Reviewer 1

In this study the biogeochemical and biogeophysical effects of vegetation on the climate system are analysed separately over the time span of the last 120 000 years. Spanning the last ice age inception, last glacial and the latest deglaciation, 62 "snapshot"-type (i.e., under constant forcing) simulations were integrated, distributed over that period, with HadCM3, a fully coupled atmosphere-ocean GCM with interactive vegetation. In addition, 5 transient simulations were integrated over the whole period with various versions of the cGENIE model. These simulations were based on terrestrial carbon fluxes diagnosed from the HadCM3 simulations. The authors conclude that the biogeophysical effects of vegetation account for additional mean cooling during the glacial and, in some cases, substantial regional cooling. The biogeochemical effects are smaller and of opposite sign. The authors also emphasize that different timescales are involved for these two effects on climate.

I have enjoyed reading the paper. It is well written and presents interesting results. I do have three major comments and a long list of other comments and recommend that this paper might be suitable for publication after all of my comments are addressed (i.e. major revisions).

We're very glad the reviewer enjoyed the paper and think it is interesting, and thank them for their helpful and detailed comments. Our responses to the reviewer's comments (in bold) are below (in standard text).

Major comments:

1. One underlying hypothesis with this approach is that the feedbacks considered (ocean, vegetation, atmosphere, sediments, biogeophysical, biogeochemical) add up linearly (i.e. there is no non-linear interaction between the biogeophysical and biogeochemical feedbacks). This assumption needs to be clearly stated and discussed.

We've added the following to the discussion:

Our approach here assumes that there is no non-linear interaction between the biogeochemical and biogeophysical effects. Since the biogeochemistry acts as a negative feedback and reduces over time, and the biogeophysics acts as a positive feedback and stays the same over time, there's no strong reason to believe that in equilibrium there would be any significant synergy. However, on shorter timescales and on a regional rather than global scale, it is quite possible that there could be some synergies.

2. Under which boundary conditions cGENIE was spun up? Am I right to assume that 120kaBP boundary conditions were used? Am I also right to assume that the diagnosed terrestrial carbon fluxes (calculated based on changes in terrestrial carbon) from the HadCM3 snapshot simulations were interpolated as "emission" forcing time series and applied to cGENIE? And that these "emission" forcing time series were the only changing boundary conditions during the simulations (i.e., no additional imposed CO2 changes, no continental ice sheet changes, etc)? If I understood this right, there is another assumption made by this approach: that the climate sensitivity is constant and independent of the climate state. This assumption also needs to be clearly stated and discussed.

All good questions.

Firstly, although the model configuration used was cited, without tediously reading the paper concerned, we admit that it is not clear what specific boundary conditions are assumed here,

particularly as our study considered changes occurring over a full glacial cycle. The cGENIE model configuration was modern (pre-industrial, a.k.a. late Holocene). We have now clarified this explicitly in the text.

So the overarching question (encompassing the reviewer's comments) then arises: what is the sensitivity of the (marine) carbon cycle and climate to an impulse of CO₂, considering that the boundary conditions of ice sheets, orbits, radiative forcing, sea-level, ocean circulation and chemistry, are all continually changing? This is way beyond what we can explicitly (i.e. mechanistically in the model) address here, hence the simplistic use of the modern configuration of the cGENIE model. However, we should have been more explicit in the text about the implications of this. We have now added new discussion surrounding this issue and in what ways our methodology might induce artifacts in the diagnosed contribution of biogeochemical effects to the overall glacial-interglacial climate change.

Regarding the details of the emissions forcing in cGENIE – yes, the reviewer is correct (in that they are interpolated from the GCM snap-shots). We have now made this clearer in the text.

3. As shown in Figure 3b and described on Page 8 (lines 11-15), the vegetation changes trigger a change in ocean circulation. While I agree that this is a model dependent result and not part of the core results (although certainly influencing these core results), it would be interesting to see time series of AMOC for the static and dynamic simulations. In case there is a simple feedback that can be easily crystallized (such as the one mentioned in the text), it would also be good to analyse the results further, verify that this is indeed the feedback and play and describe this feedback in more detail.

We've added the following explanation of the AMOC changes:

Although the biogeophysical changes cause cooling, there are some minima of biogeophysical temperature change seen at 30 ka, 56 ka and 100 ka (Figure 2, filled symbols). These minima have an oceanic source and are caused by vegetation interacting with thermohaline circulation changes. In our new simulations we account for the net transport of water from ocean to the ice sheets by a parameterisation that instantaneously balances any net accumulation of water on ice. This parameterisation results in fresher ocean conditions during times of precession driven N. Hemisphere summer insolation highs (less water is being used to build the ice sheets). The instantaneous nature of the parameterisation is physically unrealistic but reductions in accumulation and an increase in ablation during precession highs has been seen in fully coupled climate-icesheet EMIC simulations (e.g. Ganopolski et al, 2010). During weaker accumulation periods, the parameterisation results in a freshening of ocean surface waters and a reduction in AMOC strength from ~16Sv to 10-12Sv.

Superimposed upon this general behaviour, the addition of interactive vegetation generally does not change the AMOC strength. However, at times of weak AMOC, small changes in runoff and temperature are sufficient to cause some changes in the response. For instance, in the static vegetation simulations there is a relatively weak AMOC in the simulations for 60ka, 58ka, and 56ka. In the interactive vegetation simulations, the weakened AMOC only occurs at 60ka. Thus at 60ka the changes in climate are fairly typical of preceding times but at 58ka and 56ka there is a substantial difference between the static and dynamic vegetation simulations. The cause for this difference is associated with a combination of reduced runoff into the N. Atlantic (principally from changes in land surface in N. America) and colder temperatures, both of which act to stabilise the AMOC in all three periods but it is sufficient to prevent the AMOC weakening in the 58 and 56ka simulations.

