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Deep time evidence for climate sensitivity increase with warming

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Abstract Future global warming from anthropogenic greenhouse gas emissions will depend on climate feedbacks, the effect of which is expressed by climate sensitivity, the warming for a doubling of atmospheric CO₂ content. It is not clear how feedbacks, sensitivity, and temperature will evolve in our warming world, but past warming events may provide insight. Here we employ paleo-reconstructions and new climate-carbon model simulations in a novel framework to explore a wide scenario range for the Paleocene-Eocene Thermal Maximum (PETM) carbon release and global warming event 55.8 Ma ago, a possible future warming analogue. We obtain constrained estimates of CO₂ and climate sensitivity before and during the PETM and of the PETM carbon input amount and nature. Sensitivity increased from 3.3–5.6 to 3.7–6.5 K (Kelvin) into the PETM. When taken together with Last Glacial Maximum and modern estimates, this result indicates climate sensitivity increase with global warming.

1. Introduction

The quantification of climate sensitivity (CS) is arguably the major challenge of climate science. Present CS estimates are 1.5–4.5 K, a range mostly expressing variations in climate system feedbacks among coupled climate models, whereby cloud feedbacks are particularly uncertain [Flato et al., 2013; Bony et al., 2015; Sherwood et al., 2014]. Furthermore, climate system feedbacks may be climate dependent [Caballero and Huber, 2013]. How will such feedbacks and therewith climate sensitivity respond to ongoing global warming from anthropogenic greenhouse gas emissions?

There have been several recent attempts to constrain CS using paleo-reconstructions for different periods in the past, but the most robust of these from a data availability perspective are for cold glacial conditions [e.g., Schmittner et al., 2011]. Across the Paleocene-Eocene Thermal Maximum (PETM), the Earth experienced rapid warming of about 5°C from a mean global temperature already about 10°C warmer than present day [Dunkley Jones et al., 2013; Lunt et al., 2012]. This global warming event was driven by carbon emissions to the atmosphere leading to increased greenhouse gas forcing [Pagani et al., 2006; Zachos et al., 2008]. The PETM has been the focus of much study during the past 20 years, in part motivated by its potential relevance to ongoing warming [e.g., Dickens et al., 1995; Zachos et al., 2008]. But uncertainty remains with regard to the source and amount of the PETM carbon input and the climate response to this input [Pagani et al., 2006; Zachos et al., 2005; Zeebe et al., 2009]. Climate-carbon modeling studies with model-dependent CS values and a priori ocean carbon inventory choices and constrained by specific paleo-reconstructions have yielded different and even mutually exclusive results [Panchuk et al., 2008; Zeebe et al., 2009; Cui et al., 2011; Meissner et al., 2014]. Here we introduce a new framework for synthesizing the large and diverse body of paleo-reconstructions from the PETM and adjacent Eocene times and carry out comprehensive climate-carbon simulations using a relatively simple but well-tested and flexible Earth System Model [Shaffer et al., 2008; Eby et al., 2013]. Our results provide constrained estimates for CS dependence on climate in a warming world and shed new light upon the PETM itself.

2. Climate Sensitivity and Atmospheric CO₂

PETM ocean surface warming was about 5°C with little polar amplification [Dunkley Jones et al., 2013]. Model results indicate that mean atmospheric warming probably was slightly larger [Lunt et al., 2012].
A small fraction of this rise might be ascribed to higher concentrations of minor greenhouse gases; however, most of the warming must be explained by greater $p\text{CO}_2$ from a large carbon input to the ocean-atmosphere system, as evidenced by the PETM carbon isotope excursion (CIE) [Dunkley Jones et al., 2013; Zachos et al., 2008; Pagani et al., 2006; McInerney and Wing, 2011]. Much of this input may have been as methane that is oxidized to CO$_2$ over decadal timescales whereby enhanced $p\text{CO}_2$ radiative forcing dwarfs that from enhanced methane for the input times considered here (Text S1 in the supporting information).

We can relate PETM $p\text{CO}_2$ excursion ($\Delta p\text{CO}_2\text{(PETM)}$), pre-PETM $p\text{CO}_2$ ($p\text{CO}_2\text{(pre-PETM)}$), and PETM climate sensitivity ($C\text{S}_{\text{PETM}}$) by

$$C\text{S}_{\text{PETM}} = \Delta T\text{PETM} \ln(2) / \ln \left( \frac{p\text{CO}_2\text{(pre-PETM)} + \Delta p\text{CO}_2\text{(PETM)}}{p\text{CO}_2\text{(pre-PETM)}} \right)$$

(1)

where $C\text{S}$ refers to temperature increase after ocean heat equilibration and $\Delta T\text{PETM}$ is the mean atmospheric temperature (MAT) rise over the PETM from $p\text{CO}_2$ increase, taken here to be 5°C. An analogous expression for pre-PETM climate sensitivity ($C\text{S}_{\text{pre-PETM}}$) is

$$C\text{S}_{\text{pre-PETM}} = \Delta T\text{pre-PETM} \ln(2) / \ln \left( \frac{p\text{CO}_2\text{(pre-PETM)} + \Delta p\text{CO}_2\text{(pre-PETM)}}{p\text{CO}_2\text{(P$I}$} \right)$$

(2)

where $\Delta T\text{pre-PETM}$ is the part of the MAT difference between pre-PETM and preindustrial (PI) times due to higher $p\text{CO}_2$ and $p\text{CO}_2\text{(P$I)}$ is 280 ppm. Pre-PETM conditions can be estimated by comparing late Paleocene deep ocean and surface temperature records and considered within the context of subsequent, nonhyperthermal, early Eocene records for which more data exist. From these, we can glean that pre-PETM MAT was ~10°C warmer than PI [Huber and Caballero, 2011; Lunt et al., 2012; Zachos et al., 2008]. Much of this extra warmth may derive from albedo reduction from much reduced high-latitude ice and snow, complemented by low-latitude, albedo reductions from reduced subtropical landmass [Herold et al., 2014] (Figure S1) and increased land vegetation cover [Huber and Caballero, 2011]. Aerosol decrease and higher minor greenhouse gas concentrations may have each enhanced warming by ~1°C [Lunt et al., 2012]. From model results that include these factors [Huber and Caballero, 2011; Lunt et al., 2012], we estimate that they may account for about 4°C of the warming leaving about 6°C to be explained by greater atmospheric $p\text{CO}_2$. Thus, we take $\Delta T\text{pre-PETM}$ to be 6°C.

From equations (1) and (2) with the above choices for MAT increases due to $p\text{CO}_2$ we can deduce possible ranges for pre-PETM and PETM $C\text{S}$ from estimates of pre-PETM $p\text{CO}_2$ and the PETM $p\text{CO}_2$ excursion. Here we use proxy data and carbon cycle modeling to obtain such estimates and plot the results in a space defined by pre-PETM $p\text{CO}_2$ and the PETM $p\text{CO}_2$ excursion (Figure 1). In this way we constrain a possible area in this $p\text{CO}_2$ space and thereby also possible ranges for pre-PETM and PETM $C\text{S}$.

### 3. Mineralogical, Carbon Cycle, and Isotope Constraints

#### 3.1. Nahcolite

One constraint derives from the existence of the mineral nahcolite in the Green River Formation, USA, from the Early Eocene Climatic Optimum (EECO) about 51 Ma ago and from stability fields of sodium carbonate
forms (nahcolite, trona, and natron) as functions of temperature and \( pCO_2 \) [Lowenstein and Demicco, 2006; Jagniecki et al., 2015]. While the original work derived a lower bound on EECO \( pCO_2 \) of about 1125 ppm from the triple point where these forms are at equilibrium (in the presence of halite), the recent analysis based on new experimental data puts this point at 680 ppm and a temperature of 19.5°C. Furthermore, homogenization temperatures for fluid inclusions in halite in this formation were found to be 21 to 28°C. Together with the new stability fields, this translates to minimum and maximum estimates for EECO \( pCO_2 \) of 760 and 1260 ppm [Jagniecki et al., 2015]. Below we show how these results can constrain pre-PETM \( pCO_2 \) and the PETM \( pCO_2 \) excursion.

Pre-PETM temperatures were about 2°C cooler than during the EECO [Zachos et al., 2008], likely due to less greenhouse gas forcing given similarity between pre-PETM and EECO boundary conditions. With this and the results above, we can estimate bounds on EECO CS using

\[
C_{SECO} = \Delta T_{EECO} \ln(2) / \ln(pCO_2(EECO)/pCO_2(EECO))
\]

with \( \Delta T_{EECO} = -2°C \). Since the EECO was warmer than pre-PETM but cooler than the PETM, it is reasonable to assume that \( C_{SECO} \) lies between CS for these periods. If \( CS_{PETM} > CS_{pre-PETM} \), this and the results above imply a lower bound on pre-PETM \( pCO_2 \) when \( C_{SECO} = CS_{PETM} \) and \( pCO_2(EECO) = 760 \) ppm and an upper bound relationship between pre-PETM \( pCO_2 \) and PETM \( \Delta pCO_2 \) when \( C_{SECO} = CS_{PETM} \) and \( pCO_2(EECO) = 1260 \) ppm. If \( CS_{pre-PETM} > CS_{PETM} \), this implies a lower bound relationship between pre-PETM \( pCO_2 \) and PETM \( \Delta pCO_2 \) excursion when \( C_{SECO} = CS_{PETM} \) and \( pCO_2(EECO) = 760 \) ppm and an upper bound on pre-PETM \( pCO_2 \) when \( C_{SECO} = CS_{pre-PETM} \) and \( pCO_2(EECO) = 1260 \) ppm. These conditions and equations (1)–(3) yield the two dark blue lines in Figure 1; acceptable values for pre-PETM \( pCO_2 \) and PETM \( \Delta pCO_2 \) lie between the lines.

### 3.2. Pre-PETM Conditions

Observations of carbon isotopes and carbonate compensation depths (CCD) together with ocean carbonate chemistry and carbon cycle modeling provide further constraints. Previous PETM carbon cycle models have used a priori choices for pre-PETM \( pCO_2 \) (with associated ocean carbon inventory) and constant ocean phosphate inventories [Panchuk et al., 2008; Zeebe et al., 2009; Cui et al., 2011; Meissner et al., 2014]. We take a more general approach by considering the wide range of possible pre-PETM \( pCO_2 \) values from proxy reconstructions [Zachos et al., 2008; Beerling and Royer, 2011] and climate-dependent weathering including phosphate input to the ocean. For given lithosphere carbon emissions and weathering intensities, steady state conditions are established on million year timescales. For such conditions in the Danish Center for Earth System Science (DCESS) model used here [Shaffer et al., 2008], carbon and phosphate sources from lithosphere outgassing and/or weathering balance carbon and phosphate sinks in ocean sediment burial of organic matter and/or CaCO3 (Text S1). Given the slowly varying climate prior to the PETM [Zachos et al., 2008], we seek steady state solutions as initial conditions for PETM simulations.

Model radiative forcing was adapted to early Eocene conditions for solar forcing, albedo, high-latitude radiative forcing, and greenhouse gas concentrations to achieve in long model integrations mean atmospheric and deep ocean temperatures of 25°C and 10°C, respectively [Huber and Caballero, 2011; Lunt et al., 2012; Zachos et al., 2008] (Text S1). These temperatures were prescribed (bypassing the climate part of the model), ocean chemistry was adjusted to late Paleocene values, and biogeochemical steady states were sought for the full range of possible pre-PETM \( pCO_2 \) (Text S1 and Table S1). One difference between pre-PETM and PI simulations is the more diffuse, pre-PETM lysocline (Figure S2). A reconstruction of the pre-PETM lysocline shows a rather diffuse lysocline in the tropical Pacific Ocean [Panchuk et al., 2008]. An Indian Ocean reconstruction [Panchuk et al., 2008] appears sharper, but this may be an artifact of juxtaposition of subtropical sediment cores from ~5 km paleodepths with high-latitude cores at shallower paleodepths, in particular from well-documented Deep Sea Drilling Project site 259 [Zeebe et al., 2009; Hancock et al., 2009] (pre-PETM latitude and water depth of about 55°S and 4000 m). Another pre-PETM/PI simulation difference is high CaCO3 accumulation in deep, high-latitude pre-PETM sediments (Figure S2h). This stems from greater model biogenic CaCO3 production from warmer pre-PETM high-latitude surface layer temperatures (see parameterization in Text S1). Pre-PETM CaCO3 content (CaCO3 wt%) at 4000 m depth of about 50% are found in both the high-latitude model zone and in the data from site 259, the one high-latitude core with deep paleodepth available to test this.
Model steady state relationships between pre-PETM pCO₂ and carbon inventories for the standard pre-PETM MAT are given in Table S1 and plotted in Figure S3. For an ocean carbon inventory like PI but with late Paleocene conditions, atmospheric pCO₂ values are about 4 times greater than PI. This enhancement is due mainly to much higher ocean calcium content in the pre-PETM ocean and lower CO₂ solubility at higher temperatures. These properties and coupling to calcite saturation elevate ocean [CO₂] and depress ocean [CO₃²⁻]. Model land biosphere inventories are somewhat higher for higher pre-PETM pCO₂ from carbon fertilization (Figure S3).

3.3. Exploratory Simulations

The above steady states provide initial conditions for a series of 20,000 year simulations of the PETM initial phase for prescribed carbon inputs of different sizes but common time evolutions and for a common prescribed PETM MAT evolution with 5°C maximum warming (Text S1). This procedure again bypasses the climate part of the model to concentrate on carbon cycle response. PETM ΔpCO₂ values are given in Table S2 and contoured in pCO₂ space as carbon inputs in Figure 1. There is a significant pCO₂ increase from the warming alone (0 carbon input isoline in Figure 1) due to ocean outgassing from less CO₂ solubility and soil outgassing from more bacterial activity. Recalculations of PETM ΔpCO₂ values using constant, pre-PETM ratios carbonate to organic carbon production (rain ratios) show only modest changes (Table S2).

Carbon inputs to the atmosphere-ocean system lead to ocean acidification and CaCO₃ dissolution in the ocean sediment as well as decreased model biogenic CaCO₃ production (Text S1). These effects force shoaling of the CCD (here the depth of 10% CaCO₃ wt %). On the other hand, warming promotes increased biogenic CaCO₃ production from enhanced weathering that raises ocean phosphate concentrations and thereby new production as well as from enhanced rain ratios, an effect shown above to reproduce observed high CaCO₃ wt % in the deep, high-latitude, pre-PETM ocean. Increased biogenic CaCO₃ production forces CCD deepening. These opposing effects are reflected in model maximum, low-middle latitude CCD excursions for the PETM (Table S3 and Figure S4). For the PETM warming alone (0 carbon input) and our standard rain ratio expression (equation (S6)) the low-middle latitude CCD deepens by about 400 m. A recalculation using constant, pre-PETM rain ratios shows somewhat less shoaling (Table S3).

Observed low-middle latitude CCD shoaling across the PETM ranged from a few hundred meters in the Pacific Ocean to more than 2000 m in the South Atlantic Ocean [Panchuk et al., 2008; Zeebe et al., 2009]. Observations from the Indian Ocean indicate modest shoaling as in the Pacific [Zeebe et al., 2009]. In the late Paleocene/early Eocene, the Pacific Ocean was larger and the South Atlantic Ocean considerably smaller than now, accounting then for 57% and 7% of total, low-middle latitude ocean area, respectively (Table S4). If a South Atlantic CCD shoaling of 2000 m is taken to apply there only and a 200 m shoaling is assumed for the rest of the ocean, with basin size weighting we obtain a lower bound estimate on PETM CCD shoaling of 329 m that we round down to 300 m. If the South Atlantic shoaling is taken to apply to the whole Atlantic and a 400 m shoaling is assumed for the rest of the ocean, with this weighting we obtain an upper bound shoaling estimate of 673 m that we round up to 700 m. Thus, we adopt 300–700 m as our observation-based range for global mean, maximum low-middle latitude, PETM CCD shoaling.

For a pre-PETM carbon inventory similar to present day, this 300–700 m shoaling range is reproduced in the model for a carbon input of 2920–6530 Gt C (Figure S4; pre-PETM pCO₂ = 800). Without the warming feedbacks, i.e., with constant weathering, pre-PETM new production, and rain ratios, a much lower input of 1010–3150 Gt C forces such shoaling. This is an input range similar to that found in Zeebe et al. [2009] who did not consider warming feedbacks and thereby likely underestimated the PETM carbon input consistent with observed CCD shoaling. There are large uncertainties in modeling weathering and biosphere response to warming, but such nutrient and carbon cycle feedbacks were surely important for the workings of the PETM global warming event and should be addressed in PETM simulations.

Observed PETM CIEs vary from −2.5 ± 1.0‰ for benthic forams, −2.7 ± 1.0‰ for planktonic forams, −2.7 ± 1.1‰ for marine bulk carbonate, −4.1 ± 2.2‰ for marine organic matter, and −4.7 ± 1.5‰ for terrestrial records [McInerney and Wing, 2011]. These results are consistent with increased carbon isotope discrimination during photosynthesis for more dissolved CO₂ in the ocean surface layer, as implemented here (Text S1). PETM terrestrial records probably also reflect more carbon isotope discrimination during photosynthesis, for example, from increased humidity or pCO₂ [Bowen et al., 2004; Schubert and Jähres, 2013]. On the other hand, marine
carbonates may experience increased dissolution across the PETM and may therefore underestimate the CIE at some sites. Therefore, a “true” CIE from external carbon input is probably somewhat larger than the marine carbonate records and we choose a mean ocean-atmosphere-biosphere CIE, $C_{\text{IE}}^{\text{tot}}$, of $-3.5\%$ as our data-based target. With this and the above results we can now calculate the input $\delta^{13}C$ content, $\delta^{13}C_{\text{tot}}$, for given model carbon input and initial conditions (Text S1). The results are contoured in $pCO_2$ space in Figure S5. Possible PETM carbon input sources include organic carbon (about $-25\%$ in early Eocene times) [Falkowski et al., 2005], thermogenic methane ($-35$ to $-45\%$), and methane hydrate ($-60$ to $-70\%$). Volcanic carbon ($-5$ to $-7\%$) can probably be ruled out as a primary source: the very large volcanic carbon input needed to explain the observed CIE greatly exceeds inputs consistent with observed CCD shoaling. Therefore, we take $\delta^{13}C_{\text{tot}}$ to lie between $-25$ and $-70\%$.

4. Atmospheric CO$_2$, PETM Carbon Input, and Climate Sensitivity

The shaded area in Figure 1 defines the $pCO_2$ space for which all the above mineralogical, carbon cycling, and isotope constraints are satisfied. Pre-PETM $pCO_2$ is found to lie between 512 and 982 ppm with pre-PETM total carbon inventories between 30,780 and 40,000 Gt C. The PETM $pCO_2$ increase lies between 416 and 1500 ppm, corresponding to PETM carbon inputs of 2090 Gt C ($\delta^{13}C = -68.5\%$) to 6890 Gt C ($\delta^{13}C = -26.4\%$). Note that the range of acceptable pre-PETM $pCO_2$ is associated with pre-PETM total carbon inventories 0.76–0.99 times as large as a modern-day inventory (Figure S3). For lower initial inventories, less carbon input is needed to achieve observed PETM CCD shoaling and carbon isotope excursions (Figures S4 and S5).

With the use of equations (1) and (2), we can now calculate values for pre-PETM and PETM CS falling within the constrained $pCO_2$ space of Figure 1. These CS results and their ratios are plotted in Figure 2. We find that pre-PETM CS to lie between 3.3 and 6.8 K, while PETM CS lies between 3.4 and 6.5 K. These CS ranges for MATs of about 25 and 30°C, respectively, both extend well above upper estimates for present-day CS of 1.5–4.5 K [Flato et al., 2013] for a MAT of about 14°C, providing strong evidence for CS increase with warming. A further implication is then that as the Earth warms into the PETM, CS will not likely decrease, i.e., $C_{\text{PETM}}/C_{\text{pre-PETM}} \geq 1$. This is consistent with recent modeling indicating that any CS decrease with further warming may occur only for much warmer MAT (>40°C) [Popp et al.,...
2016]. The condition $\text{CS}_\text{PETM}/\text{CS}_\text{pre-PETM} \geq 1$ further constrains pre-PETM CS, PETM CS, and $\text{CS}_\text{PETM}/\text{CS}_\text{pre-PETM}$ to the ranges $3.3–5.6\, K$, $3.7–6.5\, K$, and $1–1.68$, respectively (Figure 2), and also puts slightly tighter bounds on our $p\text{CO}_2$ and carbon input estimates. Pre-PETM $p\text{CO}_2$ now lies between 592 and 982 ppm with pre-PETM total carbon inventories between 32,500 and 40,000 Gt C, and the PETM $p\text{CO}_2$ increase lies between 416 and 1348 ppm, corresponding to PETM carbon inputs of 2090 Gt C ($\delta^{13}\text{C} = -68.5\%$) to 6210 Gt C ($\delta^{13}\text{C} = -28.8\%$). There would be still tighter bounds on the above properties if constrained by an EECO $p\text{CO}_2$ range of 930–1260 ppm, a range consistent with both the above nahcolite constraint and very recent $p\text{CO}_2$ estimates from boron isotope data [Anagnostou et al., 2016].

The above results are quite robust to parameter value choices. For example, for a range of 5–7°C for pre-PETM warmth due to higher $p\text{CO}_2$ ($\Delta T_{\text{pre-PETM}}$ in equation (2)), our analysis yields 572–982 ppm for pre-PETM $p\text{CO}_2$, 398–1460 ppm for the PETM $p\text{CO}_2$ increase, $2020\, (\delta^{13}\text{C} = -70\%)$–$6780\, (\delta^{13}\text{C} = -26.6\%)$ Gt C for the carbon input, 2.8–6.3 K for pre-PETM CS, and 3.5–6.6 K for PETM CS. If we choose a CIE$^{\text{pat}}$ of $-4\%$ as our data-based target, we find 592–982 ppm for pre-PETM $p\text{CO}_2$, 454–1348 ppm for PETM $p\text{CO}_2$ increase, $2350\, (\delta^{13}\text{C} = -70\%)$–$6210\, (\delta^{13}\text{C} = -32.8\%)$ Gt C for the carbon input, and 3.3–6.6 K for PETM CS.

The results in Figure 1 permit a rather broad range of PETM carbon input scenarios, but some appear more likely. The highest permissible carbon input of 6210 Gt C ($\delta^{13}\text{C} = -28.8\%$) could be explained by an organic carbon injection, but such an input would be about twice the size of our pre-PETM land biosphere (Figure S3) or that in other work [Beerling, 2000]. Organic carbon reserves about half the required size have been proposed for pre-PETM permafrost [DeConto et al., 2012]. However, proxy-based reconstructions of high-latitude climate for the late Paleocene/early Eocene show mean surface temperatures well in excess of 10°C [Huber and Caballero, 2011; Lunt et al., 2012; Kemp et al., 2014], too warm to allow extensive permafrost at high latitudes even at elevations above 1000 m. Other explanations in terms of organic carbon also fall short or lack support in the paleorecord [Panchuk et al., 2008; Moore and Kurz, 2008]. If for these reasons we rule out organic carbon as the sole source of the PETM carbon input, then this input must have involved considerable amounts of methane.

The lowest permissible input of 2020 Gt C ($\delta^{13}\text{C} = -70\%$) could be explained by methane hydrate destabilization. However, there is an ongoing debate as to whether this much methane hydrate could exist for warm pre-PETM conditions [Buffett and Archer, 2004; Gu et al., 2011]. Such a scenario would require very high CS values with a CS increase into the PETM from 5.6°C to 6.5°C. One possible scenario centered within the acceptable $p\text{CO}_2$ space (star in Figure 1) would be thermogenic methane input [Svensen et al., 2004] of 3740 Gt C ($\delta^{13}\text{C} = -43.7\%$) into a pre-PETM state with 800 ppm $p\text{CO}_2$ and a total carbon inventory of 36,730 Gt C. This would raise $p\text{CO}_2$ at the PETM maximum to 1600 ppm, lead to a maximum, mean low-middle latitude CCD shoaling of 394 m, and be associated with a CS increase into the PETM from 4.0° to 5.0 K. However, other possibilities cannot be excluded like some combination of more modest amounts than considered above of methane hydrate, thermogenic methane, organic matter, and/or direct volcanic carbon input.

5. Discussion and Conclusions

Here we have taken a new approach to the problem of climate change and carbon cycling across the PETM global warming event. We use new climate reconstructions for late Paleocene/early Eocene conditions and for the PETM to set the stage for an analysis of carbon cycling and climate sensitivity. For that analysis, we compare paleoreconstructions with Earth System model simulations that consider a wide range of possible pre-PETM $p\text{CO}_2$ and associated carbon inventories for pre-PETM ocean chemistry and that include climate feedbacks on nutrient and carbon cycling. For applied pre-PETM and PETM warming, we apply joint mineralogical, carbon chemistry, and carbon isotope constraints to estimate possible ranges of pre-PETM and PETM $p\text{CO}_2$ and the size and nature of the PETM carbon input. Finally, we use these results and some consistency considerations to estimate possible ranges for pre-PETM and PETM CS.

Our pre-PETM CS range of 3.3–5.6 K is for a MAT of about 25°C, while the PETM CS range of 3.7–6.5 K is for a MAT of about 30°C. For comparison, the range of present-day CS estimates is 1.5–4.5 K for a MAT of about 14°C whereby about one fourth this CS is due to ice-albedo feedback [Flato et al., 2013], not contributing to CS in the warm late Paleocene era. Furthermore, several recent studies yield present-day CS estimates in the
In some complex climate system models, CS was found to increase for rising MAT, in particular due to some combination of fast water vapor and cloud feedbacks [Caballero and Huber, 2013; Meraner et al., 2013; Popp et al., 2016]. Our results lend some credeence to these model findings. The compilation in Figure 3 supports a CS increase with MAT but permits few conclusions on the structure of this increase. One could define a linear increase with MAT based on our results, but these results also allow for an abrupt, nonlinear CS rise for a relatively small warming as was found in the models cited above.

Our findings support the notion that as pCO2 and global temperatures continue to rise, there may be additional warming as CS may increase despite less ice-albedo feedback as snow and sea ice cover diminish. If warming increases CS, greater CS makes the Earth warm more, amplifying the warming [Bloch-Johnson et al., 2015]. Thus, our results further underline the need to limit ongoing global warming by greatly reducing anthropogenic greenhouse gas emissions as soon as possible.

References


