

## Evaluation of ocean carbon cycle models with data-based metrics

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[1] New radiocarbon and chlorofluorocarbon-11 data from the World Ocean Circulation Experiment are used to assess a suite of 19 ocean carbon cycle models. We use the distributions and inventories of these tracers as quantitative metrics of model skill and find that only about a quarter of the suite is consistent with the new data-based metrics. This should serve as a warning bell to the larger community that not all is well with current generation of ocean carbon cycle models. At the same time, this highlights the danger in simply using the available models to represent the state-of-the-art modeling without considering the credibility of each model. **INDEX TERMS:** 4255 Oceanography: General: Numerical modeling; 4805 Oceanography: Biological and Chemical: Biogeochemical cycles (1615); 4842 Oceanography: Biological and Chemical: Modeling; 4599 Oceanography: Physical: General or miscellaneous. **Citation:** Matsumoto, K., et al. (2004), Evaluation of ocean carbon cycle models with

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### 1. Introduction

[2] Three-dimensional ocean general circulation models (OGCMs) of the carbon cycle are the primary tools to characterize the ocean's response to increasing atmospheric CO<sub>2</sub> concentration. Although individual models are evaluated to various extents by their developers, how well they perform relative to other models has been difficult to assess. A primary reason is the absence of quantitative, data-based metrics of model skill. Further complications arise because models have different grids, numerics, boundary conditions, and parameterizations and because of the lack, until recently, of uniform simulation protocols. This difficulty represents a major stumbling block for the ocean carbon cycle modeling community. Outside the community, this in part causes the Intergovernmental Panel of Climate Change (IPCC), for example, to oversimplify the representation of the community with just the mean and spread of models that happened to be available at the time [Allen, 2003].

[3] We have come together under the Ocean Carbon Cycle Intercomparison Project (OCMIP) and formulated a set of standard protocols and global-scale data sets for model intercomparison and model evaluation of tracers and anthropogenic carbon [Dutay et al., 2002]. In this work, we have developed new quantitative metrics, based on the new CFC-11 and radiocarbon (<sup>14</sup>C) measurements from the World Ocean Circulation Experiment [Key et al., 1996, 2002; WOCE Data Products Committee, 2002], to assess the skill of a suite of models that have used the simulation protocols of the second phase of OCMIP (OCMIP-2). Our assessment brings to light large data-model discrepancies in ocean ventilation on decadal and centennial time scales. Our major goal is to provide some guidelines as to what constitutes a reasonable model behavior and to demonstrate that many of the models available today do not behave well.

### 2. Models and Data

[4] The suite of 19 models evaluated here includes those from the thirteen participating modeling groups of OCMIP-2 and a family of Princeton models (Table 1 and auxiliary material<sup>1</sup>). These models have dissimilar physical

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**Table 1.** OCMIP-2 and Princeton Sensitivity Models<sup>a</sup>

Model Name	Institutions (Country)
<i>OCMIP-2 Models</i>	
1) AWI	Alfred Wegener Inst. Polar Marine Res. (Germany)
2) CSIRO	CSIRO Division of Marine Research (Australia)
3) CCSR	Frontier Research for Global Change (Japan)
4) IPSL	Institute Pierre Simon LaPlace (France)
5) LLNL	Lawrence Livermore National Laboratory (USA)
6) MIT	Massachusetts Institute of Technology (USA)
7) MPIM	Max Planck Inst. Meteorologie-Hamburg (Germany)
8) NCAR	National Center for Atmospheric Research (USA)
9) NERSC	Nansen Env. and Remote Sensing Center (Norway)
10) PIUB	Physics Institute, University of Bern (Switzerland)
11) PRINCE	Princeton University (Low Kv; Low AI) (USA)
12) PRINC2	Princeton University (Intermediate Kv) (USA)
13) SOC	Southampton Oceanography Centre (UK)
14) UL	University of Liege (Belgium)
<i>Princeton Sensitivity Models (USA)</i>	
15) LH	Low Kv; High AI
16) HL	High Kv; Low AI
17) HH	High Kv; High AI
18) HiS	High Kv in Southern Ocean; otherwise PRINCE
19) SW	ECMWF reanalysis winds

<sup>a</sup>For most OCMIP-2 modeling groups the phase-2 models are not their latest. The exceptions are the Princeton models, some of which were developed more recently as part of in-house sensitivity studies [Gnanadesikan *et al.*, 2002]. The “IPSL” model presented here is the degraded version (IPSL.DM1 GM model). Kv and AI represent vertical and along-isopycnal mixing coefficients respectively.

models [Dutay *et al.*, 2002; S. C. Doney *et al.*, preprint, 2003] but use the same OCMIP-2 biogeochemistry formulations of air-sea gas exchange and seawater CO<sub>2</sub> system. Thus differences in the simulated tracer distributions can be attributed to different physical model behavior.

