How well do global ocean biogeochemistry models simulate dissolved iron distributions?

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Abstract

Numerical models of ocean biogeochemistry are relied upon to make projections about the impact of climate change on marine resources and test hypotheses regarding the drivers of past changes in climate and ecosystems. In large areas of the ocean, iron availability regulates the functioning of marine ecosystems and hence the ocean carbon cycle. Accordingly, our ability to quantify the drivers and impacts of fluctuations in ocean ecosystems and carbon cycling in space and time relies on first achieving an appropriate representation of the modern marine iron cycle in models. When the iron distributions from 13 global ocean biogeochemistry models are compared against the latest oceanic sections from the GEOTRACES program, we find that all models struggle to reproduce many aspects of the observed spatial patterns. Models that reflect the emerging evidence for multiple iron sources or subtleties of its internal cycling perform much better in capturing observed features than their simpler contemporaries, particularly in the ocean interior. We show that the substantial uncertainty in the input fluxes of iron results in a very wide range of residence times across models, which has implications for the response of ecosystems and global carbon cycling to perturbations. Given this large uncertainty, iron fertilization experiments based on any single current generation model should be interpreted with caution. Improvements to how such models represent iron scavenging and also biological cycling are needed to raise confidence in their projections of global biogeochemical change in the ocean.

1. Introduction

With the important role played by dissolved iron (DFe) in regulating ocean biogeochemical cycles well established [Boyd and Ellwood, 2010], most three-dimensional global biogeochemistry models now include a prognostic DFe tracer as standard. These models explicitly represent the DFe limitation of primary production that is prevalent across large areas of the ocean [C. M. Moore et al., 2013]. This has allowed quantitative projections regarding the impacts of environmental change in Fe-limited regions [Bopp et al., 2012], how DFe may regulate glacial-interglacial changes to the global carbon cycle [Tagliabue et al., 2009], and the wider role played by different nutrients as drivers of planktonic diversity [Ward et al., 2013]. However, the robustness of these results is reliant on how a given model represents the ocean DFe cycle. For example, a model that accounted for the hydrothermal sources of Fe was shown to be less sensitive to changes in aeolian iron supply than the same model without a hydrothermal input [Tagliabue et al., 2010]. Equally, there is a sixfold difference in the estimated impact of dust variations on glacial and interglacial changes in atmospheric CO₂ (5–28 ppm) [Kohfeld and Ridgwell, 2009] that is largely driven by details of the modeled DFe cycle.

In brief, the ocean iron cycle is regulated by a complex array of different processes [Boyd and Ellwood, 2010]. DFe is thought to be supplied to the ocean from atmospheric deposition [Jickells et al., 2005], continental margins [Erold et al., 2004], and hydrothermal vents [Tagliabue et al., 2010], with potential emerging roles for input from rivers [Rijkenberg et al., 2014], icebergs [Raiswell et al., 2008], and glaciers [Gerringa et al., 2012].
DFe is relatively insoluble in oxygenated seawater, and DFe levels are maintained to a large part due to complexation with organic ligands that bind Fe [Gledhill and Buck, 2012]. Unbound or free Fe can then precipitate as solid forms or be scavenged by particles [Bruland et al., 2014]. DFe is operationally defined by the filter size (usually 0.2 μm), and over half of the DFe pool can be colloidal [Boye et al., 2010; Fitzsimmons and Boyle, 2014; Wu et al., 2001]. This implies that the aggregation and coagulation of colloidal Fe, termed “colloidal pumping” [Honeyman and Santschi, 1989], may also be an important loss of DFe. As a divalent metal, Fe also undergoes rapid redox transformations between Fe(II) and Fe(III) species mediated by oxidation, reduction, and photochemical processes [Wells et al., 1995]. The biological cycling of Fe is also complex with varying cellular requirements for Fe [Raven, 1988; Raven et al., 1999] and the role of luxury uptake [Marchetti et al., 2009] driving a wide range in phytoplankton Fe quotas [Sunda and Huntsman, 1997; Twinning and Baines, 2013]. Equally, the recycling of DFe by bacteria, viruses, and zooplankton is emerging as a key component in governing the Fe supply to phytoplankton [Barbeau et al., 1996; Boyd et al., 2012; Hutchins and Bruland, 1994; Strzepek et al., 2005]. Lastly, process studies and basin-scale data syntheses have highlighted important specificities to the remineralization length scale and vertical profile of DFe, relative to other nutrients [Frew et al., 2006; Tagliabue et al., 2014c; Twinning et al., 2014].

The earliest global iron models were informed by the first efforts to synthesize the emerging data sets on DFe in the late 1990s [Johnson et al., 1997]. These models only considered a dust source, applied constant phytoplankton Fe demands, and inferred that the seemingly constant deep ocean DFe concentrations indicated a threshold stabilization of DFe by organic ligands [Archer and Johnson, 2000; Lefèvre and Watson, 1999]. As available DFe data sets expanded, it became clear that deep ocean concentrations were more regionally and temporally varied than accounted for by these models and that explicitly computing uncomplexed DFe led to a better model-data agreement [Parekh et al., 2004]. At the same time, assumptions regarding fixed iron solubility in dust and constant C:Fe ratios in exported organic matter were being questioned and alternatives tested [Ridgwell, 2001; Watson et al., 2000]. Toward the end of the Joint Global Ocean Flux Study era more complicated treatments of the demand for DFe from different phytoplankton groups also emerged and, when coupled to realistic models of ocean circulation, provided the first estimates of the areal extent of DFe limitation [Aumont et al., 2003; Moore et al., 2002]. In more recent years, and particularly with the advent of the GEOTRACES program (www.geotraces.org), observations of DFe have expanded rapidly [Mawji et al., 2015; Tagliabue et al., 2012]. This has driven the representation of DFe sources associated with margin sediments [Moore and Braucher, 2008] and hydrothermal vents [Tagliabue et al., 2010] in models. At the same time efforts to account for redox speciation [Tagliabue and Völker, 2011] and variability in Fe binding ligands [Misumi et al., 2013; Völker and Tagliabue, 2015] in global models have also been undertaken.

Until now there has been no comprehensive effort to evaluate how different global models represent DFe, apart from the one off model-data comparisons typical of individual publications [Moore and Braucher, 2008; Tagliabue et al., 2008]. Our maturing vision of the oceanic distribution of DFe and our deeper understanding of how it interacts with broader biogeochemical cycles now allows a more widespread intercomparison of global iron models. In conducting the first “iron model intercomparison project” (FeMIP), we aim to intercompare as broad a suite as possible of global ocean biogeochemistry models with a focus on the reproduction of features present in the full depth ocean sections emerging from the GEOTRACES program. In doing so, we highlight the challenges present for global ocean biogeochemistry models in simulating the distribution of DFe, which emerges as unique to that of other nutrients.