This threshold like behaviour of the AMOC is almost certainly highly model dependent and hence the result is not robust.

Other comments:

* Permafrost and wetlands are not (cannot be) resolved correctly. Both play important roles in terrestrial carbon feedbacks. While the Discussion briefly mentions the lack of permafrost related feedbacks in the simulations (Page 14, lines 16-23), it might be interesting to add a discussion about wetlands (changes in wetlands in the tropics, but also changes from permafrost to wetlands or vice-versa at high latitudes). Can you please broaden this discussion, including some key references, and, if possible, add an uncertainty range to your results in the Discussion section.

The following has been added to the discussion:

The soil carbon change under ice-sheets between PI and LGM is modelled as ~220 PgC. Extrapolating from the present day underestimates of the model, we could speculate that this might actually be a third too little. If the true value were ~330 PgC, this would make the total C change PI to LGM 550 PgC. This would put the change more in line with some previous estimates. It would affect the global mean annual biogeochemical contribution by ~0.1 K. This would mean the net effect of vegetation was closer to zero, but the biogeophysical effect would still dominate.

However, the exact size of the terrestrial carbon emissions is uncertain. Other carbon stores not accounted for here are potentially important, for example methane during sea level rises or changes to the wetlands in the tropics. Modelling studies that look at wetlands at the LGM suggest that although the wetland area is larger, but the methane emissions are lower compared to modern day (Kaplan, 2002). However, paleohydrological data indicates a drying in the African tropics (Gasse, 2000). Our model does not have a process based permafrost or wetlands component, and therefore the changes in methane are not accounted for. This is a particular limitation when considering the carbon stored in deep permafrost soils in Northern peatlands. Saito et al. (2013) show that, based on the temperature changes, there is a substantial expansion of permafrost area during glacial times but cannot estimate any changes in carbon storage. Zimov et al. (2006, 2009) have argued that permafrost storage could be a major source of carbon through the deglaciation, and Ciais et al. (2012) argue that there was a large extra pool of inert carbon at the LGM. Similarly, Köhler et al. (2014) have argued that large amounts of carbon were locked into permafrost which were then released rapidly at the Bolling-Allerod.

* Coarse resolution and numbers of PFTs (page 5, lines 27-32): The representation of vegetation (and therefore associated feedbacks) is crude in TRIFFID (only 5 PFTs, coarse spatial resolution, crude representation of terrestrial nutrient cycles). While this is not any worse than in most other coupled models, the implications should be discussed in more depth in the discussion section.

We've added the following about the effect of the PFTs to the discussion section:

The impacts are mainly determined by the vegetation shifts the DGVM simulates. Each gridbox has the potential for 5 PFTs, but generally the Lotka-Volterra equations used in TRIFFID mean that the gridbox is dominated by one PFT. The small number that means the range within each PFT is relatively large. Therefore the model probably underestimates the effects of small perturbations in climate, as the large definition of the PFTs allows the PFT to remain the same. Conversely, it makes an abrupt change more likely as the climate tips a girdbox from being predominantly one PFT to being predominantly another. Overall, the model could be slightly underestimating the amount of change in vegetation. However, because of the ratio of the biogeophyiscal to biogeochemical

changes, if the vegetation change is underestimated, the sign of the net effect of the terrestrial biosphere is unlikely to change. Similarly, because on the long time periods involved much of the released carbon is taken up by the ocean, the changes in carbon densities of the vegetation would need to be wrong by a lot to change the overall signal.

* It strikes me in Figure 1 that there is almost no change in tropical rain forest cover. Is that realistic?

We've added the following discussion to section 3.1:

The forest extent in the tropics at the LGM is similar to PI (see SI Figure 9 for shifts in vegetation at 21 ka). This is supported by pollen and other data (Maslin et al., 2012; Anhuf et al., 2006), and modelling (Cowling et al., 2001) which find that the although there is diminished tropical forest, there is still substantial tree cover at the LGM and little sign of widespread grasslands. Because of the PFT (rather than biome) approach of TRIFFID, and the limited number of PFTs, it's difficult to be sure whether trees in the tropics are a tropical rainforest at the LGM, because they equally could be temperate forest.

Would it be possible to include a validation of these results (present day bias + comparison to pollen data from LGM for example)?

The following has been added to the paper:

The climate model used in Hoogakker et al. 2016 is HadCM3B-M1 and the climate model used here is HadCM3B-M2.1. The climate between these two is virtually identical. Since the climate is the main aspect which determines the distribution of vegetation in a DGVM, the verification of Hoogakker's work suggests that the distributions found here are also reasonable.

Comparison with the LGM BIOME6000 dataset shows a broad agreement. The model has considerable expansion of grasses in Eurasia where BIOME6000 has grassland and dry shrubland. Broadly speaking, North America shows little change from the mid Holocene to LGM. One key weakness of the model is in western Europe, where BIOME6000 shows grassland and dry shrubland, whereas the model has shrubs and needleleaf trees.

In the present day, TRIFFID does a reasonable job, as detailed in Valdes et al. (2017).

* Page 13, lines 5-8: this is an interesting (although neither surprising nor new) result that feeds back into the discussion about climate sensitivity over long timescales. I would suggest adding a paragraph to the discussion about the different timescales involved and the implications on climate sensitivity. It would also be good to remind the reader, which of these feedbacks are usually incorporated in the state-of-the-art IPCC models (i.e. expand the second paragraph in Discussion).

The following has been added to the relevant paragraph of the discussion:

From a climate sensitivity point of view, this means that on shorter timescales, the effects of dynamic vegetation can cancel each other out. This provides some rationale for the fact that dynamic vegetation has been generally not included in the majority of state-of-the-art earth system models used in CMIP5, as it doesn't significantly affect the climate sensitivity. At longer time scales, it is more important to include dynamic vegetation, as without the positive feedback of the biogeophysical effects, the climate sensitivity would be under estimated.

