1	Ocean acidification over the next three centuries using a simple global climate carbon-cycle model:
2	projections and sensitivities.
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11	ABSTRACT
12	Continued oceanic uptake of anthropogenic CO_2 is projected to significantly alter the chemistry of the
13	upper oceans over the next three centuries, with potentially serious consequences for marine
14	ecosystems. Relatively few models have the capability to make projections of ocean acidification,
15	limiting our ability to assess the impacts and probabilities of ocean changes. In this study we examine
16	the ability of Hector v1.1, a reduced-form global model, to project changes in the upper ocean
17	carbonate system over the next three centuries, and quantify the model's sensitivity to parametric
18	inputs. Hector is run under prescribed emission pathways from the Representative Concentration
19	Pathways (RCPs) and compared to both observations and a suite of Coupled Model Intercomparison
20	(CMIP5) model outputs. Current observations confirm that ocean acidification is already taking place,
21	and CMIP5 models project significant changes occurring to 2300. Hector is consistent with the
22	observational record within both the high (>55°) and low latitude oceans (<55°). The model projects low
23	latitude surface ocean pH to decrease from preindustrial levels of 8.17 to 7.77 in 2100, and to 7.50 in
24	2300; aragonite saturation levels (Ω_{Ar}) decrease from 4.1 units to 2.2 in 2100 and 1.4 in 2300 under RCP

25	8.5. These magnitudes and trends of ocean acidification within Hector are largely consistent with the
26	CMIP5 model outputs, although we identify some small biases within Hector's carbonate system. Of the
27	parameters tested, changes in $[H^+]$ are most sensitive to parameters that directly affect atmospheric
28	CO_2 concentrations - Q_{10} (terrestrial respiration temperature response) as well as changes in ocean
29	circulation, while changes in Ω_{Ar} saturation levels are sensitive to changes in ocean salinity and Q_{10} . We
30	conclude that Hector is a robust tool well-suited for rapid ocean acidification projections, sensitivity
31	analyses, and is capable of emulating both current observations and large scale climate models under
32	multiple emission pathways.

1. INTRODUCTION

Human activities have led to increasing anthropogenic emissions of greenhouse gases to the atmosphere. In the first decade of the 21st century CO₂ emissions from anthropogenic sources and land use changes accounted for ~9 Pg C yr⁻¹, with future emission projections of up to 28 Pg C yr⁻¹ by 2100 under RCP 8.5 (Riahi et al., 2011). The world's oceans have played a critical role in lessening the effects of climate change by absorbing 25-30% of the total anthropogenic carbon emissions since 1750 (Le Quéré et al., 2013; Sabine et al., 2011).

40 In response to this increasing atmospheric burden of CO_2 and increasing oceanic uptake, the oceans are experiencing both physical and biogeochemical changes: surface and deep water warming, 41 42 reduced subsurface oxygen, and a reduction in calcium carbonate saturation levels and pH (Doney, 43 2010). Mean surface ocean pH has already decreased by 0.1 units relative to preindustrial times 44 (Caldeira et al., 2003). If current emission trends continue, ocean acidification will occur at rates and 45 extents not observed over the last few million years (Feely et al., 2004; Feely et al., 2009; Kump et al., 46 2009; Caldeira et al., 2003). Ocean acidification occurs when atmospheric CO₂ dissolves in seawater $(CO_2(aq))$, forming carbonic acid (H_2CO_3) ; dissociating into carbonate (CO_3^{2-}) and bicarbonate (HCO_3^{-}) , 47 and releasing protons (H^+). The net effect of adding CO₂ to the system increases the concentrations of 48 $[H_2CO_3]$, $[HCO_3^-]$, and $[H^+]$, while decreasing $[CO_3^{2-}]$ concentrations and lowering the pH. The sum of 49 50 $[HCO_3^-]$, $[CO_3^{2-}]$ and $[CO_2^*]$, where $[CO_2^*] = [CO_2(aq)] + [H_2CO_3]$ represents the dissolved inorganic (DIC) of the system. As $CO_2(aq)$ continues to increase in the ocean it reacts with CO_3^{2-} , forming HCO_3^{-} , decreasing 51 52 the fraction of CO_2 that can be readily absorbed by the oceans. Because of this capacity of the ocean to 53 buffer chemical changes, a doubling of atmospheric $[CO_2]$ will not correspond to a doubling of $[CO_2^*]$ but 54 instead will result in an increase on the order of 10% (Dickson and Millero, 1987). Due to both chemical 55 and physical changes (e.g., warming and stratification), the oceans may become less efficient in the

uptake of anthropogenic CO₂ as the climate continues to change (Sarmiento and Le Quéré, 1996; Matear
and Hirst, 1999; Joos et al., 1999; Le Quéré et al., 2010).

58 Numerous experiments and observations indicate that ocean acidification will have significant 59 effects on calcifying marine organisms. For example, the rate of coral reef building may decrease, 60 calcification rates of planktonic cocolithophores and foraminifera may be suppressed, and significant 61 changes in trophic level interactions and ecosystems may occur (Cooley and Doney, 2009; Silverman et 62 al., 2009; Fabry et al., 2008; Riebesell et al., 2000). Some coral reefs are believed to already be eroding 63 for parts of the year due to ocean acidification (Yates and Halley, 2006; Albright et al., 2013). Global 64 surface pH is projected to drop by up to 0.33 units (Gehlen et al., 2014; Orr et al., 2005) and all existing 65 coral reefs will be surrounded by conditions well outside of preindustrial values and even today's saturation levels (Ricke et al., 2013) under the RCP8.5 scenario. 66

67 These model projections of ocean acidification come primarily from Earth System Models 68 (ESMs) that integrate the interactions of atmosphere, ocean, land, ice and biosphere to estimate the 69 present and future state of the climate. ESMs are computationally expensive and typically run using 70 stylized experiments or a few RCPs (greenhouse gas concentration trajectories used in the Intergovernmental Panel on Climate Change 5th Assessment Report (IPCC, 2013)). This generally limits 71 72 ESM-based analyses to those scenarios. The occurrence of ocean warming and acidification is consistent 73 across CMIP5 ESMs, but their rates and magnitudes are strongly dependent upon the scenario (Bopp et 74 al., 2013).

