to be >1 in the western Atlantic (Figure 2a), as well as near continental margins in the Eastern Atlantic. A few sites in the Nordic Seas, southeast of the Labrador Sea, and the equatorial Atlantic show focusing factors <1, but sites with positive focusing factors are much more common, reflecting intentional sampling bias toward regions with rapidly accumulating sediments. LGM focusing factors are lower than 1 in broad regions of the North Atlantic, with focusing only occurring at a few sites in the central western Atlantic or at continental margins in the Eastern Atlantic. There appears to be a latitudinal divide in the North Atlantic, with all but one site north of 50°N having an LGM/Holocene ratio of <1, indicating less focusing in the LGM relative to the Holocene in this region.

3.2. Pacific Ocean

Holocene mass fluxes in the Pacific (Figures 1a and S5 and Table 2) are highest along the continental margins and in the Bering Sea, where fluxes reach up to ~8 g/cm\(^2\)kyr. The lowest mass fluxes occur in the North and South Pacific gyres (<0.5 g/cm\(^2\)kyr). It is possible that even lower mass fluxes may exist in the centers of the gyres that have not yet been sampled and where accumulation rates are so low (0.5 cm/kyr or less, Schmitz et al., 1986) that LGM and Holocene sediments are mixed by bioturbation. Mass fluxes are generally higher in the western Pacific (120°E to 180°E, >1 g/cm\(^2\)kyr) than in the eastern Pacific (−180°E to −70°E, <1 g/cm\(^2\)kyr). Along the equatorial Pacific, Holocene mass fluxes average about 1 g/cm\(^2\)kyr, with a
latitudinal gradient that mirrors the decreasing productivity trend with increasing distance from the nutrient-rich zone of equatorial upwelling. For example, at the Line Islands (approximately −160°E), Holocene mass fluxes along a latitudinal transect of nine sites steadily decrease from ~1.8 g/cm²kyr at the equator (0.2°S) to 0.8 g/cm²kyr at the northernmost site (7.0°N) (Costa et al., 2016b, 2017a; Jacobel et al., 2017b), a trend that is not captured in age model-based mass accumulation rates. The equatorial Pacific

Figure 1. Compiled ²³⁰Th-normalized mass fluxes for (a) the Holocene (0–5 ka), (b) the Last Glacial Maximum (LGM, 18.5–23.5 ka), and (c) the LGM/Holocene mass flux ratio. Dark gray shaded area in (b) and (c) shows exposed land when sea level is 120 m lower. All data and references are provided in supporting information Table S1. Raw ²³⁰Th concentrations maps are provided in supporting information Figure S4.
also manifests a distinct zonal distance effect (supporting information Figure S6), with the lowest mass fluxes occurring in the central equatorial Pacific (~0.5 g/cm²kyr) and increasing more or less monotonically toward the continental margins.

The existing data show that LGM mass fluxes (Figure 1b) were high along the continental margins and low within the North and South Pacific gyres. Unlike in the Atlantic, proximity to ice sheets had only a minor impact on adjacent marine mass fluxes, specifically in the northeastern Pacific near the Cordilleran Ice Sheet (Figure 1b). A more systematic shift in mass fluxes occurred in the Okhotsk Sea and western subarctic Pacific, where glacial mass fluxes were generally >2 g/cm²kyr and as high as 5.8 g/cm²kyr. Along the equator, glacial mass fluxes averaged 1.3 g/cm²kyr and displayed the same zonal and meridional mass flux trends as in the Holocene. The LGM/Holocene mass flux ratio was greater than 1.1 for the majority of the Pacific Basin (Figure 1c). Only parts of the South Pacific and western equatorial Pacific have LGM/Holocene mass flux ratios that are less than or equal to 1. There is no significant trend in LGM/Holocene mass flux ratios with core site water depth.

Constraints on focusing factors in the Pacific are spatially limited, with coverage of the subtropical gyres practically absent. Holocene focusing factors are generally greater than 1 (Figure 2a), and only five sites record winnowing (Ψ = 0.67–0.98) in the Holocene, on the Ontong Java Plateau, the Sulu Basin, and the eastern Japanese coast. Sites in the equatorial Pacific have slightly higher average rates of focusing during the Holocene (Ψ = 2.8) than in the LGM (Ψ = 2.4), but zonal and/or meridional trends in focusing appear less pronounced than those of mass fluxes. In fact, almost all sites in the Pacific show lower rates of sediment focusing during the LGM relative to the Holocene (Figure 2c).

### 3.3. Indian Ocean

Data coverage in the Indian Ocean is relatively low compared to that in other ocean basins (Figures 1, 2, and S5 and Table 2). Coverage in this region is also 4 times greater for the Holocene (n = 83) than for the LGM (n = 21). Holocene mass fluxes increase near the continental margin in the northern Indian Ocean, in the eastern Indian Ocean along the coast of Australia, and in the western Indian Ocean near the southeast coast of Africa (Figure 1a). The few sites that approach the subtropical gyre suggest that Holocene mass fluxes are quite low there, down to 0.15 g/cm²kyr. LGM mass fluxes generally show similar spatial patterns, albeit with far fewer data (Figure 1b). High glacial mass fluxes occurred in the Red Sea (up to 3.46 g/cm²kyr) and

### Table 2

<table>
<thead>
<tr>
<th></th>
<th>Atlantic</th>
<th>Pacific</th>
<th>Indian</th>
<th>Southern</th>
<th>Arctic</th>
<th>Global</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Holocene</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean mass fluxes (g/cm²kyr)</td>
<td>1.92</td>
<td>1.38</td>
<td>1.54</td>
<td>1.16</td>
<td>2.39</td>
<td>1.56</td>
</tr>
<tr>
<td>Median mass fluxes (g/cm²kyr)</td>
<td>1.66</td>
<td>0.84</td>
<td>1.17</td>
<td>0.94</td>
<td>1.48</td>
<td>1.13</td>
</tr>
<tr>
<td>1σ mass fluxes (g/cm²kyr)</td>
<td>1.27</td>
<td>2.62</td>
<td>1.24</td>
<td>1.05</td>
<td>1.82</td>
<td>—</td>
</tr>
<tr>
<td>N</td>
<td>334</td>
<td>136</td>
<td>83</td>
<td>275</td>
<td>12</td>
<td>840</td>
</tr>
<tr>
<td>95% confidence</td>
<td>1.78–2.06</td>
<td>0.93–1.83</td>
<td>1.27–1.81</td>
<td>1.03–1.28</td>
<td>1.34–3.44</td>
<td>1.48–1.65</td>
</tr>
<tr>
<td><strong>LGM</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean mass fluxes (g/cm²kyr)</td>
<td>3.41</td>
<td>1.60</td>
<td>1.83</td>
<td>1.30</td>
<td>0.38</td>
<td>2.00</td>
</tr>
<tr>
<td>Median mass fluxes (g/cm²kyr)</td>
<td>2.03</td>
<td>1.22</td>
<td>1.42</td>
<td>0.72</td>
<td>0.26</td>
<td>1.38</td>
</tr>
<tr>
<td>1σ mass fluxes (g/cm²kyr)</td>
<td>4.73</td>
<td>2.20</td>
<td>1.17</td>
<td>1.38</td>
<td>0.28</td>
<td>—</td>
</tr>
<tr>
<td>N</td>
<td>92</td>
<td>108</td>
<td>21</td>
<td>36</td>
<td>5</td>
<td>262</td>
</tr>
<tr>
<td>95% confidence</td>
<td>2.42–4.39</td>
<td>1.18–2.03</td>
<td>1.32–2.34</td>
<td>0.84–1.76</td>
<td>0.13–0.64</td>
<td>1.81–2.19</td>
</tr>
<tr>
<td><strong>LGM/Hol mass flux ratio</strong></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Mean ratio</td>
<td>1.84</td>
<td>1.39</td>
<td>1.21</td>
<td>1.16</td>
<td>1.60</td>
<td>1.45</td>
</tr>
<tr>
<td>Median ratio</td>
<td>1.37</td>
<td>1.18</td>
<td>1.13</td>
<td>0.93</td>
<td>1.60</td>
<td>1.22</td>
</tr>
<tr>
<td>1σ ratio</td>
<td>1.62</td>
<td>0.63</td>
<td>0.64</td>
<td>0.71</td>
<td>0.84</td>
<td>—</td>
</tr>
<tr>
<td>N</td>
<td>71</td>
<td>67</td>
<td>17</td>
<td>25</td>
<td>2</td>
<td>182</td>
</tr>
<tr>
<td>95% confidence</td>
<td>1.46–2.22</td>
<td>1.23–1.54</td>
<td>0.90–1.52</td>
<td>0.88–1.44</td>
<td>0.42–2.78</td>
<td>1.38–1.53</td>
</tr>
</tbody>
</table>