* Table 1: I am certainly misunderstanding something here. . . Why does the vegetation C change differ for all 4 set-ups? Shouldn't vegetation C only be affected by new land (especially because the atmosphere does not "see" the carbon released from under the ice)? In which case the two ELE simulations should be very similar, if not the same (same for the two ELI simulations)?

Looking again at table 1, it could be clearer. The vegetation C differs in all four scenarios because all four are compared to the PI veg and soil carbon change, and all have a different combination of land. Since they're all compared to PI, where there is vegetation and soil carbon where at the PI there is glaciers, both are included. Critically, areas that at the LGM had ice-sheets, had forests etc. at PI, so there is some vegetation carbon difference. The rounded values for each aspect are on page 11, lines 3-5. To help make this clearer, we've amended these so the exact values as calculated in the model are given.

* HadCM3 simulations: it is my understanding that the HadCM3 simulations are run under constant external forcing, initialized from the previous MOSES 1 simulation, then run for 300 years with equilibrium TRIFFID (50 years of TRIFFID for each 5 years of the climate model run), and finally integrated 300 years with dynamic TRIFFID (where TRIFFID is called every 10 days). I am puzzled by this approach – I would assume that TRIFFID is in equilibrium after the first 300 years of "equilibrium setting". I would also assume that atmospheric conditions are fairly close to equilibrium after these first 300 years, so TRIFFID in "equilibrium setting" saw internal forcing with little drift. Why integrate TRIFFID in dynamic setting for the last 300 years, if the forcing that TRIFFID "sees" is more or less constant in a climatological sense and if TRIFFID is already in equilibrium? This does not change the results presented here, it just seems weird (e.g. why not run TRIFFID in equilibrium over the whole 600 years of constant boundary conditions?). See also text on Page 11, lines 15-16 "the model is run for sufficient length of time for the soil and vegetation carbon to reach equilibrium".

This is an interesting question, and we agree that it's not immediately evident in the paper what the rationale was for our methodology. We hope the following will help clarify.

In equilibrium mode TRIFFID is prone to considerable fluctuations around the 'true' equilibrium determined by the dynamic mode, as the 50 years it runs with can be prone to bias due to interannual variability. Therefore, we consider equilibrium mode a tool for spin-up, and wherever possible use dynamic mode in simulations for publication – that's why we don't just run it for 600 years in equilibrium mode. In dynamic mode TRIFFID can take a long time to fully equilibrate the vegetation, and the soil carbon can only be in equilibrium after the vegetation (because of the litter changes). However, these changes are relatively small.

Therefore, you're right that it probably doesn't make that much difference to the final simulation results. But we consider it due diligence, and have reported as clearly and honestly as possible what we have done to make to create these simulations.

* Set of simulations with static vegetation based on PI simulation (page 4, lines 11- 14): I assume that the vegetation in this simulation is masked out under continental ice sheets (adjusted for the ice sheet extension of the period of interest). Can you please add this here.

You're absolutely right that in the Static simulations the land cover is the PI vegetation cover, with the time-appropriate continental ice-sheets imposed. We specify on line 16 that the ice-sheets are the same in both simulations, we think it's clearer to talk about the differences only, and the

similarities in a separate paragraph. Therefore, to help ease any potential confusion, we've added to lines 11 -14 a note to see below for the details of aspects which are the same in the two sets.

* Why are the EPICA anomalies halved? Is that to get a representation of global temperature changes (versus changes in local temperature)? Can you please clarify this in the text?

You're correct, we use the halved EPICA anomalies as an approximation of global temperature changes. We mention this on page 7, line 2-3, but have also added this to the figure caption.

* I do not understand what is shown in Figure 4a. Is it the globally integrated (or globally mean) albedo anomalies? If this is the case; why don't they add up? Or are these means over land/ocean versus global mean? Can you please clarify?

We apologise, Figure 4 had the incorrect key. We've rectified this, and hopefully it now makes more sense.

* Figure 5: can you please show these plots for simulations that treat carbon under ice sheets the same way? These plots should only show the difference in C and albedo due to changes in vegetation and soils away from ice sheets. The large purple areas in 5e and f take the attention away from the results.

We've changed these plots to mask over the carbon under ice-sheets, and added the original plots to the supplementary information.

* cGENIE simulation TRCE needs to be described in more detail in Section 2.2.

The TRCE is currently explained on lines 26 - 29 and we acknowledge that this could lead to readers thinking it is a cGENIE simulation. To remedy this, we have included the explanation about TRCE in the overview of the methodology at the top of section 2.

Editorial changes:

* Page 4, line 6: I would prefer to go forward in time: 120 – 0 kaBP. Done

* Page 4, line 21-24: again, please go forward in time; e.g. 120 to 80 kaBP,... Done

* The filled points are hard to see in Figure 2a

Whilst we appreciate the filled points are difficult to see in this particular plot, the x axis is the same as on figure 2b, and so the filled points should be easy to infer. Also, the filled points are discussed in relation to figure 2b. 2a is already a 'busy' plot, so we're reluctant to make the points larger, or to break with the pattern set of using the filled points to indicate the time periods of interest.

* Figure 4: I find it hard to discern the different lines – would it be a good idea to plot these time series in different colours?

We've put the lines in different shades of red, so hopefully it's easier to see which are which. We plotted these line in red to tie in with the colour scheme in other plots, as they all use the GCI-ELE scenario. We're reluctant to use a third colour scheme in the paper, as we feel it could be confusing to readers and reduce the visual cohesiveness of the paper.

* Page 9, line 6: "is" missing Done.

* Figure 6: the red and pink lines are hard to discern

We've replaced the pink with a green.

* Figure 7: Figure caption should read GCI_ELE and not GLI_ELE

Corrected.

Comment on Colin's comment: I agree with the authors that this is a sensitivity study, showing the whole range of possible scenarios (including unlikely ones). I would not recommend reducing the numbers of scenarios shown and discussed in the text. However, I agree with Colin that it should be made clear in the abstract that the higher terrestrial carbon loss scenarios are more likely (this is the one the cGENIE time series are shown for in Figure 7).

We have added to following proviso in the abstract:

In addition, depending on the assumptions about soil carbon under ice-sheets and sea level rise, we find a range in terrestrial carbon storage change from a reduction in LGM carbon storage of -440 PgC, to a gain of +37 PgC, *though we consider the negative part of the range more likely*.

Side note: a newer estimate of total terrestrial carbon glacial/interglacial change based on benthic d13C data is given by Menviel et al. (2017).

Thank you for drawing our attention to this paper. We have included it in the discussion.

Reviewer 2

General comments

In this paper the authors present results of 62 equilibrium simulations with a coupled atmosphereocean-vegetation general circulation model covering the last glacialinterglacial cycle. They performed simulations with and without dynamic vegetation to quantify the effect of the terrestrial biosphere on glacial-interglacial climate variability in terms of both biogeophysical and biogeochemical effects. Although the results represent an important contribution to our understanding of the role of the terrestrial biosphere to glacial-interglacial variability, I have some major comments that should be addressed before this paper can be considered for publication in Climate of the Past.

We thank the reviewer for their view that this work is an important contribution to the literature. We address their comments (in bold) in standard font below.

Since changes in vegetation cover are key to the results presented in this study, the modelled vegetation should be compared with available reconstructions, where available. The BIOME6000 dataset for the last glacial maximum provides a unique reconstruction of vegetation cover for glacial climate conditions and could be used to evaluate the model performance. As a minimum requirement, model results and reconstructions should at least be compared qualitatively and discrepancies discussed. Comparison with other modelling studies would also be useful (e.g. (Prentice et al. 2011; Hoogakker et al. 2016)).

The following has been added to the paper:

The climate model used in Hoogakker et al. 2016 is HadCM3B-M1 and the climate model used here is HadCM3B-M2.1. The climate between these two is virtually identical. Since the climate is the main aspect which determines the distribution of vegetation in a DGVM, the verification of Hoogakker's work suggests that the distributions found here are also reasonable.

Comparison with the LGM BIOME6000 dataset shows a broad agreement. The model has considerable expansion of grasses in Eurasia where BIOME6000 has grassland and dry shrubland. Broadly speaking, North America shows little change from the mid Holocene to LGM. One key weakness of the model is in western Europe, where BIOME6000 shows grassland and dry shrubland, whereas the model has shrubs and needleleaf trees. Similarly, assessment of the PI vegetation cover of HadCM3 by Valdes et al. (2017) shows good agreement with reconstructions of 1800 vegetation.

The representation of vegetation cover in Figure 1 in terms of dominant PFTs can be misleading. For example it seems from Figure 1 that tropical forest remains practically unchanged during glacial conditions. Is this just an artifact of the dominant PFT representation or is it a real feature of the model (in which case the discrepancy with available reconstructions has to be discussed)? In any case I would instead suggest showing fractions of all 5 PFTs separately, at least for preindustrial and LGM.

We've included the PFT changes PI-LGM separately in the supplementary information. The extent of change of broadleaf forest in the tropics changes relatively little. However, as discussed above, this is similar to many other estimates.

Figure 1 shows the dominant PFT, which inevitably 'hides' not only decrease in the proportion of broadleaf trees, (the dominant PFT is simply the one with the largest proportion, even if that

proportion is low), but also shifts within that climatic envelope. In particular, a broadleaf tree is not necessarily 'tropical rain forest', but equally can be a temperate broadleaf forest, or even savannah-type trees.

We've also added the following discussion to section 3.1:

The forest extent in the tropics at the LGM is similar to PI (see SI Figure 9 for shifts in vegetation at 21 ka). This is supported by pollen and other data (Maslin et al., 2012; Anhuf et al., 2006), and modelling (Cowling et al., 2001) which find that the although there is diminished tropical forest, there is still substantial tree cover at the LGM and little sign of widespread grasslands. Because of the PFT (rather than biome) approach of TRIFFID, and the limited number of PFTs, it's difficult to be sure whether trees in the tropics are a tropical rainforest at the LGM, because they equally could be temperate forest.

The results about the biogeophysical feedback are presented in a rather superficial way, which makes it difficult to get a quantitative understanding of the processes responsible for the positive feedback. As the authors mention in the introduction, the biogeophysical feedback results primarily from the vegetation controlling the surface energy fluxes. The results section in the paper focuses almost exclusively on the effect of changes in surface albedo. Latent heat effects are only mentioned once when referring to Figure 4b but are not discussed further, and sensible heat flux changes are not mentioned at all. Albedo changes are probably the dominant effect, but the other effects should also be quantified. I would suggest to add panels showing the changes in latent and sensible heat flux to Figure 3 and to move the albedo plots from Figure 5 to Figure 3 (it is not clear to me why albedo maps are shown together with vegetation and soil carbon). Also, I would suggest replacing the albedo figures with net shortwave radiation absorbed at the surface. Shortwave radiation is a more appropriate measure because it accounts for changes in insolation and can moreover directly be compared to the latent and sensible heat fluxes.

We have added figures of the net shortwave radiation, sensible heat, and latent heat in the supplementary information. Whilst we understand the reasons for the reviewer wishing to see figures of net shortwave radiation absorbed at the surface, we hope that by showing these other metrics in the SI, you'll understand why we feel the albedo is more informative, as well as a more generally understood and used metric.

The authors show that the vegetation is interacting with the thermohaline circulation. It would be interesting to understand how this is happening. Can anything be said about possible causal relations, given the available simulations? Is vegetation affecting runoff into the North Atlantic, or are vegetation and THC interacting via changes in atmospheric circulation?

We've added the following explanation of the AMOC changes:

Although the biogeophysical changes cause cooling, there are some minima of biogeophysical temperature change seen at 30 ka, 56 ka and 100 ka (Figure 2, filled symbols). These minima have an oceanic source and are caused by vegetation interacting with thermohaline circulation changes. In our new simulations we account for the net transport of water from ocean to the ice sheets by a parameterisation that instantaneously balances any net accumulation of water on ice. This parameterisation results in fresher ocean conditions during times of precession driven N. Hemisphere summer insolation highs (less water is being used to build the ice sheets). The instantaneous nature of the parameterisation is physically unrealistic but reductions in accumulation and an increase in ablation during precession highs has been seen in fully coupled climate-icesheet EMIC simulations (e.g. Ganopolski et al, 2010). During weaker accumulation periods, the

parameterisation results in a freshening of ocean surface waters and a reduction in AMOC strength from ~16Sv to 10-12Sv.

Superimposed upon this general behaviour, the addition of interactive vegetation generally does not change the AMOC strength. However, at times of weak AMOC, small changes in runoff and temperature are sufficient to cause some changes in the response. For instance, in the static vegetation simulations there is a relatively weak AMOC in the simulations for 60ka, 58ka, and 56ka. In the interactive vegetation simulations, the weakened AMOC only occurs at 60ka. Thus at 60ka the changes in climate are fairly typical of preceding times but at 58ka and 56ka there is a substantial difference between the static and dynamic vegetation simulations. The cause for this difference is associated with a combination of reduced runoff into the N. Atlantic (principally from changes in land surface in N. America) and colder temperatures, both of which act to stabilise the AMOC in all three periods but it is sufficient to prevent the AMOC weakening in the 58 and 56ka simulations. This threshold like behaviour of the AMOC is almost certainly highly model dependent and hence the result is not robust.

In the model description section, no information is given on the soil carbon representation in the model.

The following has been added to the model description:

The soil carbon is a single pool, increased by litterfall and decreased by respiration. The soil respiration is controlled by moisture and temperature and returns carbon dioxide to the atmosphere unless, as is the case here, the atmospheric carbon dioxide is fixed. The litterfall is an area-weight sum of the litterfall of the five PFTs in each gridcell.

I expect the results of the biogeochemical part to strongly depend on how soil carbon is represented in the model. In particular, a proper representation of carbon stored in permafrost is probably crucial to model land carbon storage during glacial times. In the discussion section the authors mention that the model does not have a process-based permafrost component, but this should also clearly be stated in the description section.

We have added a statement to the model description that there is no permafrost component.

The amount of carbon which can potentially be buried below the ice sheets will strongly depend on how carbon in frozen soils is treated. The authors should discuss this in more detail. How does the carbon stored in permafrost in the model compare to observational estimates (Hugelius et al. 2014) for the present day?

The following has been added to the discussion:

Carbon in frozen soils is treated essentially the same way as any other soil. The soil carbon respiration is dependent on temperature and soil moisture. Where the temperature is near or below freezing, little or no respiration occurs, causing a build up or retention of soil carbon. There is no soil carbon in the model under icesheets, therefore the values are extrapolated from the soil carbon present when there aren't icesheets (e.g. at PI).

For present day, Hugelius et al. 2014 shows around 75 – 100 kg carbon m2 far north Siberia, 20 -40 further south. Far northern Canada is much more heterogeneous, with values from 20 – 150 kg C m2. The modelled PI values are on the low side, and much more homogeneous, around 15 – 20 kg C

m2, but is similar to Hugelius et al. 2014 in that it shows far north America to be less consistent, with some higher areas of 35-40 kgC m2 in the far north. (See the now supplementary figure of the loss of soil carbon under icesheets at LGM.)

What this suggests is that while on the correct order of magnitude, the model has a very modest amount of soil carbon that could be considered permafrost. Therefore, we think it's reasonable to include this low estimate of soil carbon in the uncertainties.

The soil carbon change under icesheets between PI and LGM is modelled as ~220 PgC. Extrapolating from a comparison with Hugelius et al. 2014, we could speculate that this might actually be a third too little. If the true value were ~330 PgC, this would make the total C change PI to LGM 550 PgC. This would put the change more in line with some previous estimates. It would affect the global mean annual biogeochemical contribution by ~0.1 K. This would mean the net effect of vegetation was closer to zero, but the biogeophysical effect would still dominate.

In the biogeochemistry results section the effect of dynamic vegetation on land carbon storage is not discussed, although the differences in vegetation and soil carbon between static and dynamic simulations at LGM is shown in Figure 5 (the 30 kyr maps of vegetation and soil carbon in Figure 5 seem redundant to me). If I understand correctly, in this section only the dynamic vegetation simulations are discussed. This should be clarified in the text.

In section 3.3 the Dynamic simulations are discussed in relation to the Static simulations. As mentioned in the methods, the Static simulations have the PI vegetation cover and carbon stores (i.e. there is no terrestrial carbon cycle in the Static simulations). Therefore, we consider discussing the difference between the Static and Dynamic simulations to be discussing the effects of dynamic vegetation.

We include the 30 ka maps to highlight that whereas the albedo shows a very different pattern between 21 ka and 30 ka, the overall pattern of carbon changes remains similar but slightly smaller, which accounts for why the net biogeophysical and biogeochemical effects vary considerably between these two simulations. Since the focus of the paper is understanding the net effects of vegetation, we feel they are informative to include.

A figure showing the differences in land carbon storage between LGM and present day would be helpful.

We show the carbon storage at the LGM (and all other time points) in the supplementary information (figure 8 in the discussion manuscript). In addition, the differences in land carbon storage can be seen in Figure 5c and 5e for the vegetation and soil respectively (which together make up the land in this model) carbon maps for the LGM. As explained in the methods, because the Static simulations have the PI vegetation cover, they also have the PI carbon storage. Therefore, for the carbon the PI – LGM anomaly is the same as the Static – Dynamic. We have added this information to the Figure 5 caption.

Specific comments

Page 1, line 7: specify that the 62 simulations are 'equilibrium' simulations. Done.

Page 1, line 18: ocean/atmosphere

Done.

Page 3, line 13: vegetation carbon -> land carbon Done.

Page 4, line 14: remove brackets Done.

Page 4, line 29: timer -> time Done.

Figure 1: legend is hard to read We've increased the size of the text in the legend.

Page 6, line 7: -0.91°C was -0.84°C in the abstract (if I understand correctly)

Our apologies, the abstract incorrectly used the LGM rather than the largest value. The abstract has been corrected to -0.91 (as shown in Figure 2b) rather than the LGM value of -0.84.

Figure 3: Are the figures for annual mean characteristics? Please specify.

Yes, they are mean annual values. This has been added into the figure caption.

Figure 4: use of different colors for different lines would improve readability

We've changed the lines to different shades of red as well as the patterns, so hopefully it's easier to read. We plotted these line in red to tie in with the colour scheme in other plots, as they all use the GCI-ELE scenario. We're reluctant to use a third colour scheme in the paper, as we feel it could be confusing to readers and reduce the visual cohesiveness of the paper.

Page 9, line 6: it IS unclear Added.

Page 10, line 2: carbon stores changes -> carbon stores Done.

Page 14, lines 8-11: shelf carbon stocks values should be positive Done.

Page 14, lines 13-15: check sentence Corrected.

Quantifying the Influence of the Terrestrial Biosphere on Glacial-interglacial Climate Dynamics

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Abstract. The terrestrial biosphere is thought to be a key component in the climatic variability seen in the paleo record. It has a direct impact on surface temperature through changes in surface albedo and evapotranspiration (so called biogeophysical effects) and in addition, has an important indirect effect through changes in vegetation and soil carbon storage (biogeochemical effects) and hence modulates the concentrations of greenhouse gases in the atmosphere. The biogeochemical and biogeophysi-

- 5 cal effects generally have opposite signs meaning that the terrestrial biosphere could potentially have played only a very minor role in the dynamics of the glacial-interglacial cycles of the late Quaternary. Here we use a fully-coupled dynamic atmosphere-ocean-vegetation General Circulation Model (GCM) to generate a set of 62 equilibrium simulations spanning the last 120 ka. The analysis of these simulations elucidates the relative importance of the biogeophysical versus biogeochemical terrestrial biosphere interactions with climate. We find that the biogeophysical effects of vegetation account for up to an additional -0.91°C
- 10 global mean cooling, with regional cooling as large as -5° C, but with considerable variability across the glacial-interglacial cycle. By comparison, while opposite in sign, our model estimates of the biogeochemical impacts are substantially smaller in magnitude. Offline simulations show a maximum of +0.33°C warming due an increase of 25 ppm above our (pre-industrial) baseline atmospheric CO₂ mixing ratio. In contrast to shorter (century) time-scale projections of future terrestrial biosphere response where direct and indirect responses may at times, cancel out, we find that the biogeophysical effects consistently
- 15 and strongly dominate the biogeochemical effect over the inter-glacial cycle. In addition, depending on the assumptions about soil carbon under ice-sheets and sea level rise, we find a range in terrestrial carbon storage change from a reduction in LGM carbon storage of -440 PgC, to a gain of +37 PgC, though we consider the negative part of the range more likely. We suggest that prevailing uncertainties allow for only a small net transfer of carbon between terrestrial biosphere and ocean/atmosphere implying that explaining the observed CO₂ ice core record could be rather simpler than previously thought.

20 1 Introduction

Terrestrial vegetation interacts with the climate in complex ways, both responding to and impacting climate conditions and hence creating an important feedback in the Earth system (e.g., Claussen, 2009; Davies-Barnard et al., 2014b; Harrison and Prentice, 2003; Jahn et al., 2005; Matthews et al., 2003; Pongratz et al., 2010). The influence of the terrestrial biosphere on

climate occurs in two distinct ways. Firstly, there are a number of biogeophysical mechanisms such as changes in albedo or evapotranspiration that provide a direct physical influence on surface climate via changes in net solar radiation transfer, infrared loss, roughness length, latent heat loss, and less directly, via changes in moisture exchange and hence transport. Climate feedbacks driven by these changes in terrestrial vegetation have been hypothesised to be partially responsible for some

- 5 of the major past climate states (e.g., Bradshaw et al., 2015; Claussen et al., 2006; Crucifix and Loutre, 2002; de Noblet et al., 1996; Zhou et al., 2012), with many studies particularly focussing on the biogeophysical effects at the last glacial maximum (LGM) (e.g., Hopcroft and Valdes, 2014; Jahn et al., 2005; Kageyama et al., 2012; O'ishi and Abe-Ouchi, 2013). The second way in which the terrestrial biosphere can influence climate is via variations in the carbon stored in vegetation and soil. This is a crucial component for understanding changes in the carbon cycle through the last glacial-interglacial cycle (Montenegro
- 10 et al., 2006) and numerous attempts have been made to estimate the total carbon storage using a range of methods, such as inferences from marine and terrestrial carbon isotopes (e.g., Shackleton et al., 1977; Bird et al., 1994), databases of pollen (e.g., Adams and Faure, 1998; Crowley, 1995), and simple and complex modelling (e.g., Prentice et al., 1993; Kaplan et al., 2002; Köhler and Fischer, 2004; Brovkin et al., 2012; O'ishi and Abe-Ouchi, 2013). The resulting range of carbon storage change estimates is from a few hundred to about 1000 PgC (Ciais et al., 2012). One could add to this changes in the weathering of
- soil minerals and hence CO_2 uptake from the atmosphere, and nutrient, particularly phosphate, supply to the ocean and hence changes in in the ocean productivity. For simplicity, we will not address these further here (except to include a basic silicate weathering feedback in our model analysis of the impacts of terrestrial carbon storage change).

Simulations of future vegetation changes show that the biogeochemical aspect can globally be around the same magnitude as the biogeophysical effects (e.g., Davies-Barnard et al., 2014b) meaning that there is uncertainty even in the sign of the

net feedback with climate change. Both biogeophysical and biogeochemical effects likely also play an important role in past climate change and potentially the same fundamental uncertainty in the sign of the climate feedback might arise. However, model simulations have generally focussed on either the biogeophysical impacts of vegetation changes (e.g., Bradshaw et al., 2015; Claussen et al., 2006; Jahn et al., 2005; O'ishi and Abe-Ouchi, 2013; Shellito and Sloan, 2006) or biogeochemical impacts (e.g., Kaplan et al., 2002; Ciais et al., 2012) and the question of the overall feedback on climate rarely addressed, although Claussen (2009) argues that the net effect at the LGM is dominated by the biogeophysical effects.

One of the few examples where both have been combined and the net effects of vegetation on past climate estimated over long time periods, is Brovkin et al. (2012). They used an earth system model of intermediate complexity (EMIC) to suggest that the net effect of vegetation is to decrease global temperatures during the last glacial-interglacial cycle. But the model used is relatively coarse in resolution (10° in latitude and 51° in longitude) and reduced in physical process complexity. This may be

30 important because of the local and spatially heterogeneous nature of biogeophysical effects and depending on the location of the forest, the biogeophysical and biogeochemical effects of forest change can be very different (Bonan, 2008). For this reason, fully coupled General Circulation Models (GCM) are commonly used in quantifying future climate changes to vegetated land surface (Brovkin et al., 2013a, b; Davies-Barnard et al., 2015; Davin and de Noblet-Ducoudré, 2010). The importance of considering both biogeophysical and carbon cycle impacts together at finer scale when assessing the climate impacts of

35 vegetation is illustrated by work quantifying the climate impacts of forest changes. Studies have found that deforestation

would cause local high latitude cooling (Betts, 2000), global warming (Davin and de Noblet-Ducoudré, 2010), or even slight global cooling (Davies-Barnard et al., 2014b, a). These outcomes are not predictable from looking at the biogeophysics or terrestrial biogeochemistry alone at coarse resolutions.

Here we present the first model analysis using a fully-coupled dynamic atmosphere-ocean-vegetation GCM over the last
120 ka that quantifies the net effect of vegetation on climate. (A prior study – Singarayer and Valdes (2010) – did not have dynamic vegetation and hence could not directly evaluate the biogeophysical effects.) We separate the biogeophysical and biogeochemical effects of vegetation to understand the overall climate effect of vegetation over the last glacial cycle. We show that over the whole period the biogeophysical is the dominant effect, and that the biogeochemical impacts may have a lower possible range than typically estimated. We also highlight how the temporal scale affects the net impact of terrestrial biosphere
changes.

2 Methods

We use the GCM HadCM3 to run a series of simulations with and without dynamic vegetation to provide the biogeophysical changes and the land carbon changes. To look at the climate impact of those vegetation carbon changes, we then use the GCM terrestrial carbon changes as an input to the EMIC cGENIE to calculate the resulting change in atmospheric CO_2 and global termeasture

15 temperature.

For future climate changes studies, the response of atmospheric CO_2 concentrations (and hence climate) to changes in terrestrial carbon storage can be calculated using the Transient Response to Cumulative Emissions (TRCE) approach (Gillett et al., 2013), which demonstrated proportionality between carbon emissions and temperature rise (Goodwin et al., 2015). We include these estimates, for completeness. However, this approach is only valid for relatively rapid changes. On the longer time

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scales of glacial-interglacial change, we need to take into account the full changes in ocean carbon chemistry and including the interactions of ocean and atmosphere with the solid Earth (e.g. weathering). To do this, we employ the 'cGENIE' Earth system Model of Intermediate Complexity (EMIC).

2.1 Climate Model Description

The GCM used in the simulations in this study is the UK Met Office Hadley Centre's HadCM3B-M2.1a and HadCM3B-M2.1a
 M2.1aD (Valdes et al., 2017). Though not from the latest generation of climate models, HadCM3 remains an extensively used model for many research applications around the world due to its computational efficiency, which means that long integrations and many ensemble members can be run.

HadCM3 is a three dimensional, fully coupled, fully dynamic ocean, non-flux adjusted global climate model (Collins et al., 2001). The atmosphere component, HadAM3, has a cartesian grid with a horizontal resolution of 2.5° x 3.75° , 19 vertical

30 levels and a time step of 30 minutes (Pope et al., 2000). The ocean and sea-ice component has the same horizontal resolution as the atmosphere, with 20 vertical ocean levels.

The land surface scheme used for the atmosphere component of HadCM3 is the Met Office Surface Exchange Scheme, MOSES2.1 (Gregory et al., 1994; Cox et al., 1999). MOSES can also use an additional vegetation and terrestrial carbon model, TRIFFID (Top-down Representation of Interactive Foliage and Flora Including Dynamics) (Cox, 2001; Cox et al., 1998). TRIFFID predicts the vegetation based on plant functional types using a competitive, hierarchical model. TRIFFID has two

5 modes, equilibrium mode, which quickly brings the vegetation cover into equilibrium by running fifty years of TRIFFID for each five years of the climate model run, and dynamic, which runs TRIFFID every ten days. TRIFFID and MOSES have nine land surface types, five of which are vegetation: broadleaf trees, needle leaf trees, shrubs, C_3 grasses and C_4 grasses. These are known as plant functional types (PFTs) and have different leaf area index limits and other phenological differences in the model. Soil moisture in the model is represented on 4 layers of thicknesses (measured from the top) of 0.1, 0.25, 0.65 and 2.5

10 m (Essery et al., 2001).

The soil carbon is a single pool, increased by litterfall and decreased by respiration (Cox, 2001; Cox et al., 1998). The soil respiration is controlled by moisture and temperature and returns carbon dioxide to the atmosphere unless, as is the case here, the atmospheric carbon dioxide is fixed. The litterfall is an area-weight sum of the litterfall of the five PFTs in each gridcell (Cox, 2001; Cox et al., 1998). There is no permafrost component in the model, and soil in frozen regions is treated the same as

15 in any other.

2.1.1 GCM Simulations and Experimental Methodology

The simulations used here are revised versions of those described in Singarayer and Valdes (2010), who used HadCM3 version HadCM3-M1, which has an older surface scheme (MOSES1) than the MOSES2.1 used here, and no dynamic vegetation. Two sets of 62 simulations were performed, covering the time period 120 - 0 ka BP:

- The first set of 62 simulations used TRIFFID to predict vegetation changes. Each individual simulation was initialised from the previous MOSES1 simulations (which were run for 600 years) and were then run for a further 300 years with 'equilibrium' TRIFFID and a final 300 years with fully dynamic vegetation. This set will be referred to as the Dynamic set.
 - A second set of simulations uses static vegetation based on the pre-industrial simulation of the dynamic set (extrapolated
- 25

to new land areas using a simple nearest neighbour algorithm). They are otherwise identical to the Dynamic set (see details below). These will be referred to as the Static set.

The differences between Dynamic and Static allows us to evaluate the biogeophysical and biogeochemical responses of terrestrial carbon cycle change.

Both sets of simulations are forced with the same changes in orbit, greenhouse gases (CO₂, CH₄, and N₂O) and ice sheets,
as in Singarayer and Valdes (2010) except that we use a revised ice sheets extent and elevation, as discussed in Singarayer et al. (2011).

We have also added a parameterisation of water transport from ocean to ice sheet in order to ensure that ocean salinity is conserved during each simulation. In the normal configuration of HadCM3, salinity is conserved by the numerical scheme but

water that accumulates as snow on ice sheets is not interactively considered. A predefined (spatially varying) flux of water is prescribed into the model which minimises the salinity drift for the pre-industrial simulation but this is not normally changed for other time periods. In our new parameterisation, we continue to add the predefined flux but also add an additional flux which is spatially uniform but temporally variable to ensure that the volume integral ocean salinity is relaxed back to its initial value, with a relaxation time scale of 10 years. This prevents any spurious long term drifts in ocean salinity.

Note that this model does not have a closed carbon cycle. There is no representation of carbon in the ocean and terrestrial carbon changes do not feedback to the atmosphere (since the greenhouse gas forcings are prescribed). However, the carbon that would have returned to the atmosphere can be inferred from the change in the carbon stores in the soil and vegetation, allowing the biogeochemical impact of vegetation to be understood, as well as the biogeophysical. From the 22 ka to pre-industrial,

10 simulations are run for every 1000 years. From 80 ka to 22 ka, simulations are run for every 2000 years. For 120 ka to 80 ka, simulations are run for every 4000 years. (See grey points in Figure 2 for a representation of the temporal distribution of the 62 simulations.) Reported final climatologies are based on the last 30 years of each simulation.

2.2 EMIC Description

The cGENIE Earth system model is used to calculate the impacts on atmospheric CO₂ over the glacial cycle and hence make
a time-varying estimate of the contribution of biogeochemical changes to glacial-interglacial climate change. The model is based around a fast energy-balance based atmosphere model coupled to a 3D ocean circulation component and dynamic-thermodynamic sea-ice (Edwards and Marsh, 2005), plus representations of ocean-atmosphere (Ridgwell and Hargreaves, 2007), ocean-sediment (Ridgwell et al., 2007), and atmosphere-land (terrestrial weathering) (Colbourn et al., 2013) carbon cycling. As employed here: the non-seasonally forced ocean has 8-levels and the configuration and selection of model parameterisations and parameter values is identical to that described in Lord et al. (2016). These choices are made to minimise

experiment run-time and provide maximum traceability (to a previously used and in-depth analysed configuration), respectively.

2.3 cGENIE Carbon Cycle Simulations

The evolution of terrestrial carbon storage simulated by HadCM3 from 120 ka to pre-industrial was used to derive a forcing for cGENIE. In this, we created a continuous time-series of the carbon flux from the terrestrial biosphere by calculating the difference in carbon storage calculated at the end of each HadCM3 time-slice and then assuming a linear interpolation between these points. For the 'Full' simulations, cGENIE was then run for 120 ka using this forcing and starting from a fully spun-up state of global carbon cycling including an initial balance between the rate of silicate rock weathering and volcanic CO₂ outgassing (see Lord et al. (2016) for details). For the 'Carbonate' simulations, the model was run with just carbonate compensation only, as per Ridgwell and Hargreaves (2007); For the 'Closed' simulations, the carbon remains in the atmosphere. Both the resulting history of atmospheric CO₂ as well as annual mean global surface air temperature were

extracted and calculated as anomalies relative to the late Holocene (pre-industrial).

Using the 'Full' setup, cGENIE simulations were run using four different carbon estimations from the GCM simulations (see Table 1). For 'