Reconciling Atmospheric and Oceanic Views of the Transient Climate Response to Emissions

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Abstract The Transient Climate Response to Emissions (TCRE), the ratio of surface warming and cumulative carbon emissions, is controlled by a product of thermal and carbon contributions. The carbon contribution involves the airborne fraction and the ratio of ocean saturated and atmospheric carbon inventories, with this ratio controlled by ocean carbonate chemistry. The evolution of the carbon contribution to the TCRE is illustrated in a hierarchy of models: a box model of the atmosphere-ocean and an Earth system model, both integrated for 1,000 years, and a suite of Earth system models integrated for 140 years. For all models, there is the same generic carbonate chemistry response: An acidifying ocean during emissions leads to a decrease in the ratio of the ocean saturated and atmospheric carbon inventories and the carbon contribution to the TCRE. Hence, ocean carbonate chemistry is important in controlling the magnitude of the TCRE and its evolution in time.

Plain Language Summary The increase in surface temperature with the amount of carbon emitted to the atmosphere depends on the uptake and storage of heat and carbon. Ocean heat uptake acts to strengthen surface warming, as the ocean becomes more stratified in time. Carbon uptake by the ocean and terrestrial system acts to weaken surface warming by removing carbon from the atmosphere. The proportionality of surface warming to carbon emissions may be written in terms of a thermal contribution multiplied by a carbon contribution. The carbon contribution depends on the increase in the atmospheric carbon inventory plus the maximum amount of carbon that the ocean may hold. To understand the role of ocean chemistry, we diagnose the response of climate models of differing complexity over centennial and millennial timescales. In all the models, there is a similar carbon response: During emissions, the ocean surface acidifies and the maximum amount of carbon that the ocean can hold decreases, which weakens the carbon contribution to the proportionality of surface warming to carbon emissions. Hence, ocean carbonate chemistry is important in controlling the proportionality of surface warming to carbon emissions and its evolution in time.

1. Introduction

Climate model projections reveal that surface global warming is nearly proportional to the cumulative carbon emission (Allen et al., 2009; Gillet et al., 2013; Matthews et al., 2009; Zickfeld et al., 2009). This relationship is encapsulated in the near constancy of an empirical climate metric, the Transient Climate Response to cumulative carbon Emissions (TCRE), which is defined by the ratio of the change in global mean, surface air temperature since the preindustrial era, ΔT(t), to the cumulative carbon emissions, Iem(t),

\[ \text{TCRE} = \frac{\Delta T(t)}{I_{\text{em}}(t)} \]  

(1)

The TCRE is usually considered for radiative forcing only from atmospheric CO₂, although this relationship may be generalized to include the effects of non-CO₂ radiative forcing (Williams et al., 2016). Two complementary views have been invoked to understand the TCRE. In the first view, emphasizing the atmospheric response, the TCRE is explained in terms of the airborne fraction, ΔIatm(t)/Iem(t), and the ratio of surface warming to changes in the atmospheric carbon inventory, ΔT(t)/ΔIatm(t) (Matthews et al., 2009), which may be reexpressed in terms of a thermal contribution, ΔT(t)/R(t), the airborne fraction, and the ratio of radiative forcing and changes in the atmospheric carbon inventory, R(t)/ΔIatm(t) (Ehlert & Zickfeld, 2017).
In this study, we reconcile these two different perspectives of the TCRE. Our approach is to focus on the ratio of the changes in the ocean saturated carbon inventory, defined by how much carbon the ocean may take up relative to the instantaneous atmospheric CO₂, and the atmospheric carbon inventory. A new expression for the TCRE is provided including both the airborne fraction and the ocean saturated carbon inventory, which highlights the effect of carbonate chemistry in controlling the ratio of changes in ocean saturated and atmospheric carbon inventories (section 2). This new TCRE expression is applied to understand the climate response during emissions and after emissions cease in a hierarchy of climate models: a box model of the atmosphere-ocean and a realistic Earth system model (GFDL-ESM2M) responding to a carbon emission over nearly 100 years and integrated for 1,000 years (Frölicher & Paynter, 2015; Williams, Roussenov, Frölicher, & Goodwin, 2017) and a suite of Earth system models integrated for a 1% annual rise in CO₂ for 140 years (section 3).