[5] For our data-model comparisons, we focus on two passive tracers: CFC-11, which is measured directly, and natural <sup>14</sup>C, which is relatively straightforward to calculate from data. There is little ambiguity associated with either tracer when interpreting data-model discrepancies. In addition, these two tracers provide information on the ocean circulation that brackets the decadal to centennial timescales relevant for the uptake of anthropogenic CO<sub>2</sub>. The need to separate natural <sup>14</sup>C, which is produced only in the atmosphere, from the measured abundance [Broecker *et al.*, 1995; Rubin and Key, 2002] arises because oceanic <sup>14</sup>C has been significantly altered in the upper ocean over the last decades with the introduction of thermonuclear bomb-produced <sup>14</sup>C from the atmosphere. Over most of the ocean, the penetration of the bomb <sup>14</sup>C is shallow and does not extend beyond several hundred meters. Therefore, with the exception of the North Atlantic where North Atlantic Deep Water (NADW) is formed, and locations around Antarctica where Antarctic Bottom Water is formed, the deep ocean can be assumed to be free of bomb <sup>14</sup>C, eliminating most of the uncertainties associated with the separation.

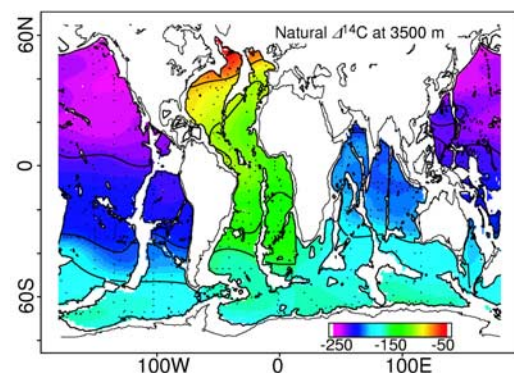
### 3. Results and Discussion

[6] The deep ocean ventilation is readily assessed by the distribution of natural <sup>14</sup>C, whose abundance is a direct measure of ocean ventilation on centennial time scales. We make data-model comparison of the deep natural <sup>14</sup>C distribution by examining the natural <sup>14</sup>C abundance of NADW, the North Pacific Deep Water (NPDW), and the

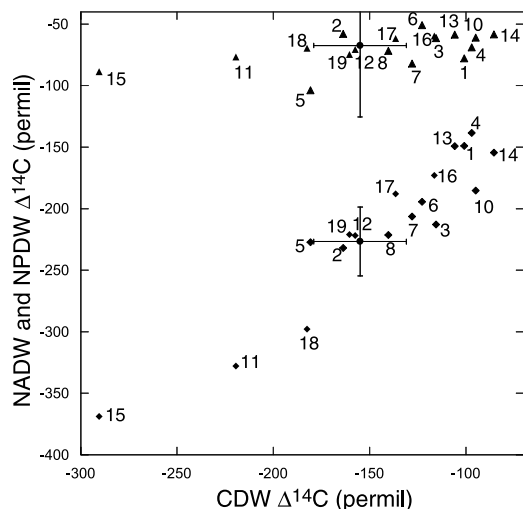
Circumpolar Deep Water (CDW). The three water masses are important “end-members” that can characterize the deep ocean. Their typical natural <sup>14</sup>C abundance, expressed with  $\Delta^{14}\text{C}$  in per mil (‰), as determined from measurements, are  $-70\text{‰}$ ,  $-160\text{‰}$ , and  $-230\text{‰}$  respectively for NADW, CDW, and NPDW (Figure 1; see auxiliary material for water mass definition). The oldest NPDW has the most negative and least  $\Delta^{14}\text{C}$ , and the most recently ventilated NADW has the highest <sup>14</sup>C abundance. Figure 2 shows that simulated natural  $\Delta^{14}\text{C}$  varies substantially [Orr, 2002], particularly in the North Pacific and Southern Ocean. In the North Pacific,  $\Delta^{14}\text{C}$  varies between  $-138\text{‰}$  and  $-380\text{‰}$  from the most ventilated (OCMIP-2 IPSL) model to the most stagnant (Princeton LH), respectively. Many OCMIP-2 models ventilate the deep Pacific excessively. Interestingly, natural  $\Delta^{14}\text{C}$  is correlated rather well between NPDW and CDW (diamonds, Figure 2) unlike that between NADW and CDW (triangles, Figure 2), suggesting the important role that CDW plays in these models in ventilating the deep Pacific.