### 2. Methodology

#### 2.1. Intercomparison Process

The goal of this study was to include as many global iron models as possible in order to ensure a “state of the art” view on their representation of Fe cycling. In that regard, our 13 models (Table 1) range from those used in the recent Intergovernmental Panel on Climate Change report for coupled climate-carbon studies to those focused on global patterns of Fe cycling and effects on ocean biogeochemical cycles and phytoplankton diversity and to those concerned with geological timescales. This inclusive design thus did not impose a rigid set of guidelines regarding the model forcings, as done for the ocean carbon cycle model intercomparison and climate model intercomparison projects. While imposing identical ocean circulation or external forcing scenarios would have permitted a more direct cross comparison of the different iron models, the extra
Table 1. A Summary of the FeMIP Modelsa

<table>
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<tr>
<th>Model</th>
<th>Spin-Up</th>
<th>Dust</th>
<th>Sediment</th>
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constraints would have drastically reduced the number of Fe models able to participate and hinder our aim to account for the full diversity of Fe models. Groups submitted their best representation of the dissolved iron distribution in netCDF format at monthly frequency for a canonical year on their standard model grid, alongside additional requested information (temperature, salinity, nitrate, phosphate, and silicic acid concentrations, where available). We compiled model data from 13 model configurations: BEC [J. K. Moore et al., 2013], BFM [Vichi et al., 2007], BLINGv0 [Galbraith et al., 2010], COBALT [Stock et al., 2014], GENIE [Fe scheme as summarized by Matsumoto et al. [2013], MEDUSA1 [Yool et al., 2011], MEDUSA2 [Yool et al., 2013], MITecco [Dutkiewicz et al., 2015], MITigsm [Dutkiewicz et al., 2014], PISCES1 [Aumont et al., 2015], PISCES2 [Resing et al., 2015; Völker and Tagliabue, 2015], REcoM [Hauck et al., 2013], and TOPAZ [Dunne et al., 2013], all implemented at the global scale. All models were then regridded onto a 1°×1° horizontal grid with 33 vertical levels (bounded by 0, 10, 20, 30, 40, 50, 75, 100, 125, 150, 200, 250, 300, 400, 500, 600, 700, 800, 900, 1000, 1100, 1200, 1300, 1400, 1500, 1750, 2000, 2500, 3000, 3500, 4000, 4500, 5000, and 5500 m) as a common FeMIP grid.

2.2. Observational Data Sets

Observations of dissolved iron are taken from two sources. First, we use an updated version of a global DFe database [Tagliabue et al., 2012] with approximately 20,000 individual observations. This database was gridded at monthly resolution on the FeMIP grid to compare models and observations grid cell by grid cell and month by month, with no volume weighting. Second, we extracted DFe data from recent GEOTRACES sections from the 2014 intermediate data product [Mawji et al., 2015]. For comparison purposes (section 3.2) the modeled DFe from the longitude, latitude, and month of each sampling station was then extracted, and the observed data were regridded on the same 33 vertical levels as the models (averaging where more than one observation was present in a particular depth bin). We use data sets collected on the GA-02 West Atlantic cruise [Rijkenberg et al., 2014], the GA-03 North Atlantic zonal transect [Hatta et al., 2014], the CoFeMUG south Atlantic zonal cruise [Saito et al., 2013], the GIPY-6 Atlantic sector of the Southern Ocean cruise [Chever et al., 2010; Klunder et al., 2011], and the recently completed GP-16 Equatorial Pacific zonal section [Resing et al., 2015] that is not yet in the GEOTRACES data product. We note that all IDP2014 GEOTRACES data [Mawji et al., 2015] are also included in the global data set.

2.3. Brief Introduction of the Different Iron Models

The goal here is not to exhaustively describe the FeMIP models for which we refer to the original publications. Rather, we seek to summarize how the models treat key components of the Fe cycle and to highlight important differences (Table 1). In our summary we focused on how each model treated the sources of Fe, the chemistry of Fe (including the representation of Fe-binding ligands, how free Fe is computed, and whether...
scavenging is a first-order rate or a second-order function of particle concentrations), biological cycling of Fe (if Fe/C ratios were variable and if zooplankton excretion of Fe depends on the Fe content of prey), and particle Fe dynamics (how many particle pools were simulated and whether the Fe regeneration efficiency was unique or coupled to organic matter).

All models considered a dust source of Fe. Only BFM, GENIE, and MEDUSA1 did not consider sedimentary Fe supply; only BEC, BFM, PISCES1, and PISCES2 include river input of Fe; and BEC, PISCES1, and PISCES2 are the only models that represent hydrothermal Fe input. All models except BEC compute the free Fe concentration that can be scavenged based on Parekh et al. [2004] and all except BFM, COBALT, MEDUSA1, and MEDUSA2 have a second-order scavenging rate, i.e., a dependency on particle concentrations. Only PISCES1 and PISCES2 include a representation of colloidal losses of DFe, based on aggregation of dissolved organic matter [Aumont et al., 2015]. It is notable that despite a maturing understanding of the variations in the concentrations of Fe binding ligands [Gledhill and Buck, 2012] most FeMIP models still assume a constant ligand concentration (as per the earliest Fe models) that is 1 nM for all models except BFM and PISCES1 which use 0.6 nM. Two exceptions in this regard are PISCES2 and TOPAZ. TOPAZ applies an empirical relationship to dissolved organic carbon (DOC) to derive ligand concentrations (5 × 10^{-3} mol ligand per mol DOC). PISCES2 is the only FeMIP model to represent a dynamic ligand pool with explicit sources and sinks [Völker and Tagliabue, 2015] and a variable computation of the colloidal Fe fraction [Liu and Millero, 1999], modified to account for hydrothermal ligand supply [Resing et al., 2015]. BLING switches off Fe scavenging when oxygen drops below 1 mmol m^{-3} [Galbraith et al., 2010], and both BLING and COBALT reduce the stability of Fe-ligand complexes in the presence of light [Galbraith et al., 2010; Stock et al., 2014]. Both the MITecco and MITgsm models cap DFe to a maximum value of 1.3 nM with any excess Fe being numerically deleted. Due to the noted flexibility in planktonic demands for Fe [Sunda and Huntsman, 1997; Twining and Baines, 2013], almost all FeMIP models have variable Fe/C ratios, with only MEDUSA1, MEDUSA2, MITecco, and MITgsm retaining fixed Fe/C ratios. Recycling by zooplankton is variable in some FeMIP models and thus dependent on an assumed zooplankton Fe quota, except for BEC, BLING, MEDUSA1, MEDUSA2, and REcoM where there is a fixed rate of recycling. Lastly, all models include one particulate Fe pool, except PISCES1 and PISCES2, which consider two, and BEC, which represents sinking implicitly (accounting for ballasting). Only COBALT invokes reduced regeneration efficiency relative to organic material that elongates the regeneration depth scale beyond that for sinking organic material [Stock et al., 2014].

Finally, it is notable that several models were only run for a few decades or centuries (BEC, BFM, COBALT, MEDUSA1, MEDUSA2, MITecco, and MITgsm), a time comparable to the respective residence time of Fe in the model in some cases, making them potentially more sensitive to their initial conditions. This issue is discussed in more detail in section 3.1.1.

3. Results

3.1. Intermodel Differences in Dissolved Iron Distributions and Cycling

3.1.1. Iron Fluxes and Residence Times

Beginning with an integrated view, there is substantial variability in the modeled Fe residence times across the FeMIP models with two broad groupings of a few years and a few hundred years (Table 2). Across the 13 models, all include dust sources, 10 include sediment sources, but only three include hydrothermal and riverine Fe sources, respectively (Table 2). Even for a given source, there is substantial intermodel difference in its strength. For example, dust fluxes of dissolved iron range from ~1 to >30 Gmol Fe yr^{-1} between models (Table 2, accounting for any intermodel variations in solubility and mineral fraction). These intermodel differences across all input fluxes result in a wide range of total iron inputs to the ocean (66.9 ± 67.1 Gmol Fe yr^{-1}, Table 1). In contrast, we find a surprising degree of agreement in the mean ocean iron concentration (0.58 ± 0.14 nM, Table 2) from the models, with slightly greater intermodel differences in the total integrated inventory of Fe reflecting different model grid sizes (e.g., some models do not include the Arctic Ocean or the Mediterranean). Ultimately, this results in a wide range of residence times of dissolved iron in the models (~5 to >500 years, Table 2) that reflects different assumptions regarding the strength of the sources of DFe to the ocean compensated by variable scavenging rates in order to reproduce the observed DFe concentration.