Note. Global data are computed by weighting each of the five ocean basins by volume. The 95% confidence range is calculated as the mean ± 2 standard errors, where 1 standard error is equivalent to the standard deviation divided by the square root of the number of data points.
Arabian Sea (4.03 g/cm²kyr), while low glacial mass fluxes still characterized the sites off Madagascar (0.90 g/cm²kyr) and near the subtropical gyre (0.23 g/cm²kyr). Of the sites with both LGM and Holocene data, about half experienced lower mass fluxes during the LGM relative to the Holocene (LGM/Hol < 1, Figure 1C), with the possible exception of the core near the subtropical gyre.

**Figure 2.** Sediment focusing for (a) the Holocene (0–5 ka) and (b) the Last Glacial Maximum (LGM, 18.5–23.5 ka), and (c) the LGM/Holocene ratio of sediment focusing ratio. The generally high rates of focusing (>1) are largely due to the sampling bias toward high-accumulation rate sites. Dark gray shaded area in (b) and (c) shows exposed land when sea level is 120 m lower.
Sediment focusing in the Indian Ocean is poorly constrained (Figure 2), and thus, it is difficult to draw any robust conclusions about the remobilization of sediment along the seafloor and how it affects $^{230}$Th burial in this region as a whole. In the Red Sea, sediment focusing is between 1 and 2.2 in the Holocene, and two of the three records have high sediment focusing ($\Psi = 3.9$–6.5) during the LGM. Extreme winnowing ($\Psi = 0.25$) is calculated during the Holocene for one site just to the north of Madagascar. During the LGM, several sites along the west coast of Australia showed no or relatively low degrees of sediment focusing ($\Psi = 0.95$–1.9).

### 3.4. Southern Ocean

The Southern Ocean is defined here as regions south of the Subantarctic Front, comprising all records south of 55°S in the Pacific sector and 50°S in the Atlantic and Indian sectors. Holocene mass fluxes in the Southern Ocean (Figures 1a and S5 and Table 2) are fairly low, with just over half of the sites having values greater than or equal to 1 g/cm²kyr. The highest mass fluxes in the Holocene occur in the Indian sector, at 4.9 g/cm²kyr. Within the Atlantic sector, adjacent sites sometimes show inconsistent results. For example, at 5°–6°E, 50°–53°S, three different mass fluxes are reported: 3.30 g/cm²kyr at ODP1094 (Jaccard et al., 2013; Robinson et al., 2009), 1.64 g/cm²kyr at TN57-13PC4 (Anderson et al., 2009), and 0.66 g/cm²kyr at PS1759 (Geibert et al., 2005; Walter et al., 1997).

Data coverage during the LGM is considerably reduced, dropping to about 13% of what is available for the Holocene (Figure 1b and Table 2). Most of these records (26 of 35) are in the Pacific sector, with only one in the Indian sector. Mass fluxes are highest (up to 5.7 g/cm²kyr) in the Atlantic sector, particularly at the sites near the Weddell Sea, which may have received enhanced delivery of ice-rafted debris during the glacial period. Many sites in the Atlantic sector had higher LGM mass flux relative to the Holocene (Figure 1c). In the Pacific sector, glacial mass fluxes were generally lower in the Antarctic Zone (LGM/Hol = 0.70–0.97, 95% confidence) and higher in the Subantarctic Zone (LGM/Hol = 0.93–2.10, 95% confidence).

As in the Indian Ocean, sediment focusing in the Southern Ocean is poorly constrained (Figure 2). In the Holocene (Figure 2a), sediment focusing is generally above 1, with particularly high values ($\Psi \geq 10$) at two sites in the Atlantic sector and two sites in the Indian sector. The three sites with sediment winnowing ($\Psi = 0.34$–0.86) in the Holocene are all in the Pacific sector, and they range from the margin of the Southern Ocean (50°S) into the Antarctic Zone (64°S). In the LGM, sediment focusing ($\Psi > 1$) occurs at all sites except the Weddell Sea ($\Psi = 0.13$) and south of New Zealand ($\Psi = 0.88$). This same site from New Zealand is the only one that shows greater sediment focusing during the LGM (LGM/Hol of 1.3). The other five sites that have focusing factors in both the Holocene and the LGM all show substantially lower rates of sediment focusing during the glacial period (LGM/Hol = 0.09–0.52).

### 3.5. Arctic

In the Arctic Ocean, mass flux varies from 0.13 to 7.24 g/cm²kyr during the Holocene (Figures 1a and S5 and Table 2), and the highest mass fluxes are located close to the coast in the Canadian Arctic Shelf. In contrast, cores located in the central Arctic ocean have mass fluxes ranging from 0.2 to 1.5 g/cm²kyr, with no distinction between Amerasian and Eurasian basins. Variations in mass flux within the central Arctic appear to be linked to both water depth and physiographic features of the core location (e.g., proximity to ridge). The spatial difference in mass fluxes between central Arctic and coastal area reflects the large difference of sediment input within the different parts of the Arctic Ocean.

Sedimentation in the Arctic Ocean during the LGM was limited (Figure 1B), and several cores may even contain a hiatus during this period (Not & Hillaire-Marcel, 2012; Poore et al., 1999). Therefore, mass flux data for the LGM are quite sparse. Generally lower mass fluxes (0.12–0.80 g/cm²kyr) occurred during the LGM in comparison with the Holocene, which is consistent with a slowdown of the sedimentation process in the Arctic Ocean during colder periods characterized by extended ice cover. Near the Canadian Arctic Shelf, where Holocene mass fluxes are relatively high for the Arctic, the resulting LGM/Holocene mass flux ratios are all below 1 (LGM/Hol = 0.37–0.92; Figure 1c).

Sediment focusing in the Arctic is poorly constrained (Figure 2), and thus, it is difficult to draw any robust conclusions about the remobilization of sediment along the seafloor and how it affects $^{230}$Th.
burial in this region as a whole. At 140–150°E, sediment focusing in the Holocene is minimal (Ψ = 1.09–1.38; Figure 2a), but higher degrees of focusing (Ψ = 1.46–3.34) are found near the Bering Strait (−175°E to 175°E). Only one site, north of western Greenland (−61°E), contained sufficient data to assess sediment focusing in the LGM (Figure 2b), during which this site demonstrated sediment focusing (Ψ = 2.03) within the range observed in the Holocene.

4. Sediment Fluxes Under LGM Climate Conditions

The response in sedimentary mass fluxes to glacial climate conditions varied among individual ocean basins (Table 2). The Atlantic is the only ocean with significantly higher mass fluxes during the LGM (2.42–4.39 g/cm²kyr; all ranges are at the 95% confidence level) relative to the Holocene (1.77–2.05 g/cm²kyr). The Pacific, Indian, and Southern Oceans also had higher mass fluxes during the LGM relative to the Holocene, but they were not significantly greater. In contrast to the other basins, the Arctic is the only ocean with significantly lower mass fluxes during the LGM (0.13–0.63 g/cm²kyr) relative to the Holocene (1.34–3.44 g/cm²kyr).