2. Theory

2.1. A Definition for the TCRE Including the Airborne Fraction and Ocean Saturation

To understand the link between the two formulations of TCRE in (2) and (3), consider the changes in the global carbon inventory, where cumulative carbon emissions, \( l_{em}(t) \), drive changes in the sum of the global carbon inventories for the atmosphere, \( \Delta l_{atm}(t) \), ocean, \( \Delta l_{ocean}(t) \), and terrestrial, \( \Delta l_{ter}(t) \), systems relative to the preindustrial era,

\[
l_{em}(t) = \Delta l_{atm}(t) + \Delta l_{ocean}(t) + \Delta l_{ter}(t). \tag{4}
\]

By connecting changes in the ocean carbon inventory, \( \Delta l_{ocean}(t) \), to changes in the ocean saturated and ocean undersaturated components (defined relative to the instantaneous atmospheric CO₂), \( \Delta l_{ocean}(t) = \Delta l_{sat}(t) - \Delta l_{sat}(t) \), then the carbon inventory changes in (4) may be rearranged as

\[
l_{em}(t) - \Delta l_{ter}(t) + \Delta l_{sat}(t) = \Delta l_{atm}(t) + \Delta l_{sat}(t). \tag{5}
\]

For the present day, the atmospheric CO₂ and the increase in the ocean carbon inventory, \( \Delta l_{ocean} \), are close to 400 ppm and 100 PgC, respectively, so that the increase in the saturated ocean carbon inventory, \( \Delta l_{sat} \), is about 900 PgC and in the ocean carbon undersaturation, \( \Delta l_{sat} \), is about 800 PgC.

Assuming that the changes in the radiative forcing, \( R(t) \), are driven only by changes in atmospheric CO₂, the radiative forcing is expressed in terms of changes in either the net carbon emission to the combined atmosphere and ocean, \( l_{em}(t) - \Delta l_{ter}(t) \), plus the ocean carbon undersaturation, \( \Delta l_{sat}(t) \) (Goodwin et al., 2015)
(see supporting information), or equivalently from (5), in the atmospheric, \( \Delta I_{atm}(t) \), plus the ocean saturated carbon, \( \Delta I_{sat}(t) \), inventories,

\[
R(t) = \frac{a}{I_B} (I_{em}(t) - \Delta I_{sat}(t)) \cdot \frac{\Delta I_{atm}(t) + \Delta I_{sat}(t)}{I_{em}(t)}.
\]

By substituting (6) into either (2) or (3), a new theoretical expression is derived for the TCRE,

\[
TCRE = \frac{\Delta T(t) a}{R(t) I_B} \left( \frac{\Delta I_{atm}(t) + \Delta I_{sat}(t)}{I_{em}(t)} \right) = \frac{\Delta T(t) a}{R(t) I_B} \left( \frac{\Delta I_{atm}(t)}{I_{em}(t)} \right) \left( 1 + \frac{\Delta I_{sat}(t)}{\Delta I_{atm}(t)} \right),
\]

where the ratio \( \Delta I_{sat}(t)/\Delta I_{atm}(t) \) represents the sensitivity of changes in the saturated ocean carbon inventory to changes in the atmospheric carbon inventory relative to the preindustrial era. This new expression for the TCRE including the airborne fraction (7) is equivalent to the expression for the TCRE including ocean undersaturation (3). Next, we explore how the ratio of the ocean saturated and atmospheric carbon inventories, \( \Delta I_{sat}(t)/\Delta I_{atm}(t) \), is controlled by ocean carbonate chemistry.