The derivation of the residence time for Fe from each model allows us to evaluate the impact of the shorter runs performed for some models. Taken at face value, even the relatively short runs performed by almost all
Table 2. A Summary of the Magnitude of the Fe Sources, the Total and Average Fe Inventories, and the Residence Time of Fe Across the FeMIP Models

<table>
<thead>
<tr>
<th>Model</th>
<th>Dust</th>
<th>Sediment</th>
<th>Hydrothermal</th>
<th>Rivers</th>
<th>Total</th>
<th>Fe Inventory (x10^11 mol)</th>
<th>Average Fe (nmoles L^-1)</th>
<th>Residence Time (years)</th>
</tr>
</thead>
<tbody>
<tr>
<td>BEC</td>
<td>21.9</td>
<td>84.6</td>
<td>17.7</td>
<td>0.34</td>
<td>124.5</td>
<td>10.1</td>
<td>0.74</td>
<td>8.1</td>
</tr>
<tr>
<td>BFM</td>
<td>1.4</td>
<td>0</td>
<td>0</td>
<td>0.06</td>
<td>1.4</td>
<td>8.8</td>
<td>0.65</td>
<td>626.3</td>
</tr>
<tr>
<td>BLING</td>
<td>3.3</td>
<td>9.1</td>
<td>0</td>
<td>0</td>
<td>12.4</td>
<td>5.3</td>
<td>0.37</td>
<td>42.4</td>
</tr>
<tr>
<td>COBALT</td>
<td>32.5</td>
<td>155</td>
<td>0</td>
<td>0</td>
<td>182.5</td>
<td>6.8</td>
<td>0.50</td>
<td>3.7</td>
</tr>
<tr>
<td>GENIE</td>
<td>1.8</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>1.8</td>
<td>10.1</td>
<td>0.48</td>
<td>560.0</td>
</tr>
<tr>
<td>MEDUSA1</td>
<td>2.7</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>2.7</td>
<td>6.3</td>
<td>0.46</td>
<td>232.0</td>
</tr>
<tr>
<td>MEDUSA2</td>
<td>3.4</td>
<td>2.9</td>
<td>0</td>
<td>0</td>
<td>6.8</td>
<td>4.8</td>
<td>0.35</td>
<td>69.9</td>
</tr>
<tr>
<td>MITigsm</td>
<td>3.5</td>
<td>104</td>
<td>0</td>
<td>0</td>
<td>107.5</td>
<td>8.8</td>
<td>0.65</td>
<td>8.2</td>
</tr>
<tr>
<td>MITecco</td>
<td>1.4</td>
<td>194</td>
<td>0</td>
<td>0</td>
<td>195.4</td>
<td>9.0</td>
<td>0.66</td>
<td>4.6</td>
</tr>
<tr>
<td>PISCES1</td>
<td>32.7</td>
<td>26.6</td>
<td>11.3</td>
<td>2.5</td>
<td>71.0</td>
<td>8.1</td>
<td>0.59</td>
<td>11.5</td>
</tr>
<tr>
<td>PISCES2</td>
<td>32.7</td>
<td>26.6</td>
<td>11.3</td>
<td>2.5</td>
<td>71.0</td>
<td>11.2</td>
<td>0.81</td>
<td>15.7</td>
</tr>
<tr>
<td>ReCoM</td>
<td>3.7</td>
<td>0.6</td>
<td>0</td>
<td>0</td>
<td>4.3</td>
<td>12.5</td>
<td>0.73</td>
<td>291.6</td>
</tr>
<tr>
<td>TOPAZ</td>
<td>13.8</td>
<td>74.8</td>
<td>0</td>
<td>0</td>
<td>88.6</td>
<td>6.8</td>
<td>0.50</td>
<td>7.6</td>
</tr>
</tbody>
</table>

Mean 66.9 8.3 0.58 144.7 175.8

the models (except BFM, MEDUSA1, and perhaps also MEDUSA2) are more than twice the residence time for Fe in that particular model. Nevertheless, it should be noted that many of these residence times for the global ocean are likely skewed toward lower values due to strong local sources that have a muted wider influence. For example, much of the interior Fe distribution in the PISCES1 model has been shown to be linked to a subducted preformed component [Tagliabue et al., 2014b], suggesting that the deep ocean equilibration timescale in this model, at least, must be much longer than the 11 years of its average residence time. This is likely to be the case for models that employ a formulation for the rate of DFe scavenging that depends on particulate fluxes, as biogenic fluxes in the ocean interior are considerably slower than near the surface where sedimentary and dust sources are dominant. Feedbacks will also exist between DFe inventory and biological fluxes, meaning that a ~1000 year timescale component to the overall equilibrium adjustment will exist that involves the redistribution of major nutrients globally. As such, this raises questions regarding the distributions of Fe in the ocean interior for models that are only run for a few decades, even if that is longer than the average residence time.

3.1.2. Statistical Assessment of FeMIP Models

In order to provide a general picture of variability amongst the models, we examine correlations between observed and simulated DFe at the same locations (Table 3). When viewed globally throughout the entire water column, correlations between observations and simulations can be as high as 0.51, while some are even

Table 3. Correlation Coefficient (R) and in Parentheses the Mean Bias (nM) Between the Different FeMIP Models and the Expanded Database of Tagliabue et al. [2012] Across Different Depth Bins