This basin-specific variability in glacial mass flux suggests that multiple mechanisms were simultaneously active but heterogeneously distributed in altering mass fluxes. For example, in the North Atlantic, mass fluxes were likely high due to enhanced glacial terrigenous input including ice-rafted debris (e.g., McManus et al., 1998) that more than compensated for lower CaCO₃ burial (e.g., Crowley, 1985). Lower glacial sea level may have allowed more efficient transport of sediments to the deep sea rather than storage on the continental shelves (Francois & Bacon, 1991). This process would have been more effective at increasing basin-wide mass fluxes in the Atlantic, where the narrow basin width would concentrate these “additional” sediments into a more confined region than in, for example, the Pacific. Globally higher dust flux (e.g., Kienast et al., 2016; Kohfeld & Harrison, 2001) also may have contributed to higher mass fluxes in the Atlantic, particularly downwind of the Sahara. In other ocean basins, windblown dust deposition is only a small net contributor to sediment fluxes. In the equatorial Pacific, mass fluxes were likely higher due to enhanced glacial CaCO₃ preservation (e.g., Anderson et al., 2008; Cartapanis et al., 2018; Farrell & Prell, 1989), and in the western Pacific warm pool, due to land exposure and erosion. Mass fluxes may have been lower in the Arctic and certain sites in the Southern Ocean due to the inhibiting effects of sea ice formation on the biological production of particles. Generally, glacial sediment fluxes were higher almost everywhere in the Northern Hemisphere, possibly because of the erosive presence of continental ice sheets. Large portions of the South Pacific, South Atlantic, Southern Ocean, and Indian Ocean have few if any constraints during the LGM. Whether or not mass fluxes in these regions may have changed in the past is still an open question for future research.

5. Modeling ²³⁰Th: State of the Art

5.1. From Simple 1-D Scavenging Models to Integration of ²³⁰Th Into Earth System Models

Modeling ²³⁰Th in the ocean began with 1-D analytical models (Bacon & Anderson, 1982; Clegg et al., 1991; Nozaki et al., 1981; Nozaki & Horibe, 1983; Roy-Barman et al., 1996), which demonstrated that only reversible scavenging was able to reproduce the observations of both the dissolved and particulate vertical profiles. To explain complex (nonlinear) ²³⁰Th profiles, more elaborate box models were developed that could account for different transport conditions and particle regimes under different ocean conditions, such as upwelling of deep water masses in the Southern Ocean (Chase et al., 2003; Roy-Barman et al., 2019; Rutgers van der Loeff et al., 2016; Rutgers van der Loeff & Berger, 1993; Venchiarutti et al., 2011), convection of deep water masses in the North Atlantic (Moran et al., 1995, 1997; Vogler et al., 1998), and lateral exchange between open ocean and ocean margin regimes (Anderson et al., 1983; Lao et al., 1992; Roy-Barman et al., 2009). More recently, ²³⁰Th has also been integrated into complex geographic schemes in 2-D models (Luo et al., 2010; Marchal et al., 2000), in 3-D models of intermediate complexity (Henderson et al., 1999; Missiaen et al., 2019; Rempfer et al., 2017; Siddall et al., 2007, 2005), and in global climate models (GCMS) (Dutay et al., 2009; Gu & Liu, 2017; Rogan et al., 2016; Van Hulten et al., 2018). The models of intermediate complexity are generally computationally efficient (i.e., able to produce 1,000 years of simulations in a few hours), but their spatial resolution is rather coarse (e.g., Henderson et al., 1999; Siddall et al., 2005) and/or the particle representation contains strong simplifications (e.g., Missiaen et al.,...
Thorium removal to sediments is primarily driven by two major parameters: (1) the particle fluxes (concentrations and settling speed) and (2) the partition coefficients, or the affinity of each particle type for scavenging Th. Most early models did not parameterize different particle types but instead used a homogeneous particle field (Henderson et al., 1999; Luo et al., 2010). Now, most 3-D models consider at least three different biogenic particle types: calcium carbonate (CaCO$_3$), particulate organic carbon, and opal. Some models also include lithogenic (dust and fluvial) particles (Siddall et al., 2005; Van Hulten et al., 2018) or other aerosols like volcanic ash (Rogan et al., 2016), but uncertainty regarding their influence on Th scavenging has justified their omission in other models (e.g., Gu & Liu, 2017; Missiaen et al., 2019). At the same time, all models face challenges in reproducing the observed particle size spectrum. Many models still employ one single particle size class with a uniform settling speed of 1,000 m/yr, which is consistent with estimates derived from observed particulate $^{230}$Th profiles (Anderson et al., 2016; Gdaniec et al., 2018; Krishnaswami et al., 1976). NEMO-PISCES (Dutay et al., 2009; Van Hulten et al., 2018) accounts for two particle size classes: fast (18,200 m/yr) and slow (730 m/yr) sinking particles. On the other hand, HAMOCC has a parametrization of the scavenging coefficients that implicitly accounts for the observed variability in particle sizes (Heinze et al., 2006; Henderson et al., 1999).

Thorium scavenging has been represented in two distinct ways in models. The first method considers only one tracer, the total $^{230}$Th activity for transport (advection and diffusion), and partitions it into dissolved and particulate activities using equilibrium partition coefficients (e.g., Dutay et al., 2009; Gu & Liu, 2017; Siddall et al., 2005, 2007). The second method considers dissolved and particulate $^{230}$Th activities as two tracers transported by the model and regulates the exchange between the two phases using adsorption and desorption rate constants (e.g., Marchal et al., 2000; Missiaen et al., 2019; Rempfer et al., 2017) or partition coefficients (Henderson et al., 1999; Van Hulten et al., 2018). Most models (Gu & Liu, 2017; Rempfer et al., 2017; Siddall et al., 2005, 2007) initiate with partition coefficients ($K_d$) that were determined on the Joint Global Ocean Flux Study (JGOFS) campaigns in the Pacific Ocean (Chase et al., 2002), but due to the large uncertainties on these observations (Chase et al., 2002; Chase & Anderson, 2004; Hayes, Anderson, Fleisher, Vivancos, et al., 2015; Luo & Ku, 2004; Roy-Barman et al., 2005), the models subsequently treat the $K_d$ as tunable parameters. HAMOCC (Heinze et al., 2018) is the only model currently updated with the newer scavenging coefficients from the Atlantic GEOTRACES section (Hayes, Anderson, Fleisher, Huang, et al., 2015). In NEMO-PISCES (Dutay et al., 2009; Van Hulten et al., 2018) and iLOVECLIM (Missiaen et al., 2019), the $^{230}$Th scavenging coefficients are scaled to the particle fluxes rather than based on data from either JGOFS or GEOTRACES.

Incorporation of additional scavenging processes (such as described in sections 6.2 and 6.3) is only just beginning. To date, only Bern 3D (Rempfer et al., 2017) accounts for particle resuspension in benthic nepheloid layers, which they found improved their model-data agreement for dissolved and particulate water column $^{230}$Th activities. The impacts of hydrothermal scavenging have not yet been considered in any $^{230}$Th models. Finally, all the above cited studies mostly focus on reproducing the modern dissolved and particulate water column $^{230}$Th. Although some studies also performed sensitivity tests for changes in settling speed (Siddall et al., 2005), scavenging coefficients (Gu & Liu, 2017; Siddall et al., 2005), or circulation strength (e.g., Gu & Liu, 2017; Missiaen et al., 2019; Rempfer et al., 2017; Siddall et al., 2007), no simulations are yet available for $^{230}$Th scavenging under past climate conditions.