2.2. Control of the Ratio of the Ocean Saturated and Atmospheric Carbon Inventories

In order to understand the evolution of the ratio \( \Delta I_{sat}(t)/\Delta I_{atm}(t) \), consider an ocean buffer factor, \( B(t) \), where the fractional changes in the atmospheric and ocean saturated carbon inventories are defined relative to the preindustrial time, \( t_o \), by

\[
B(t) = \frac{\Delta I_{atm}(t)/I_{atm}(t_o)}{\Delta I_{sat}(t)/I_{sat}(t_o)}.
\]

Rearranging (8), and drawing upon basic carbonate chemistry (see supporting information), the ratio of the changes in the saturated ocean and atmospheric inventories, \( \Delta I_{sat(t)}/\Delta I_{atm}(t) \), may be approximated by

\[
\frac{\Delta I_{sat}(t)}{\Delta I_{atm}(t)} \approx \left( \frac{\rho_o V K_o}{M_o} \right) \left( \frac{K_1 K_2}{[H^+(z_{s}, t)]^2} \right) \left( \frac{CO_2(t)}{CO_2(t_o)} \right),
\]

where \( M_o \) is the moles of gas in the atmosphere, \( \rho_o \) is a referenced ocean density, \( V \) is the ocean volume, and the hydrogen ion concentration, \( H^+ \), is evaluated at the sea surface, \( z_s \). The changes in carbonate chemistry due to temperature are assumed relatively small compared with those changes due to atmospheric \( CO_2 \), and so the effect of changes in the solubility, \( K_o \), and the equilibrium coefficients, \( K_1 \) and \( K_2 \), are neglected in (9).

In this approximation for the ratio \( \Delta I_{sat(t)}/\Delta I_{atm}(t) \) in (9), each term has the following interpretation: The first term on the right-hand side represents the value that this ratio would have if \( CO_2 \) was nonreactive in seawater; the second term is inversely associated with the abundance of \( H^+ \) ions in the ocean and decreases due to the increase in \( H^+ \) ions when there is an addition of \( CO_2 \) to seawater; and the third term is associated with the increase in the atmospheric \( CO_2 \) relative to its preindustrial value. This approximation for \( \Delta I_{sat}(t)/\Delta I_{atm}(t) \) is next used to understand the carbon control of the TCRE in a range of climate models (for the validity of this approximation; see the supporting information).

3. Model Assessment of the TCRE Exploiting our Theory and Approximation

Our theory is used to interpret the control of the TCRE in a hierarchy of models forced by atmospheric \( CO_2 \) changes: a box model of the atmosphere-ocean system and an Earth system model, both integrated for 1,000 years, and a suite of Earth system models, integrated for 140 years.

3.1. Experimental Design

The box model consists of three homogeneous layers: a well-mixed atmosphere, an ocean mixed layer with 100-m thickness, and an ocean interior with 3,900-m thickness, all assumed to have the same horizontal area. The model solves for the heat and carbon exchange between these layers, including physical and chemical transfers, but ignoring biological transfers, and sediment and weathering interactions. The model is forced from an equilibrium by carbon emitted into the atmosphere with a constant rate of 20 PgC/year for 100 years and integrated for 1,000 years. Ocean ventilation is represented by the ocean interior taking up the heat and carbon properties of the mixed layer on an e-folding timescale of 200 years (see supporting information for the model closures). The radiative forcing, \( R(t) \), varies according to the increase in atmospheric \( CO_2 \): \( R(t) = a \Delta \ln CO_2(t) \) (Myhre et al., 1998), where \( a = 5.35 \text{ W/m}^2 \) and the preindustrial \( CO_2(t_o) \) is 280 ppm. This radiative forcing, \( R(t) \), is assumed to drive a global-mean radiative response, \( \lambda(t) \Delta T(t) \), plus a planetary heat uptake,
The response of an Earth system model is diagnosed over a 1,000-years integration using the global coupled, carbon-climate model developed at Geophysical Fluid Dynamics Laboratory (GFDL-ESM2M; Dunne et al., 2012, 2013). The idealized warming simulation experiment of Frölicher and Paynter (2015) is examined: forced by an annual 1% rise of atmospheric CO2 until the global-mean surface warming reaches 2 ºC at year 98 of the simulation and then the carbon emissions are set to 0. The non-CO2 greenhouse gases are kept at their preindustrial levels.