An alternative to ESMs are reduced-form models, relatively simple and small models that can be powerful tools due to their simple input requirements, computational efficiency, tractability, and thus ability to run multiple simulations under arbitrary future emission pathways. This allows for the quantification of arbitrary climate change scenarios, emulation of larger ESMs, as well as in-depth

parameter sensitivity studies and uncertainty analyses (Senior and Mitchell, 2000; Ricciuto et al., 2008;
Irvine et al., 2012).

Our goal of this study is to quantify how well Hector; a reduced-form model that explicitly treats surface ocean chemistry; emulates the marine carbonate system of both observations and the CMIP5 archive; and explore the parametric sensitivities to Hector's ocean outputs. The remainder of the paper is organized as follows; section 2, a detailed description of Hector's ocean component, the data sources and simulations run, section 3, results of the model comparison and sensitivity experiments and lastly, section 4, a discussion of the results.

87

88 2 Model Description – Hector

89 Hector is open-source and available at https://github.com/JGCRI/hector. The repository 90 includes all model code needed to compile and run the model, as well as all input files and R scripts to 91 process its output. Hector is a reduced form climate carbon-cycle model, which takes in emissions of 92 CO₂, non-CO₂s (e.g., CH₄, N₂O and halocarbons, and aerosols), converts emissions to concentrations 93 where needed, calculates the global radiative forcing, and then global mean temperature change. 94 Hector contains a well-mixed global atmosphere, a land component consisting of vegetation, detritus, 95 and soil, and an ocean component. In this study we use Hector v1.1, with an updated ocean 96 temperature algorithm to better match the CMIP5 mean. For a detailed description of the land and 97 atmospheric components of Hector, please refer to Appendix A and Hartin et al. (2015). 98 2.1 Ocean Component

99 Hector's ocean component is based on work by Lenton (2000), Knox and McElroy (1984) and
100 Sarmiento and Toggweiler (1984). It consists of four boxes: two surface boxes (high and low latitude),
101 one intermediate, and one deep box. The cold high latitude surface box makes up 15% of the ocean
102 surface area, representing the subpolar gyres (> 55°), while the warm low latitude surface box (<55°)

makes up 85% of the ocean surface area. The temperatures of the surface boxes are linearly related to
the global atmospheric temperature change, and are initialized at 2°C and 22°C for the high and low
latitude boxes respectively. This temperature gradient sets up a flux of carbon into the cold high latitude
box and a flux out of the warm low latitude box. The ocean-atmosphere flux of carbon is the sum of all
the surface fluxes (F_i, n=2).

$$F_{O}(t) = \sum_{i=1}^{n} F_{i}(t)$$
⁽¹⁾

108Once carbon enters the high latitude surface box it is circulated between the boxes via109advection and water mass exchange, simulating a simple thermohaline circulation. In this version of110Hector we do not explicitly model diffusion; simple box-diffusion models and "HILDA" (e.g., Siegenthaler111and Joos, 1992) type models are typically in good agreement with observations but are more112computationally demanding than a simple box model (Lenton, 2000). The change in carbon of any113ocean box *i* is given by the fluxes in and out (j) with F_{atm->i} as the atmospheric carbon flux of the two114surface boxes:

$$\frac{dC_i}{dt} = \sum_{j=1}^{in} F_{j \to i} - \sum_{j=1}^{out} F_{i \to j} + F_{atm \to i}$$
⁽²⁾

115 The flux of carbon between the boxes is related to the transport $(T_{i\rightarrow j}, m^3 s^{-1})$ between *i* and *j*, the 116 volume of *i* (V_i, m³), and the total carbon in *i* (including any air-sea fluxes) (C_i, Pg C);

$$\frac{dC_{i \to j}}{dt} = \frac{T_{i \to j} * C_i(t)}{V_i}$$
(3)

Volume transports are tuned to yield an approximate flow of 100 Pg C from the surface high latitude box
 to the deep ocean box at steady state, simulating deep water formation.
 Hector calculates DIC, total alkalinity (TA), CO₃²⁻, HCO₃⁻ pCO₂ and pH. DIC and TA are the two

120 carbonate variables used to solve the rest of the carbonate system. The detailed carbonate chemistry

121 equations are based on numeric programs from Zeebe and Wolf-Gladrow, (2001) (Appendix B). We

122 simplified the equations by neglecting the effects of pressure, since we are only concerned with the 123 surface ocean. Hector is run to equilibrium in a perturbation-free mode, prior to running the historical 124 period, ensuring that it is in steady state (Hartin et al., 2015; Pietsch and Hasenauer, 2006). DIC (µmol 125 kg⁻¹) in the surface boxes is a function of the total carbon (Pg C) and the volume of the box. All carbon 126 within the ocean component is assumed to be inorganic carbon. Dissolved organic matter is less than 127 2% of the total inorganic carbon pool, of which a small fraction is dissolved organic carbon (Hansell and 128 Carlson, 2001) and particulate organic carbon less than 1% of the total carbon pool (Eglinton and 129 Repeta, 2004). Therefore, for simplicity we chose not to include dissolved or particulate organic carbon 130 within Hector.

131 TA is calculated at the end of model spinup (running to equilibrium in an a historical,

132 perturbation-free mode) and held constant throughout the run, resulting in 2311.0 μ mol kg⁻¹ for the

high latitude box and 2435.0 μmol kg⁻¹ for the low latitude box. These values are within the range of

open ocean observations of 2250.0 – 2450.0 μ mol kg⁻¹ (Key et al., 2004; Fry et al., 2015). We assume

135 negligible carbonate precipitation/dissolution and no alkalinity runoff from the land surface to the open

136 ocean. Alkalinity is typically held constant with time, which is a reasonable assumption over several

thousand years (Lenton, 2000; Zeebe and Wolf-Gladrow, 2001; Glotter et al., 2014; Archer et al., 2009).

138 On glacial-interglacial time scales alkalinity and the dissolution of CaCO₃ sediments is an important

factor in controlling atmospheric [CO₂] (Sarmiento and Gruber, 2006), and thus on these scales Hector

140 will underestimate the oceanic CO₂ uptake.