5.2. Modern Th Flux to the Sediments in Models

In this section we compare the preindustrial outputs of two GCMs (Figure 3), NEMO-PISCES (Dutay et al., 2009; Van Hulten et al., 2018) and Community Earth System Model (CESM) (Gu & Liu, 2017), and two models of intermediate complexity, iLOVECLIM (Missiaen et al., 2019) and HAMOCC (Heinze et al., 2006, 2018). We evaluate the $^{230}$Th flux to the sediments ($F$) normalized by the production of $^{230}$Th in the overlying water column ($P$). $F/P$ values equal to 1 would indicate that $^{230}$Th is buried in the sediments at the rate at which it is produced, whereas $F/P$ values above or below 1 indicate that $^{230}$Th has been transported away from its production site, either by ocean advection or by diffusive fluxes along concentration gradients. Observations of $F/P$ are sparse as they require independent flux calculations, from either bottom-moored
sediment traps (below 2,000 m, e.g., Buesseler et al., 1992, 2007; Gardner, 2000) or an independent constant flux proxy (such as ^3He) in sediments. We compare the model output data with bottom-moored sediment traps primarily in the North Pacific and North Atlantic (Yu et al., 2001), the equatorial Pacific (Lyle et al., 2014), Southern Ocean (Chase et al., 2003), and the Arabian Sea (Scholten et al., 2005).

Figure 3. Modeled $^{230}$Th flux to the sediment, normalized to water column production ($F/P$). (a) iLOVECLIM (Missiaen et al., 2019). (b) HAMOCC (Heinze et al., 2018). (c) CESM (Gu & Liu, 2017). (d) NEMO-PISCES (Dutay et al., 2009; Van Hulten et al., 2018). Overlain circles show flux to production ($F/P$) ratios measured at bottom-moored sediment traps (Chase et al., 2003; Lyle et al., 2014; Scholten et al., 2005; Yu et al., 2001). Color bar is the same for all four maps and composite (f; see below). (e) Comparison of $F/P$ measured in bottom-moored sediment traps to the modeled $F/P$ at each site. Dashed black line shows 1:1. The highest correlation ($R^2 = 0.22$) occurs with the NEMO-PISCES model. Both Nemo-PISCES and CESM adequately reproduce the sign ($>1$ or $<1$) for more than 61% of the sediment traps, while iLOVECLIM and HAMOCC reproduce the sign 50% or less of the sediment traps. At the same time, iLOVECLIM, HAMOCC, and CESM more realistically predict deviations from the theoretical $F/P$ of 1 than the more extreme variability observed in NEMO-PISCES. (f) Composite of the four model outputs. Hatched regions highlight where at least three models agree on the sign ($F/P$ of $>1$ or $<1$). A full size version of the composite map (f) is provided in supporting information Figure S7.

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$^{230}$Th burial in oligotrophic and low-productivity areas. Interestingly, HAMOCC (Figure 3b) is the only model that displays a completely different pattern with spatially homogenous $F/P$ except in the North Atlantic and in the Southern Ocean, where $^{230}$Th is preferentially removed. This difference may be related to the choice of the scavenging coefficients, which are similar in iLOVECLIM, CESM, and NEMO-PISCES and different in HAMOCC, and/or to the particle fields themselves. Modeled $F/P$ broadly agrees with sediment trap observations (Figure 3e), in that continental margins tend to have high values and oligotrophic values have low values. The best correlation between data and model occurs in NEMO-PISCES, although the skill remains modest (Figure 3d; $R^2 = 0.22$). Substantial divergence between estimates is more likely at shallow depths (<2.5 km; Figure 3e); otherwise, the models tend to slightly overestimate the $F/P$ relative to the sediment traps. Overall, iLOVECLIM and HAMOCC have $F/P$ ratios that are closest to 1, suggesting minimal deviation of $^{230}$Th burial from $^{230}$Th production, while NEMO-PISCES has the largest divergence from 1.

This brief model comparison raises a few questions:

1. How do particle parameterization, settling speed, and scavenging coefficients influence the intermodel agreement or disagreement?
2. Are the models too sensitive to scavenging by biogenic particles versus lithogenic particles?
3. How do the particulate and dissolved concentrations at the bottom-ocean grid cell compare to individual GEOTRACES profiles including anomalous features like benthic nepheloid layers or hydrothermal scavenging?

Answering those questions would require a full model intercomparison project comparing the fields of dissolved and particulate activities to the available GEOTRACES data as well as core top measurements, which is beyond the scope of this paper. Yet the work presented here highlights the diversity in $^{230}$Th modeling and demonstrates that modeling studies can be helpful in evaluating the assumptions and determining the spatial efficacy of $^{230}$Th normalization.

## 6. Uncertainties and Limits of the Constant $^{230}$Th Flux Model

### 6.1. Boundary Scavenging

The application of $^{230}$Th as a constant flux proxy relies on the assumption that net lateral transport by eddy diffusion and advection in the water column are negligible components of the local $^{230}$Th mass balance. This assumption is often presumed to be justified a priori (Francois et al., 2004), since the residence time of $^{230}$Th averaged over the full water column is 20–40 years, while the time scale for basin-scale mixing and deep-ocean ventilation is on the order of centuries (Sarmiento & Gruber, 2006). However, spatial gradients in scavenging intensity throughout the ocean (Bacon, 1988) may more efficiently remove scavenging-prone elements from the water column in a high-particle flux zone compared to an adjacent low-particle-flux zone (Anderson et al., 1983, 1994; Roy-Barman et al., 2009). This situation creates a concentration gradient in the water column that in turn generates a dispersive transport (advection + eddy diffusion) of the affected element toward the high-particle-flux zone, a process called boundary scavenging, as it was first identified at continental boundaries (Bacon et al., 1976). Boundaries are now defined more broadly, and they can include productivity gradients such as those driven by upwelling in the central equatorial Pacific (e.g., Costa et al., 2017a), which can occur far from any continental margin.

Where boundary scavenging can be quantified, the offset between $^{230}$Th burial flux ($F$) and its overlying production ($P$) can be estimated. A simple particle flux module incorporated into a general circulation model suggested that 70% of the sea floor receives a $^{230}$Th flux that is within 30% of the overlying production (Henderson et al., 1999). In other words, in most of the ocean, $^{230}$Th-estimated mass fluxes are within 30% of their true value ($F/P = 0.7–1.3$). Furthermore, the deviations from overlying production are not simply a spatially uniform random error but a predictable property dependent on other oceanographic conditions, such as surface productivity and local particle composition. There was relatively little water column data available to assess this model result at the time of its publication, but the annually averaged flux of $^{230}$Th into deep sediment traps (Bacon et al., 1985; Scholten et al., 2005; Yu et al., 2001) has tended to support the roughly 30% uncertainty in the assumption of deep-sea $^{230}$Th flux.
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With higher precision and sample throughput of seawater thorium measurements afforded by modern mass spectrometry techniques, the GEOTRACES era allowed quantification of spatial concentration gradients in $^{230}$Th and lateral redistribution of $^{230}$Th associated with boundary scavenging. In the upwelling zone off the coast of west Africa, where lateral gradients in export flux are among the steepest globally (DeVries & Weber, 2017), it was concluded that roughly 40% of the water column $^{230}$Th production was being transported from the lower-particle-flux region around the Cape Verde Islands ($F/P = 0.59$) toward the high-particle-flux Mauritian margin ($F/P = 1.41$) (Hayes, Anderson, Fleisher, Vivancos, et al., 2015). In the Atlantic sector of the Southern Ocean, a scavenging gradient results in a net transport of $^{230}$Th from the ice-covered, low-particle-flux Weddell Sea ($F/P = 0.4$) toward the productive and particle-rich Antarctic Circumpolar Current region ($F/P = 1.4$) (Roy-Barman et al., 2019; Rutgers van der Loeff et al., 2016; Walter et al., 2000). In the highly productive eastern equatorial Pacific, it was concluded that roughly 25% of the water column $^{230}$Th production was being transported from the low scavenging area in the Peru Basin ($F/P = 0.76$) toward the high scavenging area in the Panama Basin ($F/P = 1.23$) (Singh et al., 2013). All of these estimates are in qualitative agreement with the modeled transport of $^{230}$Th. Future efforts to quantify boundary scavenging more precisely would benefit from more abundant observations of seawater $^{230}$Th variations across gradients in scavenging intensity in addition to more precise constraints on lateral eddy diffusivity constants, which are also spatially variable.