<table>
<thead>
<tr>
<th>Model</th>
<th>ALL</th>
<th>0–100</th>
<th>100–500</th>
<th>500–1000</th>
<th>2000–5000</th>
</tr>
</thead>
<tbody>
<tr>
<td>BEC</td>
<td>0.51 (–0.02)</td>
<td>0.48 (0.23)</td>
<td>0.52 (–0.05)</td>
<td>0.47 (–0.15)</td>
<td>0.31 (0.01)</td>
</tr>
<tr>
<td>BFM</td>
<td>0.39 (–0.48)</td>
<td>0.34 (–0.29)</td>
<td>0.36 (–0.47)</td>
<td>0.33 (–0.52)</td>
<td>–0.03 (–0.48)</td>
</tr>
<tr>
<td>BLING</td>
<td>0.37 (–0.33)</td>
<td>0.37 (–0.13)</td>
<td>0.49 (–0.17)</td>
<td>0.46 (–0.26)</td>
<td>0.01 (–0.44)</td>
</tr>
<tr>
<td>COBALT</td>
<td>0.45 (–0.25)</td>
<td>0.38 (–0.19)</td>
<td>0.48 (–0.25)</td>
<td>0.51 (–0.19)</td>
<td>–0.11 (–0.25)</td>
</tr>
<tr>
<td>GENIE</td>
<td>0.25 (–0.28)</td>
<td>0.43 (–0.02)</td>
<td>0.46 (–0.11)</td>
<td>0.43 (–0.20)</td>
<td>–0.14 (–0.40)</td>
</tr>
<tr>
<td>MEDUSA1</td>
<td>–0.01 (–0.24)</td>
<td>0.37 (0.23)</td>
<td>0.38 (0.04)</td>
<td>0.07 (–0.24)</td>
<td>0.07 (–0.37)</td>
</tr>
<tr>
<td>MEDUSA2</td>
<td>–0.14 (–0.32)</td>
<td>0.35 (0.29)</td>
<td>0.37 (0.07)</td>
<td>–0.06 (–0.30)</td>
<td>–0.10 (–0.51)</td>
</tr>
<tr>
<td>MITigsm</td>
<td>0.39 (–0.12)</td>
<td>0.34 (–0.10)</td>
<td>0.36 (–0.09)</td>
<td>0.33 (–0.14)</td>
<td>–0.03 (–0.11)</td>
</tr>
<tr>
<td>MITecco</td>
<td>0.37 (–0.14)</td>
<td>0.04 (–0.22)</td>
<td>0.42 (–0.24)</td>
<td>0.29 (–0.24)</td>
<td>–0.13 (–0.04)</td>
</tr>
<tr>
<td>PISCES1</td>
<td>0.47 (–0.23)</td>
<td>0.36 (–0.06)</td>
<td>0.47 (–0.17)</td>
<td>0.47 (–0.03)</td>
<td>0.21 (–0.27)</td>
</tr>
<tr>
<td>PISCES2</td>
<td>0.51 (–0.04)</td>
<td>0.37 (0.03)</td>
<td>0.52 (0.01)</td>
<td>0.43 (–0.03)</td>
<td>0.35 (–0.05)</td>
</tr>
<tr>
<td>ReCoM</td>
<td>0.39 (–0.05)</td>
<td>0.33 (0.25)</td>
<td>0.40 (0.01)</td>
<td>0.44 (–0.01)</td>
<td>–0.04 (–0.12)</td>
</tr>
<tr>
<td>TOPAZ</td>
<td>0.10 (–0.13)</td>
<td>0.42 (0.67)</td>
<td>0.27 (0.26)</td>
<td>0.33 (–0.12)</td>
<td>0.01 (–0.34)</td>
</tr>
<tr>
<td>Data</td>
<td>0.64</td>
<td>0.52</td>
<td>0.63</td>
<td>0.76</td>
<td>0.90</td>
</tr>
</tbody>
</table>

Average dissolved iron data (nM) for the different depth strata are presented in the final row. The iron data are gridded on the FeMIP grid as described in the text.
anticorrelated. The mean biases against observations are between $-0.02$ and $-0.48$ nM. In the 0–100 m depth stratum, where Fe is likely to be playing a role in regulating phytoplankton growth rates, all but one of the model correlations fall between $0.33$ and $0.48$, implying no clear link between model complexity and strength of correlation. On the other hand, the mean biases range from $-0.29$ to $0.67$ nM, which suggests less overall agreement in the absolute DFe levels. The 100–500 m depth slice has the overall highest correlations, and all but three models reach their highest correlations in this depth range. In the abyssal layers only the three models that consider hydrothermal iron input (BEC, PISCES1, and PISCES2) show a reasonable correlation with observations ($R = 0.20$ to $0.35$; other models are $<0.15$), highlighting the importance of this source in the deep ocean. However, the inclusion of hydrothermal iron input does not obviously lead to a
significant improvement in the surface ocean. Similarly, including (or not) sedimentary Fe input does not seem closely linked to reproducing observations in the surface or intermediate layers. For example, the two versions of MEDUSA with and without sedimentary iron input do not show much difference in their correlation coefficients. It is also important to note that we lack substantial coastal DFe data sets where sediments and/or river supply result in high DFe levels in a number of models (see section 3.1.3). Section 3.2 will more closely examine the different models using recent large-scale GEOTRACES sections as case studies in different ocean regions.

3.1.3. Intermodel Differences in Dissolved Iron

To examine the intermodel differences in dissolved iron in more detail, we compare the model mean DFe over the 0–100 m, 100–500 m, 500–1000 m, and 2000–5000 m depth slices, repeating the analysis for the boreal (30–90 N), tropical (30 N–30 S), and austral latitudes (90 S–30 S). This enables us to group the models into “high,” “moderate,” and “low” in terms of their DFe distribution, relative to the full model suite (Figure 1). Comparing Figure 1 with the statistical summary (Table 3) suggests that the intermodel trend in the average DFe concentration for the different depth slices does not always reflect good statistical agreement with the observations. However, it should be noted that while the intermodel trends in average DFe reflect full spatial and temporal averages, the statistics determined from observations only concern locations with available DFe observations (which is not spatially and temporally complete).

Beginning with the surface ocean (0–100 m) that is heavily influenced by surface sources and biological uptake, MEDUSA1, MEDUSA2, and TOPAZ are consistently relatively high in iron for all three latitudinal zones, including the Fe-limited southern latitudes. BEC is also relatively rich in Fe but only in the northern and tropical latitudes. The lowest DFe concentrations in all three geographic zones are simulated by the BLING, COBALT, and MITigsm models, with the remaining models intermediate throughout.

The relative tendencies between the different FeMIP models are generally conserved in the 100–500 m and 500–1000 m depth slices that are more heavily influenced by remineralization processes. Notable departures from this general trend are PISCES2 displaying relatively higher DFe levels in both depth bins. While both BFM and REcoM become more DFe rich in the 500–1000 m depth bin, TOPAZ stands out less as a high-DFe model.

In terms of hemispheric contrasts, BEC becomes lower in DFe in the southern region; otherwise, the intermodel trends are preserved.

In the deepest depth bin deep ocean sources such as hydrothermal vents as well as sediments are important. Unsurprisingly, the models that include hydrothermal vent DFe sources (BEC, PISCES1, and PISCES2) show high DFe levels. In contrast, the high DFe levels for BFM, MITecco, MITigsm, and REcoM cannot be ascribed to hydrothermal DFe input and may be related to initial conditions (e.g., for BFM) or deep ocean transport of high DFe levels. However, it is notable that BFM, MITecco, MITigsm, and REcoM do not perform well statistically in this depth range (Table 3). The BLING and MEDUSA1 models simulate the lowest concentrations in this depth bin. For a large number of models (BLING, GENIE, MEDUSA1, MEDUSA2, and TOPAZ), DFe concentrations decline in the 2000–5000 m bin, relative to the 500–1000 m bin.

3.1.4. Surface DFe Distributions in the Models

Due to its role as a limiting nutrient, we explore the simulated annual mean surface DFe concentrations from the FeMIP models in more detail (Figure 2, upper 50 m average). Here we see that as suggested by the range in the model biases (Table 3), there is a substantial degree of intermodel discord in the surface Fe distributions. Most models agree that the highest DFe concentrations are found underneath the Saharan dust plume in the tropical Atlantic, but others also emphasize dust supply into the Arabian Sea and enhanced DFe along the continental margins. A large number of the models suggest that the lowest DFe concentrations are found across the Pacific Ocean. Exceptions are GENIE and MEDUSA1, which have much higher DFe concentrations therein, and BEC, MEDUSA2, and TOPAZ, which restrict low DFe to the south Pacific only. The sub-Arctic Pacific is much more DFe depleted in BFM, MITecco, and MITigsm relative to the other FeMIP models. When the seasonality in DFe (presented as the maximum minus minimum DFe concentration over the year, Figure 3) is compared, strong intermodel differences also emerge. For example, some models show remarkably little seasonality (BFM, GENIE, MEDUSA1, MEDUSA2, and MITigsm), whereas others have large seasonal cycles over wide areas (>0.5 nM, BEC, MITecco, PISCES1, PISCES2, and TOPAZ). This illustrates where high annual mean concentrations in these regions are masking strong seasonal minima. For this reason it is not straightforward to compare the models against observed Fe that might have been collected during different seasons. At this stage, incomplete sampling over the seasonal
cycle is prevalent for virtually all locations with DFe measurements [Tagliabue et al., 2012], which precludes the mapping of DFe seasonality from observations. Table 3 is therefore more suited for a statistical assessment of the surface DFe for a given model against all available observations (where seasonal variations are accounted for by comparing model and data DFe at identical longitudes, latitudes, depths, and months).