[7] Ocean ventilation on shorter, decadal time scales is better characterized by the presence of CFC-11. Figure 3 shows the combined Indo-Pacific inventories of CFC-11 inside and outside the Southern Ocean relative to long term, deep Southern Ocean ventilation, as indicated by the CDW  $\Delta^{14}\text{C}$ . This figure allows us to evaluate models in their decadal and centennial time scale ventilation at the same time. We examine the Southern Ocean separately because of the large inter-model variability in CFC uptake [Dutay *et al.*, 2002] and in natural <sup>14</sup>C abundance in this region (Figure 2).

[8] In the Southern Ocean, model-based inventories of CFC-11 are positively correlated with CDW  $\Delta^{14}\text{C}$  (Figure 3c), indicating the importance of deep vertical transport (mixing and advection) in these models that carries both CFC-11 and natural <sup>14</sup>C into the deep ocean. Model-simulated inventories encompass the observations but vary roughly by a factor of 4 between the most stratified models (i.e., Princeton LH and LL) and the most vertically well-mixed models (OCMIP-2 AWI, NERSC, UL). The large inter-model variability in the Southern Ocean [Orr *et al.*, 2001] contrasts with a much smaller variability outside the Southern Ocean (Figure 3a), suggesting how sensitive Southern Ocean tracer uptake is to the formulations of surface forcing and physical parameterizations in the current generation of OGCMs.



**Figure 1.** Data-based natural  $\Delta^{14}\text{C}$  on 3500 m level [after Matsumoto and Key, 2004].



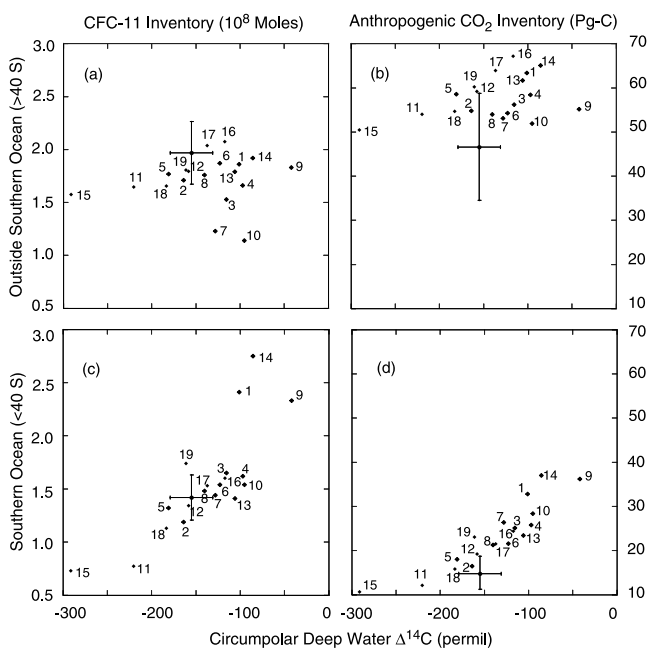
**Figure 2.** Observed and modeled deep natural  $\Delta^{14}\text{C}$ . Symbols with error bars ( $2\sigma$ ) are observations. See auxiliary material for the definitions of North Atlantic Deep Water (NADW), North Pacific Deep Water (NPDW), and Circumpolar Deep Water (CDW). Diamonds and triangles are model-predicted mean NPDW and NADW  $\Delta^{14}\text{C}$  respectively, vs. model predicted CDW. See Table 1 for model indices. The results of model #9 (NERSC) are excluded here, because the model had not been run out to equilibrium.

[9] We can then use the observed natural  $^{14}\text{C}$  distribution and CFC-11 inventories (Figures 3a and 3c) as metrics of model skill to identify those models that are most consistent with the observations. For reasonable uncertainties of  $2\sigma$  for mean CDW  $\Delta^{14}\text{C}$  and 15% for CFC-11 inventories (see auxiliary material<sup>5</sup>), there are only 4 out of 19 models that agree with observations. For smaller uncertainties of  $1\sigma$  and 10%, there is but one model. For larger uncertainties of  $3\sigma$  and 20%, there are seven models. The majority of the models analyzed in this study are thus not ventilating the ocean on decadal and centennial time scales as indicated by the observed CFC-11 and natural radiocarbon.