As a consequence of boundary scavenging, sedimentary $^{230}$Th accumulation may deviate from constant production in the overlying water column. High particle flux may yield surplus sedimentary $^{230}$Th, in which case the mass fluxes calculated using equation (7) would be biased low. In low-particle-flux zones, too little $^{230}$Th may be buried, and mass fluxes may be biased high. The net effect is to reduce the gradient in mass fluxes observed near a boundary. For example, $^{230}$Th-normalized opal fluxes measured along a transect off west Africa would likely underestimate the difference in productivity between a nearshore site and an offshore site. This systematic bias in regions where boundary scavenging is active is likely to make comparison of absolute fluxes difficult to interpret between multiple sites across the boundary. Fortunately, boundary scavenging regions are largely defined by biogeographical provinces, which are unlikely to vary relative to one another in the past (e.g., marginal sites are likely to always have had higher absolute export productivity than gyre sites). Thus, mass fluxes may adequately capture relative trends in local fluxes at any one site over time (e.g., LGM to Holocene changes in opal flux), retaining sufficient efficacy in $^{230}$Th normalization for many paleoceanographic inquiries.

### 6.2. Nepheloid Layers

Nepheloid layers are regions of increased concentrations of suspended sediments near the seafloor. They are generated by high near-seafloor current velocities (>20 cm/s) that exceed the critical shear stress necessary for resuspension of particulates (Gardner et al., 2017; McCave, 1986). Persistent nepheloid layers were found to extend as much as 1,000 m above the seafloor at several stations along the GEOTRACES Section GA03 between Cape Cod and Bermuda (Lam et al., 2015). Because the nepheloid-layer particles in this region are primarily of lithogenic composition (Lam et al., 2015), particulate $^{232}$Th can be used as a tracer of these particles (Figure 4). Near-bottom concentrations of particulate $^{232}$Th within the nepheloid layer are 2 orders of magnitude greater than concentrations measured at middepth (Figure 4d). $^{230}$Th activities also demonstrate anomalous behaviors in the nepheloid layer. While dissolved $^{230}$Th activities increase in a near-linear fashion throughout the upper water column (Figure 4a), as expected for removal of dissolved $^{230}$Th by reversible scavenging (Bacon & Anderson, 1982), $^{230}$Th activity profiles then exhibit a sharp reversal just above the upper extent of the nepheloid layer at each station. This reversal indicates enhanced scavenging and removal of dissolved $^{230}$Th by nepheloid-layer particles, and it is accompanied, in part, by increased concentrations of particulate $^{230}$Th through the same depth interval (Figure 4b).

If nepheloid layers consisted exclusively of locally resuspended sediment for which adsorbed $^{230}$Th remained fully exchangeable with dissolved $^{230}$Th in the surrounding seawater, then nepheloid layers would not be expected to enhance the removal of dissolved $^{230}$Th from bottom water. Yet there is clear evidence that enhanced removal of dissolved $^{230}$Th does occur within nepheloid layers (Figures 4a and 4b). Similar effects of nepheloid layers on dissolved and particulate $^{230}$Th profiles have also been observed in the Nansen basin (GEOTRACES central Arctic section, GN04; Gnadiiec et al., 2019). One possible
mechanism to enhance removal of dissolved $^{230}$Th within nepheloid layers would involve reduced exchangeability of sediment-bound $^{230}$Th. For example, $^{230}$Th could be immobilized on the seabed due to the growth of diagenetic coatings of Fe-Mn oxides or other authigenic phases that lock adsorbed $^{230}$Th into the particle structure. This diagenetic generation of fresh particle surfaces would also enable resuspended sediment to scavenge additional dissolved $^{230}$Th from the water column, consistent with the observed reduction of dissolved $^{230}$Th concentrations to levels well below those predicted by extrapolating trends from shallower depths (Figure 4a). Alternatively, nepheloid layers may consist of sediment resuspended from nearby topographic highs and transported laterally to the sampling locations over a time scale too short to achieve adsorption-desorption equilibrium with dissolved $^{230}$Th in ambient seawater. The observation that the specific activity (dpm/g of particles) of $^{230}$Th on particles within nepheloid layers is substantially less than that for particles above the nepheloid layers (Figure 4c) would be consistent with this mechanism. Lower specific activity of $^{230}$Th on nepheloid-layer particles would also be consistent with sources involving erosion of older sediment from which $^{230}$Th had decayed, as has been observed in the Pacific Ocean (Kadko, 1983). These conditions are not mutually exclusive.

Although the processes occurring within nepheloid layers that enhance the scavenging and removal of dissolved Th remain incompletely defined, results from these stations provide compelling evidence that these processes may impose a strong bias on $^{230}$Th-normalized fluxes. This bias can be illustrated by calculating the $^{230}$Th-normalized flux of lithogenic particles, traced using $^{232}$Th.
continents where the lithogenic material in the water column is supplied mainly as dust, the $^{230}$Th-normalized flux of particulate $^{232}$Th is expected to be uniform throughout the water column (Anderson et al., 2016). This expectation is clearly violated within the nepheloid layers of the NW Atlantic Ocean, where $^{230}$Th-normalized fluxes of particulate $^{232}$Th increase through the nepheloid layer by an order of magnitude at Station 8 and by about a factor of 40 at Station 10 (Figure 4e). Whether this reflects lateral supply of lithogenic particles or an as-yet unidentified violation of the assumptions inherent in $^{230}$Th normalization remains unknown. However, interpreting $^{232}$Th fluxes from the deepest particulate samples collected by in situ filtration (Figure 4e) as recording dust fluxes would greatly overestimate the local supply of $^{230}$Th by dust. Fully assessing the merits of $^{230}$Th normalization in regions of nepheloid layers will require identification of (1) the source(s) of the particles (resuspension of surface sediments locally, erosion and suspension of older sediments, or lateral transport from nearby topographic highs), (2) the diagenetic processes that affect the surface-adsorption properties of resuspended particles, (3) the propagated effects of variability in nepheloid thickness and particle concentration on $^{230}$Th scavenging, and (4) the 3-D mass budget of $^{230}$Th within regions of nepheloid layers. Until then, we recommend consulting global compilations of information about nepheloid layers (Gardner, Richardson, Mishonov, & Biscaye, 2018; Gardner, Mishonov, & Richardson, 2018, 2018) and interpreting $^{230}$Th-normalized fluxes in these regions with caution.

6.3. Hydrothermal Scavenging

In boundary scavenging regions, rapid $^{230}$Th removal creates concentration gradients that drive lateral diffusion of $^{230}$Th toward areas of high particle flux (section 6.1). Similar concentration gradients are found near mid-ocean ridges, where hydrothermal plumes laden with highly reactive metalliferous particles scavenge $^{230}$Th from the water column (Gdaniec et al., 2019; Hayes, Anderson, Fleisher, Huang, et al., 2015; Pavia et al., 2018; Pavia, Anderson, Black, et al., 2019; Valk et al., 2018). In sediment traps within 1 km of an active vent on the northern East Pacific Rise (EPR), $^{230}$Th fluxes were 2–10 times higher than the water column production rate (German et al., 2002), suggesting the possibility of high $^{230}$Th burial rates in near-ridge environments. If a similar process occurs at other locations along the global mid-ocean ridge system,