Figure 2. Annual mean DFe concentrations (nM) averaged over the upper 50 m from the FeMIP models. Data averaged over the period January to June and July to December are taken from the expanded Tagliabue et al. [2012] data set and have been averaged over 5° bins in latitude and longitude to improve visibility.
3.2. Comparison to Recent GEOTRACES Ocean Sections

To more closely examine how the different DFe models represent the observed distribution of DFe, we focus on a range of recent GEOTRACES sections. As described above (section 2.2), each model is extracted at the exact location of the sampling locations, with the observations regridded onto the same vertical grid. We refer the readers to the cited papers below for a more complete discussion of each observational section and additional interpretation. In this assessment we emphasize the key features observed on each section and how different models are able to reproduce them. Because of this goal and because a given model may do a good job of reproducing one feature, but not another, we did not perform statistical assessments of the individual models for each section.

Figure 3. Annual maximum minus annual minimum DFe concentrations (nM) averaged over the upper 50 m from the FeMIP models.

3.2. Comparison to Recent GEOTRACES Ocean Sections

To more closely examine how the different DFe models represent the observed distribution of DFe, we focus on a range of recent GEOTRACES sections. As described above (section 2.2), each model is extracted at the exact location of the sampling locations, with the observations regridded onto the same vertical grid. We refer the readers to the cited papers below for a more complete discussion of each observational section and additional interpretation. In this assessment we emphasize the key features observed on each section and how different models are able to reproduce them. Because of this goal and because a given model may do a good job of reproducing one feature, but not another, we did not perform statistical assessments of the individual models for each section.
Figure 4. DFe concentrations (nM) from the GA-02 [Rijkenberg et al., 2014] cruise and extracted from the FeMIP models.
3.2.1. West Atlantic
The GA-02 West Atlantic meridional section provides unprecedented coverage of DFe concentrations along the Atlantic Ocean, as well as insights into different mechanisms that control the cycling, regeneration, and supply of DFe [Rijkenberg et al., 2014]. The key features of this section are (i) low surface DFe in both the northern and southern end-member surface waters, (ii) a surface DFe enrichment around 20°N in the tropics associated with a subsurface DFe minima, (iii) a strong DFe regeneration maxima at 5°-10°N centered around 500–1000 m, (iv) a hydrothermal signal at around 5°S and between 2000 and 3000 m depth, and (v) a hot spot of DFe that is present over much of the water column associated with the confluence of the Brazil and Falklands current at around 35–40°S.

The model representation of key features is as follows (Figure 4):

1. Almost all models capture low DFe in the southern end-member surface waters, except MEDUSA1 and MEDUSA2 and perhaps also REcoM and TOPAZ. However, it is only BFM and COBALT, and to a lesser degree BEC, BLING, MITigsm, PISCES1, and PISCES2, that reproduce the observed low DFe concentrations associated with the northern end-member surface waters.
2. A surface DFe enrichment (presumably from dust) around 20°N is clearly present in BEC, MEDUSA1, MEDUSA2, MITecco, MITigsm, PISCES1, PISCES2, REcoM, and TOPAZ but is less apparent in other models (BFM, BLING, COBALT, and GENIE). Nevertheless, in MEDUSA2, REcoM, and TOPAZ the influence of surface dust deposition appears to be much greater than is observed. Only PISCES1, PISCES2, and COBALT show the observed subsurface minima in DFe below the dust signal.
3. With respect to the strong DFe regeneration maxima at 5°-10°N centered around 500–1000 m, COBALT displays a regeneration maximum at around the right depth level, while in BFM high concentrations appear to be smeared from surface to the sea floor. In all other models the regeneration signal in DFe is generally too small or absent and where it is present (e.g., BFM, BLING, and GENIE) it is generally too shallow in the water column.
4. Concerning the hydrothermal signal at around 5°S and between 2000 and 3000 m depth, of the three models that include hydrothermal DFe input, only PISCES2, with a greater longevity of hydrothermal Fe [Resing et al., 2015], shows a hint of DFe enrichment in the right location. MEDUSA2 underestimates DFe in the ocean interior along the entire Atlantic section.
5. No models capture the elevated DFe over almost the entire water column around 35–40°S. In the observations, this is ascribed to the offshore export of Brazilian shelf waters or DFe input from the dissolution of particulate Fe associated with the Rio de la Plata River [Rijkenberg et al., 2014].

3.2.2. Subtropical North Atlantic
The GA-03 North Atlantic zonal section crossed the subtropical North Atlantic between Cape Verde and Woods Hole (U.S.) via Bermuda. Key signals in the data set [Hatta et al., 2014] are (i) strong enhancements in DFe associated with DFe regeneration and also coastal input along the eastern and western margins, (ii) a surface enrichment along with a subsurface minimum in DFe, and (iii) a strong hydrothermal anomaly over the mid-Atlantic ridge.

The model representation of key features is as follows (Figure 5):

1. Enhanced DFe in the subsurface along the margins is represented to different degrees by the FeMIP models. BLING, COBALT, MITecco, and PISCES1 have hints of subsurface maxima in DFe along the eastern margin. It is encouraging that the addition of ligand production during remineralization in PISCES2 clearly improves the intensity of the remineralized DFe signal. However, none of these models have a broad homogenous signal (down to >2000 m) of elevated DFe that is observed on the eastern margin, except perhaps BEC, which has a strong subsurface maximum that spreads over all depth levels.
2. The subsurface minima in DFe underlying a surface (presumably dust) enrichment is captured clearly by COBALT, PISCES1, and PISCES2 and slightly less clearly by BEC, BFM, and BLING.
3. A hydrothermal anomaly is present in PISCES1 but closer in magnitude to the observations in PISCES2, while BEC also displays a strong hydrothermal signal. COBALT displays a sediment signal at depth that is not reproduced by the observations. It also notable that many of the models present an “inverted” DFe profile, with decreasing DFe concentrations toward the ocean interior (GENIE, MEDUSA1, MEDUSA2, REcoM, and TOPAZ), which could be indicative of too great a residence time for DFe at the ocean surface. Also, BLING, COBALT, MEDUSA2, and TOPAZ seem to be systematically too low in terms of their interior ocean DFe levels across this section.
Figure 5. DFe concentrations (nM) from the GA-03 cruise [Hatta et al., 2014] and extracted from the FeMIP models.
Figure 6. DFe concentrations (nM) from the CoFeMUG cruise [Noble et al., 2012] and extracted from the FeMIP models.
3.2.3. Subtropical South Atlantic

The CoFeMUG section traversed the south Atlantic between Namibia and Brazil and had the following notable signatures [Noble et al., 2012; Saito et al., 2013]: (i) a remineralization signal and/or sediment input on the eastern margin, (ii) low overall surface concentrations, and (iii) a strong hydrothermal signal at depth.