[10] Presumably these smaller groupings of the “better” models make more reliable estimates of the global oceanic uptake of anthropogenic carbon than larger groupings that include models that are not as consistent with data. The uptake estimates from the subset of four models, which are essentially the same as those from the subsets of one and seven models, are  $1.9 \pm 0.2$  and  $2.2 \pm 0.2$  Pg-C/yr ( $\text{Pg} = 10^{15}\text{g}$ ; the uncertainty of  $\pm 0.2$  represents half the range of the model spread) during the 1980s and 1990s respectively. We made a 3% downward correction [Orr, 2002] to the 1990s estimates, which are from simulations using the IPCC S650 scenario, whose 1990s atmospheric  $\text{CO}_2$  concentrations were slightly higher than observed. The new estimates are somewhat lower and, as expected, have smaller spread than those from the full set of OCMIP-2 models for both the 1980s ( $2.0 \pm 0.4$  Pg-C/yr) and 1990s ( $2.4 \pm 0.5$  Pg-C/yr) [Orr, 2002]. For the 1980s, the model-based estimates are consistent with the most recent IPCC estimate of  $1.9 \pm 0.6$  Pg-C/yr, based on atmospheric  $\text{CO}_2$  and  $\text{O}_2$  measurements [Houghton et al., 2001], and with  $1.6 \pm 0.4$  Pg-C/yr, based on global CFC distribution [McNeil et al., 2003]. It is

difficult to directly translate the new model-based estimates into reality, because we do not account for a number of important processes, including interannual variability and climate-carbon feedbacks. Therefore, our new estimates should simply be regarded as improved model-based estimates without necessarily implying an improved understanding of the true global carbon budget.

[11] Finally, we draw attention to the comparison of the model-simulated anthropogenic  $\text{CO}_2$  inventories with those reconstructed from ocean carbon observations using the  $\Delta\text{C}^*$  method [Gruber et al., 1996]. In Figures 3b and 3d, we show only the Indian [Sabine et al., 1999] and Pacific [Sabine et al., 2002] basins, because the paucity of WOCE data in the far North Atlantic makes it difficult to determine the preformed properties of NADW (i.e., properties of seawater when it was last at the surface before it became part of NADW) and hence anthropogenic carbon. The comparison suggests that the models that successfully simulate the correct CDW  $\Delta^{14}\text{C}$  and CFC-11 inventories tend to simulate higher anthropogenic  $\text{CO}_2$  inventories than the data-based estimates. This is clearly evident north of  $40^\circ\text{S}$ , where simulated anthropogenic carbon inventories are on average 10 Pg-C higher than the data-based inventory of  $47 \pm 7$  Pg-C. Our analysis shows that much of the discrepancy originates from the Pacific inventories. It may result from possible model deficiencies, such as misrepresentation of the oceanic buffering capacity or of the air-sea gas exchange, or possible biases in the data-based estimates.



**Figure 3.** Observed and modeled CFC-11 and anthropogenic carbon inventories in the Indian and Pacific Ocean versus Southern Ocean natural  $\Delta^{14}\text{C}$ . Observed inventories are based on WOCE CFC-11 measurements [WOCE Data Products Committee, 2002] and anthropogenic carbon reconstructions [Sabine et al., 2002, 1999]. Error bars represent  $2\sigma$  for natural  $^{14}\text{C}$ , 15% uncertainty for CFC-11 inventories, and 25% uncertainty for anthropogenic carbon inventories (see auxiliary material). Southern Ocean is defined here as south of  $40^\circ\text{S}$ .

At the same time, our results highlight the importance of better understanding the fidelity of the data-based estimates of anthropogenic carbon, which have recently been suggested as being too low [Hall et al., 2004].

#### 4. Conclusions

[12] We have developed quantitative, data-based metrics of ocean carbon cycle model skill and show that only about a quarter of the 19 models we examined are reasonably consistent with those metrics. While the simple metrics we introduced here are certainly not the best or the only ones, our study illustrates clearly that some models are inadequate. This recognition is timely for the larger community today, when ocean carbon cycle models are becoming more numerous, and relatively easy metrics to assess them are needed to ensure quality control. Identifying what in the models are responsible for this inadequacy will require sensitivity studies, where important model parameterizations and physics are changed one at a time.

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