Hector solves for pCO₂, pH (total scale), and [HCO₃⁻], [CO₃²⁻], and aragonite (Ω_{Ar}) and calcite saturation states(Ω_{Ca}) in both the high and low latitude surface ocean boxes. pCO₂ is calculated from the concentration of [CO₂^{*}] and the solubility of CO₂ in seawater, based on salinity and temperature. [CO₂^{*}] is calculated from DIC and the first and second dissociation constants of carbonic acid from Mehrbach et al. (1973), refit by Lueker et al. (2000) (Appendix B).

Carbon fluxes between the atmosphere and ocean are calculated (Takahashi et al., 2009):

$$F = k \alpha * \Delta p CO_2 = Tr * \Delta p CO_2$$
(4)

147 where k is the CO₂ gas-transfer velocity, α is the solubility of CO₂ in seawater (K₀, Appendix B), and the 148 ΔpCO_2 is the difference in pCO₂ between the atmosphere and ocean. The product of k and α results in Tr, the sea-air gas transfer coefficient, where Tr (g C m⁻² month⁻¹ μ atm⁻¹) = 0.585 * α * Sc^{-1/2} * U₁₀², 0.585 149 is a unit conversion factor (from mol liter⁻¹ atm⁻¹ to g-C m⁻³ µatm⁻¹ and from cm h⁻¹ to m month⁻¹) and Sc 150 151 is the Schmidt number. The Schmidt number (Appendix B) is calculated from Wanninkhof (1992) based 152 on the temperature of each surface box. The average wind speed (U_{10}) of 6.7 m s⁻¹ is the same over 153 both surface boxes (Table 1). pH (total scale), $[HCO_3^-]$, and $[CO_3^{-2}]$ are calculated using the $[H^+]$ ion and 154 solved for in a higher order polynomial (Appendix B). 155 Aragonite and calcite are the two primary carbonate minerals within seawater. The degree of 156 saturation in seawater with respect to aragonite (Ω_{Ar}) and calcite (Ω_{Ca}) is calculated from the product of 157 the concentrations of calcium $[Ca^{2+}]$ and carbonate ions $[CO_3^{2-}]$, divided by the solubility product (K_{sp}) . 158 The $[Ca^{2+}]$ is based on equations from Riley and Tongudai (1967) at a constant salinity of 34.5. If $\Omega = 1$,

159 the solution is at equilibrium, and if $\Omega > 1$ ($\Omega < 1$) the solution is supersaturated (undersaturated) with 160 respect to the mineral.

$$\Omega = \frac{[Ca^{2+}] [CO_3^{2-}]}{K_{sp}}$$
(5)

161 **2.2. Simulation and experiments**

Hector is run under prescribed emissions from 1850-2300 for all four Representative
Concentration Pathways (RCP 2.6, RCP 4.5, RCP 6, RCP 8.5) (Moss et al., 2010; van Vuuren et al., 2007).
We compare how well Hector can emulate the carbonate system of the CMIP5 median. Our results
section will mainly focus on RCP8.5 exploring the response of the carbonate system under a high
emissions scenario.

167 We also ran a series of model sensitivity experiments to quantify how influential some of 168 Hector's parameter inputs are on its outputs (in particular, $[H^+]$ and Ω_{Ar}). Such sensitivity analyses are 169 important to document model characteristics, explore model weaknesses, and determine to what 170 degree the model behavior conforms to our existing understanding of the ocean system. We do not 171 sample Hector's entire parameter space, a computationally demanding exercise, but instead choose a 172 list of the parameters that we expect, a priori, to be important in calculating the marine carbonate 173 system. We selected those parameters directly influencing atmospheric CO₂ concentrations (beta and 174 Q_{10}), parameters involved in the calculation of temperature (albedo and climate sensitivity) and those 175 parameters involved in the uptake of carbon in the surface ocean (ocean surface temperature, salinity, wind stress, and ocean circulation). These parameters are varied by ±10% relative to the RCP8.5 control, 176 177 and we compare the percentage change from the reference and the perturbation cases in 2005, 2100, 178 and 2300. The reference, RCP8.5, refers to the tuned set of parameters found in Hector v1.1, resulting 179 in Figures 2-6. 180 2.3 Data Sources 181 All RCP input emission data are available at http://tntcat.iiasa.ac.at/RcpDb/. Comparison data are

obtained from a suite of CMIP5 Earth System Models (Table 2) (Taylor et al., 2012). The CMIP5 output is
 available from the Program for Climate Model Diagnostics and Intercomparison

184 (http://pcmdi3.llnl.gov/esgcet/home.htm). We took the 0-100m (depth) mean for all available CMIP5

data for six output variables, computing the monthly mean for all years in the historical (1850-2005) and

186 RCP 8.5 (2006-2300) experiments. All outputs were regridded to a standard 1-degree grid using bilinear

- 187 interpolation in CDO version 1.7.1rc1, and then high latitude (-90 to -55 and 55 to 90 degrees), low
- 188 latitude (-55 to 55), and global area-weighted means computed using R 3.2.4. All CMIP5 comparisons
- used in this study are from model runs with prescribed atmospheric CO₂ concentrations. We
- acknowledge that this is not a perfect comparison, as Hector is emissions-forced being compared to the

concentration-forced CMIP5 models, but very few of the latter were run under prescribed emissions.
We use a combination of root mean square error (RMSE), rates of change (Δ) and bias (degree of
systematic over- or underestimation) to characterize Hector's performance relative to the CMIP5
median.