The model representation of key features is as follows (Figure 6):

1. Interestingly, more models are able to simulate a remineralization signal on the eastern side of the basin (COBALT, MEDUSA1, MEDUSA2, MiTecco, PISCES1, PISCES2, REcoM, and TOPAZ) for this section than for the GA-03 section, although for some models this feature is too weak or spread over too many depth levels.
2. All models, except MEDUSA1, MEDUSA2, REcoM, and TOPAZ, are able to reproduce the overall low DFe conditions in the surface waters.
3. BEC and PISCES1 represent a DFe anomaly over the ridge as observed, but this is underestimated. PISCES2 represents a stronger hydrothermal signal, but it appears to spread too far off axis relative to that which is observed. Again, COBALT displays a strong sediment signal in the deep ocean that is not observed. BFM, BLING, MEDUSA2, and to some extent TOPAZ underestimate interior ocean DFe levels.

3.2.4. Southern Tropical Pacific

The GP-16 cruise ran from Ecuador to Tahiti [Resing et al., 2015] and displays the following key features: (i) DFe enrichment along the eastern margin over almost the entire water column, (ii) low surface concentrations, and (iii) a remarkable hydrothermal plume propagating westward for > 4000 km from the East Pacific Rise to at least 150°W.

The model representation of key features is as follows (Figure 7):

1. BEC, COBALT, PISCES2, and TOPAZ are the only models able to produce the broad signal of elevated DFe throughout the entire water column on the eastern margin. BLING, MEDUSA1, MEDUSA2, and REcoM display an enrichment in DFe, but this remains more tightly localized than observed.
2. All models capture the low DFe levels typical of Pacific surface waters, but for some models (BFM, BLING, COBALT, GENIE, MEDUSA2, and TOPAZ), low DFe is also too prevalent in the ocean interior.
3. BEC and PISCES1 capture a local hydrothermal signal above the East Pacific Rise, but only PISCES2 goes any way toward reproducing the degree of off-axis transport. As seen previously, MITgsm and COBALT show DFe increases near the sea floor, but these are more widespread than seen in the observations. As noted previously, BFM, BLING COBALT, MEDUSA2, and TOPAZ show too little DFe in the ocean interior (<0.3 nM), relative to the observations (>0.6 nM away from the hydrothermal plume).

3.2.5. Southern Ocean-Atlantic Sector

Both the GIPY-4 and GIPY-5 cruises ran from Cape Town (South Africa) to the Antarctic continent along the so-called “GoodHope” line during the International Polar Year [Chever et al., 2010; Klunder et al., 2011]. These cruises sampled at different resolutions north and south of the Polar Front and have been blended to form one section. Notable features in this data set include (i) low but nonzero concentrations at the surface that propagate into the subsurface, (ii) a strong remineralization signal at around 500 m near 60°S, and (iii) a strongly local hydrothermal signal over the Bouvet region ridge crest at around 54°S and more widespread elevated DFe in the abyssal ocean north of the ridge (i.e., between ~54°S and the northern end of the transect).

The model representation of key features is as follows (Figure 8):

1. Most models display low overall DFe concentrations at the surface. GENIE, MEDUSA1, MEDUSA2, and to a lesser degree REcoM and TOPAZ overestimate surface DFe concentrations. But even the models that have low surface DFe show rapid increases with depth, indicating that the ferricline is too shallow in all models.
2. No FeMIP model captures the remineralization signal seen in the subsurface just south of the Polar Front.
3. Despite including a hydrothermal source, BEC is unable to represent the local hydrothermal enrichment. While PISCES1 represents a slight hydrothermal anomaly that appears to be from an adjacent source, the longer lifetime of hydrothermal Fe in PISCES2 leads to the anomaly being too widespread in the abyssal ocean. On the other hand, both BEC and PISCES2 show elevated DFe in the abyssal ocean north of the main ridge at 54°S that compares well with the data. COBALT, MiTecco, and MITgsm again show a sediment signal in DFe at depth, while COBALT and TOPAZ show very high values near the Antarctic coast. None of these features are observed in the data set. The BFM stands out from the other models with the large underestimation of DFe in the Southern Ocean interior as already seen for the GA-02 section.
Figure 7. DFe concentrations (nM) from the GP-16 cruise [Resing et al., 2015] and extracted from the FeMIP models.
Figure 8. DFe concentrations (nM) from the GIPY-4 and 5 cruises [Chever et al., 2010; Klunder et al., 2011] and extracted from the FeMIP models.
4. Discussion

4.1. Examining Intermodel Differences in Fe Distributions Relative to Other Nutrients

In short, we find a wide range of simulated DFe distributions from current global ocean biogeochemical models that reflects an apparent lack of intermodel agreement in the processes that control the oceanic distribution of DFe. When assessed against the best DFe data sets, most models perform modestly both quantitatively in terms of magnitudes and patterns and qualitatively in representing the inferred mechanisms. This has important implications for how models are used to understand biogeochemical cycles [Galbraith et al., 2010; Moore et al., 2002; Tagliabue et al., 2014a], planktonic diversity, and resource competition [Dutkiewicz et al., 2012; Ward et al., 2013], as well as the ocean response to fluctuations in the environment in general [Bopp et al., 2013; Dutkiewicz et al., 2013; Tagliabue et al., 2009]. It is noteworthy that this intermodel disagreement appears to be solely driven by the particular way in which different models represent the Fe cycle. If we examine the models in terms of macronutrients (nitrate and phosphate), then, taking the long meridional GA-02 section as example, we see a much stronger intermodel and model-data agreement (Figures 9 and 10). Although intermodel differences due to specific physical models are visible in the Atlantic water mass structure, the mechanisms driving the N and P cycles are similar.

We further contextualize the intermodel Fe differences by examining how they represent the relative inventories of Fe and NO$_3$ in the ocean interior by plotting the Fe$^+$ tracer (Fe-NO$_3$ × $r_{Fe/NO_3}$). Defining $r_{Fe/NO_3}$ in the same way as for the GA-02 section [Rijkenberg et al., 2014] (based on the observed Fe apparent oxygen utilization relationship, which results in a Fe/N ratio of 0.47 mmol/mol) and using PO$_4$ (and a NO$_3$/PO$_4$ ratio of 16/1) for GENIE and BLING, which do not simulate NO$_3$, allows us to examine DFe concentrations relative to NO$_3$ (Figure 11). The data show relatively replete waters originating from the Northern Hemisphere linked to North Atlantic Deep Water (NADW), which becomes flanked above and below by relatively Fe-poor water from the Southern Hemisphere linked to Antarctic Intermediate Water and Antarctic Bottom Water. There is also a zone of relatively depleted Fe in the subsurface overlying the NADW signal in the Northern Hemisphere likely linked to northern subtropical mode water. In these sections we can see that NADW is relatively impoverished in DFe in MEDUSA1, MEDUSA2, and TOPAZ, despite these models generally overestimating surface DFe. This may indicate an overly short lifetime for Fe away from the surface and subsequent lack of permanence in the NADW signal. Looking at southern sourced waters, all models except BFM perform well (notwithstanding the northern sourced water biases). Obviously, this comparison should only be taken as indicative since different models are underpinned by different relationships between NO$_3$ and Fe and the actual planktonic Fe:N ratio can vary from the value chosen in the Rijkenberg et al. [2014] study [Twining and Baines, 2013]. Nevertheless, it does provide an additional means to assess the relative transport of Fe and NO$_3$ through the ocean interior.