Figure 5. Compilation of $F/P$ results from the Southern East Pacific Rise (SEPR, blue), Juan de Fuca Ridge (JdFR, red), and Mid-Atlantic Ridge (MAR, yellow). $^{230}$Th flux to production ($F/P$) values are calculated by $^3$He-normalized $^{230}$Th fluxes that are then normalized to the production in the water column, and they are equivalent to flux-to-production ($F/P$) values for sediment traps, as shown in Figure 3. JdFR and MAR data are from Middleton et al. (2020) and SEPR data are from Lund et al. (2019). Legend applies to both panels (a) and (b). (a) $F/P$ versus distance from the ridge crest, where distance is calculated using the corresponding sediment age and spreading rate for each ridge. (b) $F/P$ versus hydrothermal Fe ($Fe_{HT}$) flux ($^3$He normalized). Dashed lines identify $F/P$ of 1 and $Fe_{HT}$ flux of 20 mg/cm²kry (see text). (c) Conceptual model of $^{230}$Th burial, where the $F/P$ at the ridge axis is greater on the SEPR (blue curves) than on the JdFR red curve. Near-axis fluxes are likely supplied by $^{230}$Th diffusion from off-axis, causing $^{230}$Th fluxes less than the water column production rate on the ridge flanks ($F/P < 1$). The off-axis reach of $^{230}$Th deficits is likely greater on the SEPR ($x_2$ or $x_3$) than on the JdFR ($x_1$). The SEPR deficits are shown as dashed lines as they are currently unconstrained.
it is possible that ridges act as an important boundary sink for $^{230}$Th in the open ocean (German et al., 2002; Frank, Eisenhauer, Kubik, et al., 1995). While early efforts to quantify $^{230}$Th burial rates in sediments were limited by a lack of independent flux constraints (Dymond & Veeh, 1975; Shimmield & Price, 1988), more recently, the use of extraterrestrial $^3$He to determine sedimentary $^{230}$Th fluxes has permitted the independent assessment of $^{230}$Th scavenging by hydrothermal plumes. $^3$He-normalized $^{230}$Th fluxes, when compared to production in the water column, provide flux to production ratios ($F/P$) similar to sediment traps but with the unique ability to record changes in $^{230}$Th burial over geologic time scales.

On the southern EPR, $F/P$ values in ridge crest sediments are highly variable (Figure 5). Although most $F/P$ ratios fall in the range of 1.0–1.5, values as high as 4 were found at stratigraphic levels corresponding to the highest hydrothermal iron fluxes (Lund et al., 2019). The strong correlation between $^{230}$Th and Fe fluxes indicates that hydrothermal fallout was the primary driver of the $^{230}$Th signal. The data also imply that scavenging of $^{230}$Th may vary over millennial time scales on the southern EPR, with maximum $^{230}$Th burial rates occurring during the last deglaciation. Elevated deglacial burial fluxes of $^{230}$Th occurred at all three southern EPR sites examined thus far, which are located 8 to 28 km from the ridge crest. If the surplus $^{230}$Th associated with hydrothermal scavenging originated from off-axis locations, the flanks of the southern EPR may have experienced $^{230}$Th deficits during the deglaciation (Lund et al., 2019).

Hydrothermal scavenging can also influence $^{230}$Th burial on mid-ocean ridges with substantially less plume coverage than the southern EPR. For example, on the Juan de Fuca Ridge (JdFR), $^{230}$Th burial rates are lower than the water column production rate ($F/P < 1$) within 10 km of the ridge crest, while $^{230}$Th burial is similar to the production rate ($F/P ~ 1$) in cores located more than 10 km off-axis (Figure 5a) (Costa et al., 2017b; Middleton et al., 2020). Furthermore, surplus $F/P$ values (i.e., $F/P > 1$) are not as high on the JdFR as on the southern EPR, with $F/P$ values not exceeding ~2 in samples deposited within 18 km of the ridge crest. The most important result from the JdFR, however, is the clear documentation of $^{230}$Th deficits ($F/P < 1$), presumably due to lateral diffusion of $^{230}$Th from the water column at the core site location toward areas of high hydrothermal particle flux on the ridge axis (Middleton et al., 2020). Hydrothermally influenced sediment cores recovered within the axial valley of the Mid-Atlantic Ridge also exhibit deficit $^{230}$Th burial rates ($F/P < 1$), as observed on the JdFR (Middleton et al., 2020). However, $^{230}$Th systematics in the Mid-Atlantic Ridge environment may be additionally complicated by along-axis currents and bottom scavenging induced by the unique bathymetry of slow spreading ridges (Middleton et al., 2020).

Interestingly, the hydrothermal threshold for surplus $F/P$ ratios appears to occur at Fe fluxes of ~20 mg/cm$^2$kyr at both the JdFR and the southern EPR. $F/P$ values greater than 1 correspond to hydrothermal Fe fluxes >20 mg/cm$^2$kyr, while $F/P$ values less than 1 correspond to Fe fluxes <20 mg/cm$^2$kyr (Figure 5b). Whether or not this observed threshold is coincidental or meaningful will be borne out through continued research into $^{230}$Th burial in a range of mid-ocean ridge settings. The JdFR and southern EPR data capture two primary consequences of hydrothermal scavenging for $^{230}$Th, including higher than expected fluxes near the ridge axis and lower than expected fluxes at more distal locations.

Based on the results from each ridge, we present a conceptual model for how hydrothermal activity modifies $^{230}$Th burial rates (Figure 5c) (Lund et al., 2019; Middleton et al., 2020). Surplus $^{230}$Th burial where hydrothermal particle flux is high must be supplied by lateral diffusion from further off-axis, which creates $^{230}$Th deficits where Fe fluxes are relatively low. Reduced $^{230}$Th burial is clearly observed on the JdFR, at distances of approximately 5 to 12 km off-axis. On the southern EPR, the magnitude and spatial footprint of ridge flank $^{230}$Th deficits remains unknown, and it would require analyzing $^{230}$Th burial fluxes from an array of sites from the ridge crest and flanks. We suggest that a full quantitative model for the effects of hydrothermal scavenging on $^{230}$Th burial be developed that combines $^{230}$Th and $^3$He analyses from a range of hydrothermal environments with variable Fe flux, plume incidence, spreading rate, and ridge geometry. In the interim, we suggest that both $^{230}$Th and $^3$He normalization be used to assess the potential influence of hydrothermal scavenging on bulk sedimentation rates in ridge proximal locations.

6.4. Grain Size Effects and Focusing: Does Particle Size Bias $^{230}$Th-Normalized Sedimentary Fluxes?

Because small particles (<10 μm) have a large specific surface area relative to their mass, they tend to bear higher $^{230}$Th concentrations relative to other particle classes in sediment (Kretscher et al., 2010; Loveley
et al., 2017; McGee et al., 2010). For example, in sediments from the Southern Ocean and southeast Atlantic, it was found that $^{230}$Th concentrations were 1.6–2.2 times higher in the $<2 \mu m$ fraction than in the bulk sediment (Kretschmer et al., 2010), while in the eastern equatorial Pacific, $^{230}$Th concentrations in the $<4 \mu m$ fraction ranged from 0.7 to 2.1 times the $^{230}$Th concentrations in the bulk sediment (Loveley et al., 2017). Any process that preferentially affects fine grain size classes thus has the potential to decouple bulk sedimentary $^{230}$Th concentrations from the overlying integrated water column inventory.

One such process is sediment redistribution along the seafloor, during which near-bottom flow speeds $>10$–15 cm/s will preferentially remove fine grains ($<16 \mu m$) from areas of winnowing and redeposit them further downstream in areas of focusing (e.g., Law et al., 2008; McCave et al., 1995, 2017). The consequences of such fine-fraction redistribution on $^{230}$Th-normalized fluxes have been modeled by Kretschmer et al. (2010), with the assumptions that (i) the original vertical flux is coarser than the lateral sediment flux; (ii) lateral sediment flux is controlled by preferential transport of fines; and (iii) both vertical and lateral fluxes contain the same $^{230}$Th activity in the fine fraction. This model demonstrated that, in the Southern Ocean, the particle size effect may lead to an underestimation of vertical fluxes in areas of focusing and an overestimation of vertical fluxes in areas of winnowing. The estimated bias scales with both the degree of sediment focusing and the mean grain size of the focused sediment, ranging from as low as a 6% underestimation of true mass fluxes to as much as 80% underestimation (Kretschmer et al., 2010). Similarly, in the eastern equatorial
This resilience can be demonstrated by comparing mass fluxes of bulk sediment, fine material, and coarse material from multiple cores that experience different degrees of sediment remobilization (Figure 6), including both focusing and winnowing. A common proxy for lithogenic material, $^{232}$Th is primarily carried in the fine fraction, so that any grain size fractionation of $^{230}$Th will be compensated by the inverse effect on $^{232}$Th activities. In contrast, coarse material (>63 μm) will be particularly insensitive to sediment remobilization, as focusing and winnowing generally do not act on these larger grain sizes. The JdFR, in the northeast Pacific ($-45°N$, $-135°E$, 2,655–2,794 m), contains six sites within 50 km of one another and suffers from a range of sediment focusing caused by the rough bathymetry of the mid-ocean ridge (Costa et al., 2016a; Costa & McManus, 2017). The Line Islands, in the central equatorial Pacific ($-0$–$7°N$, $-160°E$, 2,798–3,542 m), contain nine sites along a rough carbonate ridge (Lyle et al., 2016), and although the core transect spans over 1,000 km, their extremely low dust content leads to only minor spatial trends in dust flux on glacial-interglacial time scales (Costa et al., 2016b; Jacobel et al., 2017a). Focusing factors were calculated within Marine Isotope Stages (MIS), e.g., Lisiecki & Raymo, 2005. Flux anomalies were calculated by first averaging (mean) fluxes over each MIS at each site and then calculating the regional average flux of all sites ($n = 6$ for JdFR, $n = 9$ for Line Islands) for each MIS, subtracting this regional average from the flux at each site and then dividing the site difference by the regional average to obtain the percent anomaly. The same procedure was applied for $^{232}$Th fluxes and coarse fluxes. Samples with uncertainties (1 s.e.) greater than ±100% have been excluded.