195 We also compare Hector to a series of observational ocean data. Surface ocean observations of 196 DIC, pCO₂, pH, Ω_{Ar} , and Ω_{Ca} are from time-series stations in both the high and latitude oceans; Hawaii 197 Ocean Time Series (HOT), Bermuda Atlantic Time Series (BATS), the European Station for Time Series in 198 the Ocean at the Canary Islands (ESTOC), the Irminger Sea, and the Iceland Sea (Table 3). The time-199 series data are annually averaged over the upper 100m of the water column. The carbonate parameters 200 not found in Table 3 are computed from temperature, salinity, and the given carbonate parameter pairs 201 using the CO2SYS software (Lewis and Wallace, 1998). The equilibrium constants (K1 and K2 from 202 Mehrbach et al., 1973 refit by Dickson and Millero, 1987) and zero total phosphorus and silica where 203 chosen to best match Hector. Lastly, a longer record (1708 – 1988) of pH and Ω_{Ar} from Flinder's Reef in 204 the western Coral Sea, calculated from boron isotope measurements, is used in the comparison 205 (Pelejero et al., 2005). We use rates of change (Δ) from 1988-2014, which overlaps the HOT and BATS 206 time series, to quantify how well Hector simulates the observed changes in the ocean carbonate 207 parameters (Table 6) (Dore et al., 2009; Bates et al., 2014).

208

209 3. Results

210 3.1 Model and Observation Comparisons

Hartin et al. (2015) conducted a thorough analysis of Hector v1.0 demonstrating that it can reproduce the historical trends and future projections of atmospheric [CO₂], radiative forcing, and global temperature change under the four RCPs. In this study we focus on the upper ocean high and low latitude inorganic carbon chemistry under RCP 8.5.

215 Hector captures the trend in DIC concentrations for both the high and low latitude surface 216 ocean with a global RMSE average of 7.0 μ mol kg⁻¹ when compared to CMIP5 models over the historical 217 period (Table 4; Figure 2). We note that there is a systematic bias in both the high and low latitude 218 surface boxes when compared to CMIP5. First, the carbon pools of the surface boxes are initialized with 219 carbon values slightly higher than the median CMIP5 values. Second, after 2100 the high latitude CMIP5 220 median begins to decline, while Hector rises and stabilizes. Only 3 CMIP5 models ran out to 2300, with 221 one model driving the decline. Regardless, this offset only results in a <3% global difference between 222 CMIP5 and Hector.

Hector accurately tracks the pCO₂ in both the high and low latitude surface ocean with similar rates of change from 1850-2300 (Figure 3). There is a low bias in Hector compared to CMIP5 models after 2100, due to the low bias in projected atmospheric [CO₂] within Hector over the same time period (Hartin et al., 2015). We do find Hector to be in closer agreement with the observation record.

Figure 4 shows the high and low latitude surface pH of Hector compared to CMIP5 and observations from BATS, HOT, ESTOC, Irminger Sea, Iceland Sea, and Flinders Reef. While the high latitude surface pH is slightly higher than the CMIP5 models, Hector is more similar to high latitude observations. Since the preindustrial, observations of surface ocean pH decreased by 0.08 units, corresponding to a 24% increase in [H⁺] concentrations and an 8% decrease in [CO₃²⁻], similar to numerous studies (Feely et al., 2004; Sabine et al., 2004; Caldeira et al., 2003; Orr et al., 2005) that

233 estimate an average global decrease in pH of 0.1 or a 30% increase in [H⁺].

The Flinder's Reef pH record provides a natural baseline to compare future trends in ocean acidification. While we did not expect the model to match exactly, as this reef site is heavily influenced by coastal dynamics and internal variability, rates of change from the preindustrial (1750) to 1988 are the same (0.0002 yr⁻¹) for both Hector and Flinder's Reef. Over the limited observational record from both the Pacific and Atlantic Oceans, Hector accurately simulates the decline in pH (-0.0017 yr⁻¹)

239 compared to observations (Table 6). Other observations in the North Pacific show surface changes of 240 pH up to 0.06 units between 1991 and 2006 with an average rate of -0.0017 yr⁻¹ (Byrne et al., 2010). 241 Recent work suggests that the North Atlantic absorbed 50% more anthropogenic CO₂ in the last decade 242 compared to the previous decade, decreasing surface pH by 0.0021 (Woosley et al., 2016). Under RCP 243 8.5, Hector projects a decrease in low latitude pH of 8.17 in 1850 to 7.77 in 2100 and down to 7.5 by 244 2300, similar to CMIP5 (Table 5). At approximately 2050, atmospheric [CO₂] is twice the preindustrial 245 concentrations, corresponding to a decrease in pH to 7.96. Shortly after this doubling, pH values are 246 well outside the natural variability found in Flinder's Reef.

247 Figure 5 illustrates the high and low latitude surface Ω_{Ar} . We only highlight Ω_{Ar} , as Ω_{Ca} is similar 248 to that of Ω_{Ar} . As with pH, Hector is slightly higher than the CMIP5 Ω_{Ar} median but closer to the 249 observational record. Hector accurately simulates the change in Ω_{Ar} (-0.0090 yr⁻¹) compared to 250 observations (Table 6). Repeated oceanographic surveys in the Pacific Ocean have observed an average 251 0.34% yr⁻¹ decrease in the saturation state of surface seawater with respect to aragonite and calcite over 252 a 14-year period (1991-2005) (Feely et al., 2012); the average decrease in Hector is between 0.19% yr⁻¹ 253 and 0.25% yr⁻¹. Saturation levels of Ω_{Ar} decrease rapidly over the next 100 years in both the high and 254 low latitude. Hector accurately captures the decline in saturation states with low RMSE values for Ω_{Ar} . 255 Under RCP8.5 Hector projects that low latitude Ω_{Ar} will decrease to 2.2 by 2100 and down to 1.4 by 256 2300. The high latitude oceans will be undersaturated with respect to aragonite by 2100 and will drop 257 down to 0.7 by 2300.

Lastly, Figure 6 highlights pH and Ω_{Ar} projections under all four RCPs from 1850 to 2300. Over the last 20 years, both pH and Ω_{Ar} have declined sharply and will continue to decline under RCP 4.5, 6.0 and 8.5, outside of their preindustrial and present day values. These RCPs represent a range of possible future scenarios, with ocean pH varying between 8.15 and 7.46 for the high latitude and Ω_{Ar} varying between 1.94 and 0.60. High latitude Ω_{Ar} saturation levels presently are much lower than the low

latitude and reach undersaturation before the end of the century. Even under a best-case scenario, RCP
2.6, low latitude pH will drop to 8.07 by 2100 and to 8.12 by 2300, with Ω_{Ar} saturation states remaining
well outside of present day values.