4.2. Identifying the Key Processes at Different Depth Strata

One important intermodel difference that clearly impacts the agreement with observations and the role of Fe on biota is the strong surface enrichments evident in some models (MEDUSA1, MEDUSA2, REcoM, and TOPAZ). In the observations, any Fe enrichments due to dust deposition are far more localized and apparently short lived in space (e.g., Figures 4 and 5). For the models surface overestimation of iron implies either too large an iron source or that the residence time for Fe at the surface is too long. The latter possibility highlights the importance of how models treat the scavenging process and could also be linked to constant Fe/C ratios that do not permit “luxury uptake” of Fe at high DFe concentrations (specifically MEDUSA1, MEDUSA2, MITecco, and MITgsm). MEDUSA1, MEDUSA2, and REcoM are three of the four models with the longest residence times (decades to centuries, Table 2), relative to the other FeMIP models, and produce high surface enrichment despite having some of the lowest dust inputs (Table 2). For MEDUSA1 and MEDUSA2 the first-order fixed scavenging rate may be too low or have not enough variability to remove Fe rapidly when concentrations are high. The constant Fe/C ratios used in these two models may also contribute to this anomalous feature. In REcoM, Fe/C ratios are variable and the scavenging is second order but may simply be too low. DFe in TOPAZ has one of the shortest residence times (~8 years, Table 2), which implies that the surface accumulation of DFe may instead be linked to relatively large sources or the variable ligand concentration. Since the ligand concentration in TOPAZ depends on DOC, which typically decays from surface to deep, there may be too much DFe stabilization occurring in the surface ocean.

At intermediate depths, the inclusion of a prognostic Fe-binding ligand pool with a particle degradation source [Völker and Tagliabue, 2015] clearly improves the reproduction of subsurface maxima in DFe.
Figure 9. NO$_3$ concentrations ($\mu$M) from the GA-02 cruise [Rijkenberg et al., 2014] and extracted from the FeMIP models (NO$_3$ data not provided for GENIE).
Figure 10. PO₄ concentrations (μM) from the GA-02 cruise (Rijkenberg et al., 2014) and extracted from the FeMIP models (PO₄ not provided for MEDUSA-1, MEDUSA-2, REcoM, and TOPAZ).
Figure 11. Fe*(Fe–NO3 × rFe/N) from the GA-02 cruise [Rijkenberg et al., 2014] and extracted from the FeMIP models. For models that do not provide NO3, PO4 is used and converted to NO3 assuming a ratio of 16:1.
associated with remineralization (compare PISCES2 with PISCES1) for many of the transects. Other models (COBALT and to a lesser degree BEC and BLING) are able to reproduce these features but evidently do so for different reasons. These may be related to the implicit formulation of particle flux (BEC) that ignores lateral transport of particulate Fe or the shutdown of Fe scavenging in low oxygen conditions (BLING). It is interesting that there appears to be two groups of subsurface DFe maxima seen in the observations. Sometimes these features are tightly constrained to a small depth stratum (e.g., equatorial ocean for GA-02, western margin on GA-03, and eastern margin on CoFeMUG), while in other locations the DFe enrichments span almost the entire water column (eastern margins on GA-03 and GP-16). Most models represent one or the other. For example, subsurface maxima are always tightly bounded in depth for some models (e.g., COBALT and PISCES2) or spread over depth in others (BEC) with no regional variations. Future work should explore the potential mechanisms involved, which might be linked to subsurface dissolution of dust, nutrient trapping, or impacts of low oxygen. Emerging Fe isotope work highlights the potential for nonreductive Fe release from margins [Conway and John, 2014; Homoky et al., 2013] in addition to the role of reducing sediments represented in models.

In the ocean interior the best models (in terms of their linear correlation coefficients) are those that include hydrothermal input (Table 3). While including such a source is clearly important, it is possible that this is over-emphasized in the correlations at the expense of other deep ocean structure that is evident in many of the sections. For example, many of the ocean sections do not show any “water mass”-related structure for DFe that is seen in macronutrients (e.g., Figures 9 and 10). Although adding a hydrothermal ligand seems to improve the ability of PISCES2 to reproduce the GP-16 data (Figure 7) and perhaps also the GA-02 hydrothermal signal (Figure 4), it results in too widespread a hydrothermal anomaly in the Southern Ocean (Figure 8) indicating too long a lifetime for this pool and the need for further refinement of the processes governing hydrothermal Fe input [Tagliabue, 2014].

### 4.3. Intermodel Differences in DFe Inputs and Cycling: The Importance of Scavenging

It is notable that there is a great deal of variability in both the total Fe input flux ($66.9 \pm 67.1$ Gmol Fe yr$^{-1}$) and the strength of a given source across the models, yet the mean ocean DFe is strikingly similar ($0.58 \pm 0.14$ nM). To a large extent, this agreement reflects the calibration of scavenging rates and the concentration of organic ligands to obtain global average iron concentrations in agreement with observations. While this relative homogeneity in modeled mean DFe would be consistent with an earlier view of the oceanic Fe inventory [Johnson et al., 1997], if anything, the emerging oceanic sections of DFe as part of the GEOTRACES program have highlighted an unexpected variability in DFe distributions in the ocean interior [Mawji et al., 2015]. This is in stark contrast to the other main limiting nutrients, which more closely reflect large-scale ocean circulation patterns and water-mass-related features (e.g., Figures 9 and 10). Thus, the apparent small differences in the mean ocean DFe between models more likely arises from a modeling community that reflects an earlier parsimonious view of the system. The relative constancy in the mean ocean DFe concentrations in the models may reflect homogenous ligand concentrations of either 0.6 or 1.0 nM, but we note that even models with varying ligand concentrations (PISCES2 and TOPAZ) show too much interior ocean uniformity.

In contrast to the mean DFe, there is a substantial degree of intermodel disagreement in the strength of different sources. For instance, BFM, BLING, GENIE, MEDUSA1, MEDUSA2, MiTecco, MiTigsm, and REcoM all have atmospheric input fluxes of <5 Gmol Fe yr$^{-1}$, whereas as in BEC, COBALT, PISCES1, PISCES2, and TOPAZ dust supply is much higher (>20 Gmol Fe yr$^{-1}$). Yet this does not drive a similar trend in mean ocean DFe (with MiTecco, MiTigsm, and REcoM showing amongst the highest DFe concentrations, Table 2). We note that these represent the total DFe flux from dust, accounting for model-specific Fe mineralogy and solubility. Equally, for those models that include sedimentary Fe input, this flux term can range from very small (e.g., <5 Gmol Fe yr$^{-1}$ in MEDUSA2 or REcoM) to very large (>70 Gmol Fe yr$^{-1}$ in BEC, COBALT, MiTecco, MiTigsm, and TOPAZ). Again, this does not map onto mean DFe trends. We note that the closer agreement for hydrothermal Fe input is more likely to reflect the fact that only two models actually include this term rather than greater confidence regarding the actual flux. Overall, the total input of DFe does not explain the intermodel variations found in mean DFe ($R^2 = 0.06$). This implies that there must be a great deal of variability in how each model treats the scavenging of Fe in order to ultimately arrive at a relatively similar mean ocean DFe concentration.