Theoretically, focused sites ($Ψ > 1$) would contain excess $^{230}$Th, creating a negative flux anomaly and plotting in the lower right quadrant (Figure 6). Winnowed sites ($Ψ < 1$) would have a deficit of $^{230}$Th, creating a positive flux anomaly and plotting in the upper left quadrant. These deviations are expected to be damped in the $^{232}$Th (fine) flux (Figure 6b) and pronounced in the coarse flux (Figure 6c). Yet no systematic bias in any of the three fluxes—bulk, fine, or coarse—emerges as a function of focusing factors, in either region. Total fluxes, fine fluxes, and coarse fluxes may be overestimated or underestimated at winnowed sites. Total

Pacific, preferential focusing of fine grain size classes has been suggested to affect the $^{230}$Th-derived vertical flux at focused sites (Marcantonio et al., 2014; Lyle et al., 2005), causing underestimations of 30% or less in most cases but maximally 70% underestimation in one sample (Loveley et al., 2017).

Importantly, in both the Southern Ocean and the eastern equatorial Pacific, even the most extreme particle size effects (e.g., up to 80% underestimation) are still considerably lower than errors associated with age model-based flux estimates in these areas. In most pelagic settings, the particle size effect is likely to be less than 30% underestimation or overestimation of mass fluxes, which is within the range of other errors associated with the $^{230}$Th activity of the bulk sediment in focused cores, corroborating the finding that the grain size effect on Th-normalized flux estimates in this area is likely small (Bista et al., 2016). This insensitivity to sediment focusing may arise because most pelagic sediment is already relatively fine (<35% coarse; McGee et al., 2010), so that there may not be any conspicuous grain size discrepancy in the lateral sediment flux relative to the vertical sediment flux, in contrast to the first assumption of the sediment remobilization model of Kretschmer et al. (2010). Furthermore, in many places, sediment focusing occurs via syndepositional transport of phyto-detritus “fluff” material (Beaulieu, 2002; Nodder et al., 2007; Smith et al., 1996) that may incorporate particles of all size classes, so that size fractionation during winnowing and transport would be minimal. In a practical sense, the application of $^{230}$Th normalization in paleoceanographic contexts does not appear to be particularly sensitive to the degree of sediment focusing that occurs in pelagic settings.

Figure 7. Changes in $^{230}$Th-normalized mass fluxes in response to changes in calcium carbonate burial. Core top mass fluxes are reconstructed from depth transects recovered from five seamounts in the tropical Atlantic (Rowland et al., 2017). Data are represented as deviations from the mass flux or calcium carbonate at the shallowest core top of each seamount; that is, a $\Delta$CaCO$_3$ of $-20\%$ indicates that the carbonate concentration is 20% lower than that of the shallowest core top on that seamount. The red expectation. Points below the red curve to mass calcium carbonate dissolution (dashed red arrows) would push the data in conjunction with the calcium carbonate particles. Loss of 230Th during fluxes that are too low relative to the flux if 230Th does not dissolve.
fluxes, fine fluxes, and coarse fluxes may be overestimated or underestimated at focused sites. In other words, the degree of sediment focusing does not appear to be a reliable predictor of flux anomalies in the sediment. Previous studies have demonstrated the resiliency of fine fraction fluxes against grain size fractionation (e.g., McGee et al., 2010), in accordance with theoretical expectations, but we demonstrate here that $^{230}$Th normalization may function adequately in coarse fraction fluxes as well. Thus, even though fine sediment may be more susceptible to sediment redistribution, its preferential mobility along the seafloor does not appear to significantly bias $^{230}$Th systematics.

6.5. Diagenesis and Calcium Carbonate Dissolution

One key assumption of $^{230}$Th normalization is that the $^{230}$Th adsorbed onto particles is unaltered by sediment dissolution. If, for example, 50% of the particles reaching the seafloor subsequently dissolve, then the affiliated $^{230}$Th, which is highly particle reactive, would be predicted to readorb onto the remaining fraction of sediment. The sedimentary $^{230}$Th concentration would then increase by a factor of 2, since the same inventory of $^{230}$Th is distributed over half the sediment mass, and the preserved $^{230}$Th-normalized mass flux would halve. But what if the $^{230}$Th did not fully readorb onto adjacent particles? If instead, some or all of the newly dissolved $^{230}$Th were permanently “lost” from the sediment by advection or diffusion, then sediment dissolution would bias $^{230}$Th concentrations too low and subsequent calculations of mass fluxes too high. Where systematic patterns of sediment dissolution occur, as they do for carbonate, especially below the lysocline, $^{230}$Th-normalized mass fluxes may be particularly susceptible to biases from loss of $^{230}$Th out of the sediment. The influence of calcium carbonate (CaCO$_3$) dissolution on $^{230}$Th-normalized mass fluxes may be investigated by looking at core top depth transects, for example, in a study from seamounts across the tropical Atlantic (Rowland et al., 2017). These sites are primarily composed of CaCO$_3$ and lithogenic components, and they are proximally located such that the principle deviations in apparent preserved mass flux with depth should result from increasing carbonate dissolution with increasing depth. For each seamount, depth-dependent CaCO$_3$ concentration and $^{230}$Th-normalized mass flux anomalies can be calculated relative to the shallowest core top available (Figure 7). If $^{230}$Th is conserved in the sediment as CaCO$_3$ dissolves, then the change in mass flux and the change in CaCO$_3$ should scale in a coherent manner (Figure 7, red line). Instead, the data scatter broadly, with little to no relation to the expected behavior ($R^2 = 0.09, p = 0.22$). Some of this inconsistency may be due to loss of $^{230}$Th during CaCO$_3$ dissolution, which has resulted in mass fluxes lower than predicted (Figure 7, red arrows), but most of the data do not deviate in the manner predicted by $^{230}$Th loss. The comparison is unfortunately imperfect, as scatter may have also been introduced by unrelated processes, such as differential bioturbation, inconsistent core top ages, or downward sediment transport along steep seamount slopes. A more rigorous assessment of the effects of dissolution on $^{230}$Th loss would require additional data from similarly clustered sites along CaCO$_3$ preservation gradients and, ideally, independent mass flux constraints from $^3$He.

A different approach to investigating the effects of CaCO$_3$ dissolution on $^{230}$Th retention in the sediment was conducted in the equatorial Pacific. Without a shallow site to benchmark the initial CaCO$_3$ content, it is impossible to infer the absolute extent of CaCO$_3$ dissolution from the sedimentary CaCO$_3$ content. Instead, the degree of dissolution would need to be estimated using an independent CaCO$_3$ dissolution proxy, for example, the fragmentation of fragile foraminiferal shells, such as those of *Globorotalia menardii* (menardii fragmentation index, or MFI; Mekik et al., 2002, 2010). In the equatorial Pacific, a comparison of MFI-based CaCO$_3$ dissolution and $^{230}$Th-normalized mass flux showed that for core tops with less than ~64% dissolution, the relationship between $^{230}$Th-normalized mass fluxes and % CaCO$_3$ dissolution (Mekik & Anderson, 2018) is consistent with reasonable $^{230}$Th retention in the sediment. As a consequence, the small variability in mean glacial CaCO$_3$ (~86%) and interglacial CaCO$_3$ (~83%) in the equatorial Pacific (Anderson et al., 2008) is unlikely to cause glacial-interglacial biases in sediment $^{230}$Th retention, even if that retention is incomplete in either time period. More extreme variability in CaCO$_3$ preservation, such as during transitional events from glacial to interglacial periods of the late Pleistocene (Anderson et al., 2008), would suggest much higher rates of CaCO$_3$ dissolution, which may be more likely to have an effect on $^{230}$Th retention in sediments on millennial time scales.

In summary, $^{230}$Th concentrations are observed to increase with increasing CaCO$_3$ dissolution, which confirms that some portion of $^{230}$Th previously associated with the dissolved sediment is indeed retained on the
Therefore, the assumption that $^{230}$Th becomes readsorbed is at least partly valid. Additionally, the wealth of water column data now available from GEOTRACES (e.g., Hayes, Anderson, Fleisher, Vivancos, et al., 2015) and other work, shows no definitive indication of increased concentrations of dissolved $^{230}$Th near the seafloor that would be expected if significant loss of $^{230}$Th from the seabed occurred (Francois et al., 2007), arguing against the widespread loss of $^{230}$Th from sediments to the water column. As yet, the scarcity of appropriate data (e.g., high-resolution depth transects and $^3$He-normalized $^{230}$Th fluxes) currently precludes a robust test of the extent of $^{230}$Th loss and the conditions under which it may be problematic for the application of $^{230}$Th as a constant flux proxy, which remains an outstanding question for further investigation.

7. Outlook on a New Development: $^{230}$Th Normalization in the Water column

In addition to quantifying fluxes at the seafloor, $^{230}$Th can also be used to estimate the sinking flux of particles in the water column (e.g., Figure 4e). Sinking particles can be expected to carry all of the overlying production of $^{230}$Th due to U decay at any depth level in the water column. The flux of a particulate component or element can be estimated simply by measuring the ratio of that component to $^{230}$Th in sinking particles. This approach is equivalent to the application of $^{230}$Th normalization to sediments, except that particles collected from the water column by filters are used instead (Hirose, 2006). Thanks to advances in analytical capabilities under GEOTRACES, particulate $^{230}$Th normalization in filter samples has now been used to generate flux profiles of trace elements (Hayes et al., 2018), particulate organic carbon (Hayes et al., 2018; Pavia, Anderson, Lam, et al., 2019), and lithogenic dust (Anderson et al., 2016).
While this method shows considerable promise to provide unprecedented constraints on regeneration rates of marine particulate constituents, there remain assumptions that require further testing. First, similar to sedimentary $^{230}$Th normalization, model estimates and more regional measurements (e.g., in annually averaged sediment traps) of $^{230}$Th burial fluxes at different depths will be required to validate where $^{230}$Th flux is equal to its integrated water column production, comparable to previous studies on sediment traps (Scholten et al., 2001; Walter et al., 2000). Second, water column $^{230}$Th normalization to date has been applied to suspended ($0.8-51 \mu m$) particles rather than sinking (e.g., $>51 \mu m$) particles due to sampling constraints on filters. Provided that the aggregation and disaggregation of small and large particles are in equilibrium (Bacon et al., 1985), fluxes determined on suspended particles will still be valid. Results from a limited number of measurements using large ($>51 \mu m$) particles indicate an offset from results obtained using the smaller size class of only a few tens of percent (Anderson et al., 2016; Pavia, Anderson, Lam, et al., 2019); however, additional measurements of $^{230}$Th on larger sinking particles will ultimately be needed to shed light on the particle dynamics involved in Th removal.

Provided that the assumptions inherent to the method can be validated, water column $^{230}$Th normalization on filtered particles could ultimately function as a more cost-effective, higher-resolution method than bottom-moored sediment traps for determining annual to multiannual particulate fluxes. These fluxes would constitute an extremely powerful tool for studying upper water column biogeochemistry (e.g., the spatial pattern of organic carbon fluxes and regeneration; Pavia, Anderson, Lam, et al., 2019), and for providing near-bottom fluxes of material arriving to the seafloor in studies of early diagenesis.

8. Takeaways and Future Recommendations: The Utility of $^{230}$Th Normalization on a Global Scale

As proxies mature, the continued evaluation of their inherent assumptions is necessary to retain their relevance to the oceanographic community. In this review, we have compiled all existing $^{230}$Th data and explored existing and new caveats to the proxy's functionality. Based on this discussion, we can summarize (1) the applicability of $^{230}$Th as a constant flux proxy and (2) useful directions for future research.

The main takeaway of this work is that $^{230}$Th normalization, as it is typically applied in paleoceanographic research, performs sufficiently well to serve the purpose for which it is intended. That is, as yet, we have no strong evidence for significantly aberrant $^{230}$Th behavior that would make age model-based mass accumulation rates or other approaches preferable. We do note, however, that there are several regions or circumstances (Figure 8) under which special consideration is recommended when interpreting $^{230}$Th-normalized mass fluxes:

1. Shallow waters ($<1,000 \text{ m}$): Low $^{230}$Th inventories produced in the shallow water column may make application of $^{230}$Th normalization analytically challenging.
2. Continental margins, particularly in eastern boundary current regions of high biological productivity and downwind of major dust sources (e.g., NW Africa and the Arabian Sea): High particle fluxes can create concentration gradients that lead to the burial of more $^{230}$Th than is produced in the overlying water column ($F/P > 1$). Absolute mass fluxes in such locations may thus be underestimated, but relative changes in mass fluxes are still likely to be robust.
3. Polar oceans: Models and water column data suggest that boundary scavenging may be quite active in the Arctic and Antarctic seas, suggesting that $^{230}$Th-normalized mass fluxes in these regions may be overestimated. We note, however, that these regions, particularly the Arctic, are the least well constrained by sedimentary data and that the complex circulation of the Southern Ocean is notoriously difficult to capture in model simulations. Future work to improve model skill and to provide better data coverage of these regions will greatly enhance our understanding of $^{230}$Th systematics in the polar oceans.
4. Hydrothermal vents: Scavenging by hydrothermal Fe-Mn particles can lead to the enhanced burial of $^{230}$Th close to active vents ($F/P > 1$) and reduced $^{230}$Th burial on the ridge flanks ($F/P < 1$). The spatial domain of these effects is poorly constrained, but it may be a function of ridge-specific hydrothermal iron flux. Mass fluxes derived using $^{230}$Th normalization are likely to be underestimated near vents and overestimated on ridge flanks, but as yet these effects have only been observed.
within ~30 km of the ridge axis. Future work in a range of hydrothermal settings with different particulate fluxes and chemistry will help establish more specific guidelines for the use of $^{230}$Th as a constant flux proxy on mid-ocean ridges.

5. Benthic nepheloid layers: Resuspension of particles from the seafloor may scavenge $^{230}$Th from the water column. Although enhanced near-bottom scavenging by resuspended sediment by itself does not violate the assumptions inherent in $^{230}$Th normalization, provided that 1-D mass balance remains intact, empirical evidence from the northwestern Atlantic nepheloid layers indicates a strong bias in $^{230}$Th-normalized fluxes. The effects of benthic nepheloid layers on the assumptions of $^{230}$Th normalization are poorly constrained, and basic questions about the processes involved remain unanswered. Future work in this area may include the incorporation of benthic nepheloid layers in modeling efforts and the application of $^3$He normalization to calculate $F/P$ ratios in regions afflicted with these nepheloid layers.

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