The ocean saturated carbon inventory is estimated as $I_{ocean}(t) = \rho_a V_{DIC}(t)$, where $V_{DIC}$ is the ocean volume-weighted dissolved inorganic carbon in mol C·kg$^{-1}$. The atmosphere and ocean buffered carbon inventory is defined by $I_o = I_{atm}(t_o) + I_{sat}(t_o)/B_{rev}(t_o)$ (Goodwin et al., 2007), where $B_{rev}$ is the Revelle buffer factor (Williams & Follows, 2011). The carbon inventories are further multiplied by 12 g/mol to be expressed in grams of carbon. The partitioning of dissolved inorganic carbon into bicarbonate, carbonate, and dissolved CO2 is solved for using the algorithm of Follows et al. (2006).

The ocean saturated carbon inventory is estimated as $I_{sat}(t) = \rho_a V_{DIC_{sat}}(t)$, where the saturated dissolved inorganic carbon, $V_{DIC_{sat}}$, is diagnosed using the ocean potential temperature, salinity, alkalinity, and the atmospheric CO2 inventory (Ito & Follows, 2005; Lauderdale et al., 2013). The saturated carbon inventory represents the amount of carbon the ocean would have if the ocean reached a chemical equilibrium with the instantaneous atmospheric CO2. The surface ocean equilibrates on an annual timescale with the atmosphere, so the surface dissolved CO2 only slightly lags atmospheric CO2. However, much of the ocean interior has not been in contact with the atmosphere since the preindustrial era when the atmospheric CO2 was 280 ppm. At present, atmospheric CO2 is 400 ppm, which corresponds to an increase in the ocean saturated carbon inventory relative to the preindustrial era, $\Delta I_{sat}$, of about 900 Pg C. In comparison, the increase in the ocean carbon inventory relative to the preindustrial era, $\Delta I_{ocean}$, is about 100 Pg C so that a further 800 Pg C is needed for the ocean to become saturated to the present atmospheric CO2.

The planetary heat uptake, $N(t)$, is dominated by the ocean heat uptake (Church et al., 2011). In the box model, more than 95% of heat passes into the ocean; henceforth, we refer to the planetary and the ocean heat uptakes as being effectively equivalent. In the box model, the climate feedback parameter, $\lambda$, is assumed to be constant and equal to 1 W·m$^{-2}$·K$^{-1}$. In the Earth system models, the climate feedback parameter is more realistically taken to be time dependent, $\lambda(t)$ (Armour et al., 2013; Gregory & Andrews, 2016; Senior & Mitchell, 2000); $\lambda(t)$ is diagnosed from the radiative forcing, $R(t)$, the surface warming, $\Delta T(t)$, and the ocean heat uptake, $N(t)$, using the empirical heat budget, $\dot{R}(t) = \lambda(t) \Delta T(t) + N(t)$ (see the supporting information). The equilibrium climate feedback parameter, $\lambda_{eq}$, is diagnosed at the end of the 1,000 years model run in the GFDL-ESM2M model. For other Earth system models without long integrations, $\lambda_{eq}$ is taken from Forster et al. (2013; Table 1 for net feedbacks).

In our diagnostics of the Earth system models, a 5-year filter is applied to remove higher-frequency variability in the ocean heat uptake and surface warming, and diagnostics are not shown for the first 10 years when the cumulative emissions are still small and the climate response is controlled by internal variability.

3.3. Analysis and Model Responses

Our aim is to understand the climate response to carbon emissions in terms of the carbon and thermal contributions to the TCRE. The box model and the GFDL-ESM2M Earth system model, integrated over 1,000 years, have broadly similar carbon and thermal responses, despite their different model complexity (Figures 1 and 3). The small differences between their model responses are associated with the more complex Earth system
Figure 1. Carbon response for the box model (left panels) and the GFDL-ESM2M Earth system model (right panels) integrated for 1,000 years: (a, b) cumulative carbon emissions, \( I_{\text{em}}(t) \), along with the carbon inventory changes relative to the preindustrial era for the atmosphere, \( \Delta I_{\text{atm}}(t) \), the ocean, \( \Delta I_{\text{ocean}}(t) \), the terrestrial system, \( \Delta I_{\text{ter}}(t) \), and the ocean saturated carbon, \( \Delta I_{\text{sat}}(t) \); (c, d) the airborne fraction, \( \Delta I_{\text{atm}}(t)/I_{\text{em}}(t) \), the oceanborne fraction for saturated carbon, \( \Delta I_{\text{sat}}(t)/I_{\text{em}}(t) \), and their sum, \( \Delta I_{\text{atm}}(t) + \Delta I_{\text{sat}}(t)/I_{\text{em}}(t) \); and (e, f) the ratio of the changes in the saturated ocean and atmospheric carbon inventories, \( \Delta I_{\text{sat}}(t)/\Delta I_{\text{atm}}(t) \), along with the three terms that control this ratio from (9): \( \rho_0 V K / M_\alpha \), \( \Delta \text{CO}_2(t)/\Delta \text{CO}_2(t) \), and \( K_1 K_2 [H^+ (z_\alpha, t)]^2 \). The thin black dotted line notes the cessation of the emissions.

GFDL = Geophysical Fluid Dynamics Laboratory.

model having a time-varying climate feedback parameter, terrestrial carbon uptake, ocean circulation changes, and its carbon cycle including weathering and sediment interactions, and the effects of ocean biology. The suite of Earth system models reveal some intermodel variability, although their model responses are broadly similar to each other (Figures 2 and 3). Hence, the essential controls of the carbon and thermal contributions to the TCRE are similar in this range of climate models despite differences in model complexity. The details of the climate response are now worked through.
Figure 2. Carbon response for the suite of Earth system models integrated for 140 years: (a) the airborne fraction, \( \Delta I_{\text{atm}}(t)/I_{\text{em}}(t) \); (b) the oceanborne fraction for saturated carbon, \( \Delta I_{\text{sat}}(t)/I_{\text{em}}(t) \); and (c) the ratio of the changes in the saturated ocean and atmospheric carbon inventories, \( \Delta I_{\text{sat}}(t)/\Delta I_{\text{atm}}(t) \).

### 3.3.1. Carbon Contribution

The carbon contribution to the TCRE from (6), \( R(t)/I_{\text{em}}(t) \), is defined in terms of changes in the airborne fraction, \( \Delta I_{\text{atm}}(t)/I_{\text{em}}(t) \), and the saturated oceanborne fraction, \( \Delta I_{\text{sat}}(t)/I_{\text{em}}(t) \),

\[
R(t) = \frac{a}{I_{\text{em}}(t)} \left( \frac{\Delta I_{\text{atm}}(t)}{I_{\text{em}}(t)} + \frac{\Delta I_{\text{sat}}(t)}{I_{\text{em}}(t)} \right) = \frac{a}{I_{\text{em}}(t)} \frac{\Delta I_{\text{atm}}(t)}{I_{\text{em}}(t)} \left( 1 + \frac{\Delta I_{\text{sat}}(t)}{\Delta I_{\text{atm}}(t)} \right). \tag{10}
\]

In all the models, during emissions, there is an increase in the atmospheric carbon inventory, \( \Delta I_{\text{atm}}(t) \), and in the ocean carbon inventory, \( \Delta I_{\text{sat}}(t) \), as some of the emitted carbon is transferred into the ocean (red and blue lines in Figures 1a and 1b). The rise in atmospheric CO\(_2\) leads to an increase in the saturated ocean carbon inventory, \( \Delta I_{\text{sat}}(t) \) (cyan lines in Figures 1a and 1b). After emissions cease, there is a decrease
Figure 3. Transient climate response to emissions (TCRE) for the box model (left panels) and the GFDL-ESM2M Earth system model (middle panels), integrated for 1,000 years, and the suite of Earth system models (right panels), integrated for 140 years: (a–c) the nondimensional, thermal contribution, \( (\lambda_{\text{eq}}/\lambda(t)(1-N(t)/R(t))) \), and carbon contribution, \( (\Delta I_{\text{atm}}(t)/\Delta I_{\text{em}}(t))(1 + \Delta I_{\text{sat}}(t)/\Delta I_{\text{atm}}(t)) \), to the TCRE; (d, e) the TCRE diagnosed from (1) (black lines), using our theory from (12) (blue lines) and using the carbonate approximation from (13) (red lines) for the box and the GFDL-ESM2M models; and (f) the TCRE diagnosed from (1) (solid lines), for the suite of Earth system models. In addition, estimates of the TCRE by artificially excluding the effect of the carbonate chemistry in (13) are denoted in (d)–(f) by dashed lines. The thin black dotted line notes the cessation of the emissions. GFDL = Geophysical Fluid Dynamics Laboratory.

in the atmospheric inventory and the saturated ocean carbon inventory, accompanied by a further increase in the ocean carbon inventory.

These carbon inventory changes are often understood in terms of the airborne, oceanborne, and landborne fractions of carbon, where \( \Delta I_{\text{atm}}(t)/\Delta I_{\text{em}}(t) + \Delta I_{\text{ocean}}(t)/\Delta I_{\text{em}}(t) + \Delta I_{\text{ter}}(t)/\Delta I_{\text{em}}(t) = 1 \) (Jones et al., 2013). The airborne fraction, \( \Delta I_{\text{atm}}(t)/\Delta I_{\text{em}}(t) \), generally decreases as carbon is transferred from the atmosphere into the ocean until an equilibrium is approached in the box and the GFDL-ESM2M models (red lines in Figures 1c and 1d). The Earth system models experience large intermodel variability in terms of the magnitude of the airborne fraction (Figure 2a), with smaller airborne fraction associated with larger terrestrial carbon uptake. The trend in the airborne fraction, \( \Delta I_{\text{atm}}(t)/\Delta I_{\text{em}}(t) \), is though similar in all the Earth system models: There is a rapid decrease in the airborne fraction over the first 50 years, followed by a slight increase, primarily associated with a slight decrease in the landborne fraction.

In all the models, there is also a decrease in the saturated oceanborne fraction, defined by \( \Delta I_{\text{sat}}(t)/\Delta I_{\text{em}}(t) \) (cyan lines in Figures 1c and 1d and colored lines in Figure 2b). Hence, there is a decrease in the sum of the airborne and saturated oceanborne fractions, \( \Delta I_{\text{atm}}(t) + \Delta I_{\text{sat}}(t)/\Delta I_{\text{em}}(t) \) (black lines/shading in Figures 1c, 1d, and 3a–3c), which is proportional to the carbon contribution, \( R(t)/\Delta I_{\text{em}}(t) \), from (10). The rate of decrease in the saturated oceanborne fraction, \( \Delta I_{\text{atm}}(t)/\Delta I_{\text{em}}(t) \), is different to that of the airborne fraction, \( \Delta I_{\text{atm}}(t)/\Delta I_{\text{em}}(t) \), due to the effect of the carbonate chemistry.

The ratio of the changes in the saturated ocean and atmospheric carbon inventories, \( \Delta I_{\text{sat}}(t)/\Delta I_{\text{atm}}(t) \), is controlled by the carbonate chemistry: \( \Delta I_{\text{sat}}(t)/\Delta I_{\text{atm}}(t) \) is proportional to \( (K_{\text{i}}K_{\text{j}}/\overline{H'_{z}}t)\overline{(CO_{2}/CO_{2})(t_{0})} \).
from (9). The magnitude and time variability of the ratio $\Delta I_{sat}(t)/\Delta I_{em}(t)$ is similar in all the models (black lines in Figures 1e and 1f and colored lines in Figure 2c). During emissions, the ratio $\Delta I_{sat}(t)/\Delta I_{em}(t)$ decreases (black lines in Figures 1e and 1f) from the increase in $H^+$ ions (red dashed lines in Figures 1e and 1f) despite the accompanying increase in atmospheric CO$_2$ (red dashed-dotted lines in Figures 1e and 1f). After emissions cease, the ratio $\Delta I_{sat}(t)/\Delta I_{em}(t)$ increases due to the decrease in $H^+$ ions, dominating over the effect of the decrease in atmospheric CO$_2$. In comparison, if the effect of carbonate chemistry is artificially excluded, the ratio $\Delta I_{sat}(t)/\Delta I_{em}(t)$ is constant and equal to $\rho_s VK_s/M_s$ (red dotted lines in Figures 1e and 1f).

### 3.3.2. Thermal Contribution

The thermal contribution to the TCRE, $\Delta T(t)/R(t)$, is understood via the empirical heat budget (Gregory & Forster, 2008; Gregory et al., 2004), where radiative forcing drives a radiative response and a heat uptake, $R(t) = \lambda(t) \Delta T(t) + N(t)$, which may be reexpressed as

$$\frac{\Delta T(t)}{R(t)} = \frac{1}{\lambda_{eq}} \left[ \frac{\lambda_{eq}}{\lambda(t)} \left( 1 - \frac{N(t)}{R(t)} \right) \right],$$

where the term in the square brackets is the realized-warming fraction (Frölicher & Paynter, 2015; Frölicher et al., 2014; Solomon et al., 2009; red lines/shading in Figures 3a–3c).

In the box model, the realized-warming fraction, $(\lambda_{eq}/\lambda(t))(1 - N(t)/R(t))$, is driven by the fraction of the radiative forcing that warms the surface versus the fraction used to increase ocean heat content, $N(t)/R(t)$. In the Earth system models, the time variation of $\lambda(t)$ also affects the realized-warming fraction. Initially, in all the models, most of the radiative forcing is used to drive ocean heat uptake and warming of the ocean interior. As the ocean interior becomes warmer, gradually a smaller fraction of the radiative forcing warms the ocean interior and the realized-warming fraction increases (red lines/shading in Figures 3a–3c). The large decline of the realized-warming fraction in the GFDL-ESM2M model during the first 20 years is associated with an initial large increase in $\lambda(t)$.

### 3.3.3. Transient Climate Response to Carbon Emissions

The TCRE may be defined in terms of a product of thermal and carbon contributions by combining (7) and (11),

$$TCRE = \frac{a}{\lambda_{eq}} \left[ \frac{\lambda_{eq}}{\lambda(t)} \left( 1 - \frac{N(t)}{R(t)} \right) \right] \frac{\Delta I_{sat}(t)}{I_{em}(t)} \left[ 1 + \frac{\Delta I_{sat}(t)}{\Delta I_{em}(t)} \right],$$

where the time-independent term, $a/\lambda_{eq}$, is the long-term equilibrium climate response to emissions (Williams et al., 2012), and the terms in the square brackets represent the nondimensional, time-dependent thermal and carbon contributions to the TCRE.

Our theoretical relationship for the TCRE diagnosed from (12) agrees well with the actual TCRE diagnosed from (1) for both the box and the GFDL-ESM2M models integrated for 1,000 years (compare the black and blue lines in Figures 3d and 3e); this agreement also holds for the other Earth system models integrated for 140 years (see the supporting information).

The TCRE is viewed as a product of the thermal and the carbon contributions. The carbon contribution follows the decrease in $(\Delta I_{em}(t)/I_{em}(t))(1 + \Delta I_{sat}(t)/\Delta I_{em}(t))$, while the thermal contribution follows the increase in the realized-warming fraction, $(\lambda_{eq}/\lambda(t))(1 - N(t)/R(t))$ (Figures 3a–3c). Thus, the thermal contribution acts to increase the TCRE, while the carbon contribution acts to decrease the TCRE. The TCRE is constant when changes in these thermal and carbon contributions compensate for each other. However, in all the models, during emissions, there is a general decrease in the TCRE after about the first 30 years, with the rate of the decrease being model dependent (Figures 3d–3f). After emissions cease, there is an increase in the TCRE in the box model and the GFDL-ESM2M (Figures 3d and 3e).

To gain further insight, now combine (9) and (12) to obtain a TCRE expression based on our carbonate approximation,

$$TCRE \approx \frac{a}{\lambda_{eq}} \left[ \frac{\lambda_{eq}}{\lambda(t)} \left( 1 - \frac{N(t)}{R(t)} \right) \right] \frac{\Delta I_{sat}(t)}{I_{em}(t)} \left[ 1 + \frac{\rho_s VK_s}{M_s} \frac{K_1 K_2}{[H^+(z_s, t)]^2} \frac{CO_2(t)}{CO_2(t_0)} \right].$$

This carbonate approximation for the TCRE underestimates the TCRE in the box model and overestimates the TCRE in the Earth system models but captures its variability in time in all the models (compare the black and red lines in Figures 3d and 3e, for the suite of Earth system models see the supporting information).
If the effect of carbonate chemistry is artificially excluded in (13), the contribution to the TCRE from \((a/\lambda(t)I_2)(1 - N(t)/R(t))(\Delta I_{\text{atm}}(t)/I_{\text{em}}(t))(1 + \rho_0 VK_a/M_o)\) is nearly constant (dashed lines in Figures 3d–3f); in the Earth system models, there is a slight increase in the TCRE without the explicit effect of the carbonate chemistry after about the first 20 years from the temporal evolution of \(\lambda(t)\), changes in ocean circulation, and changes in terrestrial carbon uptake. If the effect of carbonate chemistry is now included in (13), the term involving carbonate chemistry, \((K_{\text{atm}}/H^+(z_0, t)^2)(\text{CO}_2(t)/\text{CO}_2(t_0))\), increases the magnitude of the TCRE, and modulates the carbon contribution to the TCRE in a consistent manner in all the models. Hence, the carbonate chemistry leads to a long-term decrease in the TCRE during emissions and an increase in the TCRE after emissions cease (Figures 3d–3f).

4. Conclusions

The TCRE is a fundamental climate metric, measuring the surface warming increase for a cumulative carbon emission (Gillet et al., 2013; Matthews et al., 2009). While the TCRE may simply be diagnosed from output of climate model projections, theory may be exploited to understand how the TCRE is controlled in terms of physical and biogeochemical mechanisms (Goodwin et al., 2015; MacDougall & Friedlingstein, 2015; Williams et al., 2016; Williams, Roussenov, Frölicher, & Goodwin, 2017; Williams, Roussenov, Goodwin, et al., 2017). Our new expression for the TCRE involves a product of two time-varying thermal and carbon contributions: The thermal contribution involves ocean heat uptake and the climate feedback parameter, and the carbon contribution involves the airborne fraction (depending on ocean and terrestrial carbon uptake) and the ratio of ocean saturated and atmospheric carbon inventories (depending on ocean carbonate chemistry).

Our viewpoint of how the TCRE is controlled is assessed using a hierarchy of climate models: a box model of the atmosphere-ocean and an Earth system model, both diagnosed for integrations for 1,000 years, and a suite of Earth system models diagnosed over 140 years. For all models, there is the same ocean carbonate response: During emissions, an increase in acidity at the ocean surface leads to a decrease in the ratio of the ocean saturated and atmospheric carbon inventories, which decreases the carbon contribution to the TCRE; and after emissions cease, a decrease in acidity at the ocean surface leads to an increase in this ratio and an increase in the carbon contribution to the TCRE.

The TCRE need not remain constant in time (Krasting et al., 2014) due to changes in the time-dependent thermal and carbon contributions. While the ocean ventilated heat and carbon uptake may act in a nearly compensating manner (Solomon et al., 2009), the carbonate chemistry leads to a systematic decrease in the TCRE with increasing acidity. If the effect of the carbonate chemistry is artificially excluded in the climate model, the TCRE is smaller in magnitude and is nearly constant in time due to the compensating effects of ocean heat and carbon uptake from ventilation.

In summary, ocean carbonate chemistry is important in controlling the magnitude of the TCRE and contributing to its temporal evolution. Reassuringly, in a range of climate models with differing complexity, carbonate chemistry provides a similar control of the carbon contribution to the TCRE. Intermodel differences in the TCRE are more likely due to other factors (Williams, Roussenov, Goodwin, et al., 2017), such as the thermal contribution to the TCRE, involving the climate feedback parameter and ocean ventilation of heat, or the separate carbon contribution to the TCRE from the airborne fraction, altered by ocean ventilation of carbon and terrestrial cycling of carbon.

References