Most early Fe models that explicitly computed free Fe and sought to represent its scavenging by sinking particles treated the scavenging rate constant as a tunable parameter [Archer and Johnson, 2000; Johnson et al.,
4.4. Impact of Fe on Wider Biogeochemical Cycles: The Importance of Biological Fe Cycling

The biological cycling of DFe in a given model will dictate the net influence of a model’s DFe cycling on wider biogeochemical cycling and air-sea CO2 exchange. In that regard, the large oceanic sections, focused process studies, and laboratory experiments all provide essential and complementary information. For example, early laboratory studies demonstrated a large degree of flexibility in the phytoplankton Fe/C ratios as a function of DFe levels and cell size, as well as enhanced Fe/C ratios at lower light levels [Sunda and Huntsman, 1997]. Similar ranges in Fe/C ratios are also seen in single-cell analyses of phytoplankton from the ocean [Twining and Baines, 2013]. The enhanced Fe/C ratio seen at low light is thought to reflect so-called “biodilution,” where Fe uptake continues when phytoplankton carbon fixation is light limited, and/or a greater absolute demand for Fe at low light [Sunda and Huntsman, 1997; Sunda and Huntsman, 1998]. Almost all FeMIP models permit flexibility in the Fe/C ratio of phytoplankton (Table 1), with those that consider Fe uptake independent of C fixation able to account for any biodilution, and the BLING model considers a direct impact of Fe on photosynthesis. Emerging recent work has suggested that there are important interspecific differences in how phytoplankton Fe demands respond to light [Strzepek et al., 2012]. In their laboratory study, Strzepek et al. [2012] found that while temperate diatom species indeed showed elevated Fe/C ratios at low light, the opposite was true for Antarctic diatom species. This raises questions about how models that generally do not consider different phytoplankton species (but rather represent broader “functional types”) can account for these potentially important regional distinctions in how environmental variations impact biological Fe cycling.

Detailed process studies, mostly from the Southern Ocean, have sought to quantify Fe cycling at the ecosystem level. In doing so, the importance of regenerated Fe in the fuelling of biological productivity via the so-called “ferrous wheel” has emerged as potentially important [Bowie et al., 2009, 2015; Boyd et al., 2012, 2015; Sarthou et al., 2008; Strzepek et al., 2005]. This has been demonstrated via the development of the “fe ratio,” which represents the proportion of Fe uptake from “new” Fe sources. It has been determined for sites across the Southern Ocean by assembling Fe budgets that combine measurements of Fe pools and fluxes alongside laboratory estimates. The Fe ratio is generally around 0.1 (i.e., strongly reliant on recycled Fe) in the low-productivity regions of the Southern Ocean [Bowie et al., 2009; Boyd et al., 2005] and reaches around 0.5 and greater (i.e., less reliant on recycled Fe) in the naturally fertilized Kerguelen Island phytoplankton bloom [Bowie et al., 2015; Sarthou et al., 2008]. Lagrangian process studies have demonstrated a strong seasonal decline in the Fe ratio as the spring phytoplankton bloom declines [Boyd et al., 2012], which are consistent with
low rates of Fe input during summer [Tagliabue et al., 2014c]. In agreement, direct measurements of Fe fluxes between various components of the food web have highlighted that only regenerative fluxes can support the measured Fe demand [Boyd et al., 2012; Strzepek et al., 2005; Tagliabue et al., 2014c].

The sensitivity of a given model’s biological productivity to new or regenerated forms of Fe is crucial, as this will underpin its sensitivity to change. At present we do not know if the FeMIP models place the correct emphases on new and recycled Fe in different ocean regions. Many models rely on fixed rates of Fe regenerated by zooplankton and the remineralization of organic material, while others allow this to vary (Table 1). A key parameter in driving the turnover of Fe by the zooplankton and bacterial communities in such models is an estimate of the heterotroph demand for Fe, which is then balanced against the Fe/C provided as nutrition. New measurements of stocks and turnover of Fe from specific ocean regions are also beginning to emerge [Boyd et al., 2015], which will be invaluable in assessing the magnitude and variability of the modeled rates.

5. Future Work

A weakness of the current intercomparison is that we did not truly intercompare the Fe models but instead compared the models’ coupled physical-biogeochemical framework (including Fe). This was necessary to retain as broad a suite of models as possible for this first intercomparison. In future work, it would be useful to intercompare different Fe models within the same physical model framework (e.g., as possible in the Nucleus for European Modelling of the Ocean (NEMO) or MITgcm modeling frameworks). Additionally, a set of planned model perturbations could be performed where each individual model is subjected to a modification to its Fe supply (either as a direct fertilization event or by an alteration to one of the input fields). Much could be learned from the way the Fe cycle responds to such perturbations across the different models.

Reducing uncertainty in the input fluxes of Fe is clearly important but has proved difficult to achieve over recent years (even for long recognized Fe sources such as dust). Some progress could be made by implementing “source-specific” tracers (such as aluminum or manganese) alongside Fe to constrain individual sources. Constraining scavenging rates has emerged as a key priority, and parallel simulation of Th may help constrain rates of Fe loss and the particle pools. Moreover, many of the models used specifically for ecological questions are only run for a few decades, leading to a greater sensitivity to initial conditions. A priority for such “resource-intensive” models would be the availability of input fields based on data climatologies (such as those available for macronutrients as part of the World Ocean Atlas data sets) or consensus distributions that may emerge from improved models.

As described in section 4.4 an assessment of the different biological Fe models is also a priority, as this will underpin the carbon cycle response and has not been compared against the paradigms recently emerging from experimental work. A follow-up phase of FeMIP could include a closer comparison of the models against the detailed process study measurements made (for example) as part of the FeCycle set of experiments [Boyd et al., 2005, 2012]. A range of the Fe models could be set up in a one-dimensional Lagrangian framework and forced by observed physics to be compared rigorously against the measured Fe stocks and cycling rates.

6. Conclusions

We have compared the projected DFe distributions from 13 global ocean biogeochemistry models against each other and with available data sets. Newly available full depth sections of DFe collected from different oceanic regions as part of the GEOTRACES program have greatly facilitated this task. All models do relatively poorly in reproducing a global DFe data set of around 20,000 observations, which highlights the need for greater understanding of how the ocean Fe cycle functions and how Fe should be represented in global ocean models. We find a large degree of intermodel variability in the input fluxes of DFe, which leads to great variability in the modeled residence times. The stronger intermodel agreement in the mean ocean DFe most likely reflects earlier views of constant deep ocean DFe levels maintained by a homogenous ligand pool and requires calibration via poorly constrained scavenging rates. The way different models treat DFe scavenging has emerged as a key uncertainty that would benefit from stronger observational constraints. More detailed intermodel tests, particularly linked to process study data, are needed to assess the models’ biological components.
In closing, we reemphasize the importance of the iron cycle in global ocean biogeochemistry models, given its role, alongside NO₃, as one of the two most important limiting nutrients. Although the models analyzed here struggle to capture the detailed distribution of this highly dynamic element, it is very likely that biogeochemical models that include an iron cycle can produce a more realistic simulation than models that do not. Improving the quantitative understanding of iron cycling should be a major priority for ocean biogeochemistry research.

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Erratum

In the originally published version of this article, Figure 6 contained incorrect labeling. The model data was labeled as GA-03 when it should have been labeled CoFeMUG. This error has since been corrected, and this version may be considered the authoritative version of record.