266 3.2 Model Parameter Sensitivity

Parametric sensitives are different between $[H^+]$ and Ω_{Ar} . We use $[H^+]$ to highlight changes in pH, as pH = -log $[H^+]$. In the near term (from 2005-2100) the calculation of pH is sensitive to a combination of parameters, ocean circulation, beta (terrestrial CO₂ fertilization), and wind stress, while on longer time scales (to 2300), $[H^+]$ is most sensitive to changes in Q₁₀ (terrestrial respiration temperature response) and ocean circulation (Table 7). Global Ω_{Ar} is most sensitive to changes in salinity in both the near and long term. Similar to $[H^+]$, Ω_{Ar} becomes more sensitive to changes in Q₁₀ in the long term.

274

275 4. Discussion

276 The marine carbonate system is projected to undergo significant changes under the RCPs. pCO₂ 277 and DIC are increasing rapidly as atmospheric [CO₂] continues to rise under RCP 4.5, 6.0 and 8.5. pH, and 278 Ω_{Ar} are decreasing rapidly outside of observations and are projected to continue to decrease under all 279 scenarios (Figure 6). Only under RCP 2.6 do pH and Ω_{Ar} values begin to increase towards present day 280 values. A lowering of Ω_{Ar} from approximately 4.0 to 3.0 is predicted to lead to significant reductions in 281 calcification rates in tropical reefs (Kleypas et al., 1999; Silverman et al., 2009). By the end of the 21st 282 century low latitude ocean Ω_{Ar} will drop below 3.0, well outside of the preindustrial values of $\Omega_{Ar} > 3.5$, 283 and high latitude ocean Ω_{Ar} is close to undersaturation (Ω <1) (Figure 6). These results agree with other 284 studies that investigated changes to the carbonate system(Roy et al., 2015; Ricke et al., 2013). Accounting for seasonal variations in the Ω_{Ar} saturation levels the time of undersaturation may move 285 286 forward by up to 17 ± 10 years (Sasse et al., 2015). Due to Hector's time step of 1 year, we may be

overestimating the time when ocean acidification reaches a critical threshold. We also note that other
 factors such as eutrophication, river discharge, and upwelling will likely increase the probability that
 coastal regions will experience the effects of ocean acidification sooner than the projected open ocean
 values in Hector (Ekstrom et al., 2015).

291 Using $[H^+]$ as a proxy for pH, we find that $[H^+]$ is sensitive to Q_{10} and ocean circulation. Changes 292 in Q₁₀, (the respiration temperature response) is responsible for the release of carbon on land. 293 Uncertainties in the land carbon cycle have been attributed to uncertainties in future CO₂ projections 294 within the CMIP5 models (Friedlingstein et al., 2014). Therefore, uncertainties in the land carbon cycle 295 will also have implications for the marine carbonate system. A 10% change in the thermohaline 296 circulation parameter (Tt in Figure 1), representing a portion of the high latitude to deep ocean 297 circulation, results in \sim 3% change in [H⁺]. The dynamics of ocean uptake of CO₂ are strongly dependent 298 on this circulation of CO₂ laden waters from the surface ocean to depth. CMIP5 models project a 299 weakening in the Atlantic meridional overturning circulation by an average of 36% under RCP8.5 by 2100 300 (Cheng et al., 2013). Therefore, changes in ocean circulation may have implications on the marine 301 carbonate system, influencing the ocean pH. We also find that the high latitude surface ocean is more 302 sensitive to parameter changes than the low latitude surface ocean. The high latitude box makes up 15% 303 of the global oceans in Hector and therefore, changes of the same size are more easily felt in the high 304 latitude box compared to the low latitude box. This may have direct implications on future marine 305 carbonate projections.

Global Ω_{Ar} saturation levels are most sensitive to changes in salinity. Salinity is used to calculate (Ca²⁺) and total boron (Appendix B). Typically the carbonate system is normalized to changes in salinity to understand the chemical changes within the system, instead we show that Ω_{Ar} may be sensitive not only to future changes in atmospheric [CO₂] but also sensitive to changes in precipitation and

- evaporation. This may be important, as studies suggest significant changes in precipitation patterns
 under a changing climate (Held and Soden, 2006; Liu and Allan, 2013).
- 312
- 313
- 314 **5. Conclusions**

We developed a simple, open-source, object oriented carbon cycle climate model, Hector, that reliably reproduces the median of the CMIP5 climate variables (Hartin et al., 2015). The ocean component presented in this study calculates the upper ocean carbonate system (pCO₂, DIC, pH, Ω_{Ar} , Ω_{Ca}). Under all four RCPs, pH and Ω_{Ar} decrease significantly outside of their preindustrial values matching both observations and CMIP5. In the near future the open ocean and coral reef communities are likely to experience pH and carbonate saturation levels unprecedented in the last 2 million years (Hönisch et al., 2009).

322 This study is timely because the CMIP5 archive includes a large suite of ESMs that contained 323 dynamic biogeochemistry, allowing us to study future projections of the marine carbon cycle. Rather 324 than running the ESMs, we can use Hector to quickly emulate the global CMIP5 median for projection 325 studies under different emission pathways and sensitivity analyses of the marine carbonate system. 326 Within this study we find that numerous parameters influence $[H^+]$ and Ω_{Ar} , with both being sensitive to 327 Q₁₀. Due to Hector's simplistic nature and fast execution times, Hector has the potential to be a critical 328 tool to the decision-making, scientific, and integrated assessment communities, allowing for further 329 understanding of future changes to the marine carbonate system.

330 Appendix A: Model Description – carbon cycle

The carbon component in Hector contains three carbon reservoirs: a single well-mixed atmosphere, a land component and an ocean component. The change in atmospheric carbon is a function of the anthropogenic emissions (F_A), land-use change emissions (F_{LC}), and atmosphere-ocean (F₀) and atmosphere-land (F_L) carbon fluxes. The default model time step is 1 year.

$$\frac{dC_{atm}(t)}{dt} = F_A(t) + F_{LC}(t) - F_O(t) - F_L(t)$$
(1)

The terrestrial cycle in Hector contains vegetation, detritus, and soil, all linked to each other and the atmosphere by first-order differential equations. Vegetation net primary production is a function of atmospheric CO₂ and temperature. Carbon flows from the vegetation to detritus to soil and loses fractions of carbon to heterotrophic respiration on the way. An 'earth' pool debits carbon emitted as anthropogenic emissions, allowing a continual mass-balance check across the entire carbon cycle. Atmosphere-land fluxes at time t are calculated by:

$$F_L(t) = \sum_{i=1}^n NPP_i(t) - RH_i(t)$$
⁽²⁾

where *NPP* is the net primary production and *RH* is the heterotrophic respiration summed over userspecified n groups (i.e., latitude bands, political units, or biomes) (Hartin et al 2015).

343

344 Appendix B: Ocean Carbonate Chemistry

The ocean's inorganic carbon system is solved via a series of equations modified from Zeebe and Wolf-Gladrow (2001). TA and DIC are used to calculate the other variables of the carbonate system:

$$DIC * \left(\frac{K_1}{[H^+]} + 2\frac{K_1K_2}{[H^+]^2}\right) = \left(TA - \frac{K_BB_T}{K_B + [H^+]} - \frac{K_W}{[H^+]} + [H^+]\right) * \left(1 + \frac{K_1}{[H^+]} + \frac{K_1K_2}{[H^+]^2}\right)$$
(1)

This equation results in a higher order polynomial equation for H^+ , in which the roots (1 positive, 4 negative) are solved for. Once H^+ is solved for, pH, pCO₂, HCO₃⁻, and CO₃²⁻ can be determined. We ignore the nonideality of CO₂ in air and therefore use the partial pressure of CO₂ instead of the fugacity
of CO₂. Fugacity is slightly lower by ~0.3% compared to pCO₂ (Riebesell et al., 2009; Sarmiento and
Gruber, 2006).

$$[CO_2^*] = \frac{DIC}{\left(1 + \frac{K_1}{[H^+]} + \frac{K_1K_2}{[H^+]^2}\right)}$$
(2)

$$pCO_2 = \frac{[CO_2^*]}{K_H} \tag{3}$$

$$[HCO_3^-] = \frac{DIC}{\left(1 + \frac{[H^+]}{K_1} + \frac{K_2}{[H^+]}\right)}$$
(4)

$$[CO_3^{2-}] = \frac{DIC}{\left(1 + \frac{[H^+]}{K_2} + \frac{[H^+]^2}{K_1 K_2}\right)}$$
(5)

$$K_1 = \frac{[H^+][HCO_3^-]}{[CO_2]} \tag{6}$$

$$K_2 = \frac{[H^+][CO_3^{2-}]}{[HCO_3^{-}]}$$
(7)

K₁ and K₂ are the first and second acidity constants of carbonic acid from Mehrbach et al. (1973) and
refit by Lueker et al. (2000).

$$K_B = \frac{[H^+][B(OH)_4^-]}{[B(OH)_3]}$$
(8)

354 K_B is the dissociation constant of boric acid from DOE (1994).

$$B = 416.0 * \frac{S}{35.0} \tag{9}$$

356 Total boron is from DOE (1994)

$$K_W = \frac{[H^+]}{[OH^-]}$$
(10)

357 K_w is the dissociation constant of water from Millero (1995).

$$K_{sp} = [Ca^{2+}] * [CO_3^{2-}]$$
(11)

358 K_{sp} of aragonite and calcite is calculated from Mucci, (1983).

359

- 360 For those equations with multiple coefficients:
- 1) K_H and K₀ are similar equations calculating Henry's constant or the solubility of CO₂, but they return
- 362 different units (mol kg⁻¹ atm⁻¹ and mol L⁻¹ atm⁻¹) (see Weiss, 1974 for equations and coefficients). K_H
- 363 is used to solve pCO_2 while K_0 is used to solve air-sea fluxes of CO_2 .
- 2) The Schmidt number is taken from Wanninkhof (1992) for coefficients of CO₂ in seawater.
- 365 3) $[Ca^{2+}]$ (mol kg⁻¹) is calculated from Riley and Tongudai (1967).
- 366

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372 Author Contributions

- 373 C.Hartin designed and carried out the experiments. C. Hartin, B.Bond-Lamberty, and P.Patel
- developed the model code. A.Mundra processed the data and prepared the figures. C.Hartin
- 375 prepared the manuscript with contributions from all co-authors.

Description	Value	Notes
*Albedo ^t	-0.2 Wm ⁻²	Constant global albedo from 1950 - 2300
Area of ocean	3.6e14 m ²	Knox and McElroy, (1984)
*Beta ^t	0.36	Terrestrial CO ₂ fertilization Wullschleger et al. (1995)
*Climate Sensitivity ^t	3.0 °C	
Fractional area of HL	0.15	Sarmiento and Toggweiler, (1984)
Fractional area of LL	0.85	Sarmiento and Toggweiler, (1984) Terrestrial respiration
${}^{*}Q_{10}{}^{t}$	2.0	temperature response Davidson and Janssens (2006)
Thickness of surface ocean	100 m	Knox and McElroy, (1984)
Thickness of intermediate ocean	900 m	()
Thickness of deep ocean	2677 m	Total ocean depth 3777m
Volume of HL	5.4e15 m ³	
Volume of LL	3.06e16 m ³	
Volume of IO	3.24e17 m ³	
Volume of DO	9.64e17 m ³	
Surface Area of HL	5.4e13 m ²	
Surface Area of LL	3.06e14 m ²	
Salinity HL and LL^{t}	34.5	
Initial temperature of HL ^t	2.0 °C	

Table 1: Description and values of ocean parameters in Hector and parameters involved in the

378

sensitivity analysis.

Initial temperature of LL ^t	22.0 °C	
[*] Thermohaline circulation $(T_T)^{t}$	7.2e7 m ³ s ⁻¹	Tuned to give ~100 Pg C flux from surface to deep
*High latitude circulation (T _H)	4.9e7 m ³ s ⁻¹	Tuned to give ~100 Pg C flux from surface to deep
*Water mass exchange (intermediate to deep - E _{ID})	1.25e7 m ³ s ⁻¹	Lenton (2000); Knox and McElroy (1984)
*Water mass exchange (low latitude to intermediate - E _{IL})	2.08e7 m ³ s ⁻¹	Lenton (2000); Knox and McElroy (1984)
Wind speed HL and LL ^t	6.7 m s ⁻¹	Takahashi et al. (2009); Liss and Merlivat, (1986)
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* Parameters contained within the input file.

^t Parameters varied for the sensitivity analysis.

Table 2: CMIP5 ESM models used in this study containing ocean carbonate parameters. Ω_{Ar} , Ω_{Ca} were382calculated from the model sea surface temperature, sea surface salinity, and CO₃ concentrations.

Model	Parameters (RCP 8.5)
BCC-cm1-1	pCO ₂ *, temperature
BNU-ESM	pCO ₂
CanESM2	DIC, pH, salinity
CESM1-BGC	CO₃, DIC, pH, salinity
CMCC-CESM	pCO ₂ , temperature, CO ₃ , DIC,
	pH, salinity
CNRM-CM5	CO ₃ , DIC
GFDL-ESM2G	pCO ₂ , temperature, pH, salinity
GFDL-ESM2M	pCO ₂ , temperature, CO ₃ , pH,
	DIC, salinity
GISS-E2-H-CC	pCO ₂ , temperature, DIC,
	salinity
GISS-E2-R-CC	pCO ₂ , temperature, DIC,
	salinity
HadGEM2-CC	pCO ₂ , temperature, CO ₃ , DIC,
	pH, salinity
HadGEM2-ES	pCO ₂ , temperature, CO ₃ *, DIC*,
	pH, salinity
IPSL-CM5A-LR	Temperature*, CO ₃ *, DIC*,
	pH*, salinity*
IPSL-CM5A-MR	Temperature, CO ₃ , DIC, pH,
	salinity
IPSL-CM5B-LR	Temperature, CO ₃ , DIC, pH,
	salinity
MIROC-ESM	pCO_2 , temperature, salinity
MIROC-ESM-	pCO ₂ , temperature, salinity
CHEM	
MPI-ESM-LR	pCO ₂ *, temperature*, CO ₃ *,
	DIC*, pH*, salinity*
MPI-ESM-MR	pCO ₂ , temperature, CO ₃ , DIC,
	pH, salinity
MIKI-ESM1	pCO ₂ , temperature
NorESM1-ME	pCO ₂ , temperature, CO ₃ , DIC,
	pH, salinity

383 * Variable output to 2300.

Time-	Location	Time-Series	Reference	Ocean Carbon	Data Access
Series Site		Length		Parameters	
BATS	Sargasso	1988-2011	Bates, (2007)	TA, DIC	http://www.bios.edu/research/projec
	Sea				<u>ts/bats</u>
HOT	North	1988-2011	Dore et al.	TA, DIC, pH,	http://hahana.soest.hawaii.edu/hot/h
	Pacific		(2009)	pCO ₂ , Ω _{Ar} , Ω _{Ca}	<u>ot_jgofs.html</u>
ESTOC	Canary	1995-2009	Gonalez-	TA, pH, pCO₂	http://www.eurosites.info/estoc.php
	Islands		Davila, (2009)		
Iceland Sea	Iceland	1985-2013	Olafsson,	DIC, pCO ₂	http://cdiac.ornl.gov/oceans/Mooring
	Sea		(2007a)		<u>s/Iceland_Sea.html</u>
Irminger	Irminger	1983-2013	Olafsson,	DIC, pCO ₂	http://cdiac.ornl.gov/oceans/Mooring
Sea	Sea		(2007b)		<u>s/Irminger_Sea.html</u>
Flinders	Coral Sea	1708-1988	Pelejero et al.	pH, Ω _{Ar}	<pre>ftp://ftp.ncdc.noaa.gov/pub/data/pal</pre>
Reef			(2005)		eo/coral/west_pacific/great_barrier/fl
					inders2005.txt

384 Table 3: Observational time-series information and carbonate parameters from each location.385

Table 4: Model validation metrics for the a) high latitude and b) low latitude ocean carbonate variables

388 comparing Hector to CMIP5 from 1850-2005.

a)	RMSE	R2	Bias
DIC	10.00	0.26	47.10
pCO ₂	2.65	0.98	-31.78
рН	0.004	0.975	0.061
Ω_{Ar}	0.01	0.98	0.37
Ω_{Ca}	0.02	0.98	0.58

b)	RMSE	R2	Bias
DIC	6.50	0.76	101.28
pCO₂	3.43	0.98	-4.62
рН	0.004	0.966	0.025
Ω_{Ar}	0.02	0.97	0.36
Ω_{Ca}	0.03	0.97	0.53

- **Table 5:** Absolute values and rates of change per year (Δ) for the a) high and b) low latitude surface
- 394 ocean from 1850-2100 and 2101-2300 under RCP 8.5 for DIC (µmol kg⁻¹), pCO₂ (µatm), pH (total scsale,
- 395 unitless), Ω_{Ar} (unitless) and Ω_{Ca} (unitless).

DIC				pCO2			рН			Ω_{Ar}			Ω_{Ca}		
a)	1850	2100	2300	1850	2100	2300	1850	2100	2300	1850	2100	2300	1850	2100	2300
Hector	2107.5	2258.1	2335.5	244.7	816.6	1732.1	8.23	7.76	7.46	2.2	1.0	0.6	3.5	1.5	0.9
Δ		0.602	0.387		2.29	4.58		-	-		-	-0.002		-0.008	-0.003
								0.0019	0.0015		0.0048				
CMIP5	2104.50	2175.79	2243.41	271.62	871.00	1903.82	8.17	7.70	7.38	1.82	0.75	0.44	2.90	1.20	0.70
Δ		0.285	0.34		2.40	5.16		-	-		-	-		-	-
								0.0019	0.0016		0.0012	0.0016		0.0068	0.0025

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Table 6: Trends and standard error for the carbonate system taken from Bates et al. (2014) and values

399 calculated from Hector and CMIP5.

	Length of record	DIC (µmol kg ⁻¹ yr ⁻¹)	pCO₂ (µatm yr⁻¹)	pH (yr ⁻¹)	Ω _{Ar} (yr ⁻¹)
BATS	1983-2014	1.37 ± 0.07	1.69 ± 0.11	-0.0017 ± 0.0001	-0.0095 ± 0.0007
НОТ	1988-2014	1.78 ± 0.12	1.72 ± 0.09	-0.0016 ± 0.0001	-0.0084 ± 0.0011
ESTOC	1995-2014	1.09 ± 0.10	1.92 ± 0.24	-0.0018 ± 0.0002	-0.0115 ± 0.0023
Iceland Sea	1983-2014	1.22 ± 0.27	1.29 ± 0.36	-0.0014 ± 0.0005	-0.0018 ± 0.0027
Irminger Sea	1983-2014	1.62 ± 0.35	2.37 ± 0.49	-0.0026 ± 0.0006	-0.0080 ± 0.0040
Hector	1988-2014	0.90	1.82	-0.0017	-0.0089
CMIP5	1988-2014	0.68	1.77	-0.0018	-0.0074

- **Table 7:** Percentage change from reference (RCP8.5) for two Hector outputs a) global [H⁺] and b) global404 Ω_{Ar} for a ±10% change in eight model parameters. Results are shown for three years, 2005, 2100 and4052300.

a) Year	Parameter	+10% change	-10% change
2005	Albedo	0.13	0.00
2100		0.59	0.00
2300		0.00	-0.31
2005	Beta (terrestrial CO ₂ fertilization)	0.63	-0.50
2100		1.78	-1.78
2300		1.56	-1.88
2005	Ocean Circulation (T _T)	0.76	-0.76
2100		2.37	-1.78
2300		2.81	-3.44
2005	Q ₁₀ (terrestrial respiration		
	temperature response)	-0.13	0.13
2100		-0.59	1.18
2300		-3.13	3.44
2005	Salinity	-0.88	1.51
2100		0.59	0.59
2300		1.88	-1.25
2005	Climate Sensitivity	-0.13	0.13
2100		-0.59	0.43
2300		-2.5	2.50
2005	Surface ocean temperature	0.00	0.00
2100		0.00	0.59
2300		0.62	0.31
2005	Wind Stress	-0.38	0.63
2100		-0.59	1.78
2300		-1.25	1.25

b) Year	Parameter	+10% change	-10% change
2005	Albedo	0.01	-0.00
2100		0.01	-0.01
2300		-0.00	0.00
2005	Beta (terrestrial CO ₂ fertilization)	0.38	-0.40
2100		1.33	-1.34
2300		1.38	-1.35
2005	Ocean Circulation (T _T)	0.41	-0.45
2100		1.01	-1.05
2300		1.48	-1.55
2005	Q ₁₀ (terrestrial respiration	-0.09	0.10
2100	temperature response)	-0.87	0.95
2300		-2.40	3.00
2005	Salinity	3.80	-4.28
2100		5.60	-5.89
2300		7.17	-7.18
2005	Climate Sensitivity	0.07	-0.07
2100		0.55	-0.56
2300		0.43	-0.27
2005	Surface ocean temperature	2.07	-1.99
2100		2.41	-2.29
2300		2.43	-2.27
2005	Wind Stress	-0.18	0.25
2100		-0.65	0.88
2300		-1.13	0.88

409 Figure 1: Representation of the carbon cycle in Hector. The atmosphere consists of one well-mixed box, 410 connected to the surface ocean via air-sea fluxes of carbon. The terrestrial component consists of user defined biomes or regions for vegetation, detritus, and soil. The earth pool is continually debited to act 411 412 as a mass balance check on the carbon cycle (Hartin et al., 2015). The ocean consists of four boxes, with 413 advection (represented by straight arrows) and water mass exchange (represented by curved arrows) 414 simulating thermohaline circulation. The marine carbonate system is solved for in the high and low 415 latitude surface boxes. At steady state, there is a flux of carbon from the atmosphere to the high 416 latitude surface box, while the low-latitude surface ocean releases carbon to the atmosphere.



- 419 **Figure 2:** Dissolved inorganic carbon (DIC) for high (top) and low latitude (bottom) surface ocean under
- 420 RCP 8.5; Hector (blue), CMIP5 median, standard deviation, and model range (red, *n* = 15 (1850-2100,
- 421 with n=4 starting after 1860) and *n*= 3 (2101-2300)); and observations from BATS (teal), ESTOC (pink),
- 422 HOT (purple), Iceland (yellow) and Irminger Sea (navy). Note a doubling of CO₂ from preindustrial values
- 423 occurs around 2050.
- 424



- 426 **Figure 3:** pCO₂ for high (top) and low latitude (bottom) surface ocean under RCP 8.5; Hector (blue),
- 427 CMIP5 median, standard deviation, and model range (red, n = 15 (1850-2100) and n = 2 (2101-2300));

428 and observations from BATS (teal), HOT (purple), ESTOC (pink), Iceland (yellow) and Irminger Sea (navy).



- 430 Figure 4: pH for high (top) and low latitude (bottom) surface ocean under RCP 8.5; Hector (blue), CMIP5
- 431 median, standard deviation, and model range (red, n = 13 (1850-2100) and n = 2 (2101-2300)); and
- 432 observations from BATS (teal), ESTOC (pink), HOT (purple) Flinder's Reef (green), Iceland (yellow) and
 433 Irminger Sea (navy).





Year

435 **Figure 5:** Aragonite saturation (Ω_{Ar}) for high (top) and low latitude (bottom) surface ocean under RCP 436 8.5; Hector (blue), CMIP5 median, standard deviation, and model range (red, n = 10 (1850-2100) and n =

437 2 (2101-2300)); and observations from BATS (teal), HOT (purple) and Flinder's Reef (green).

438

440 **Figure 6:** High and low latitude a) pH and b) aragonite saturation (Ω_{Ar}) time series for Hector from 1850-441 2300 for RCP 2.6 (red), RCP 4.5 (green), RCP 6.0 (teal) and RCP 8.5 (purple). Note that even under a

strongly mitigated scenario (RCP 2.6), both Ω_{Ar} and pH at 2300 are still lower than preindustrial values.



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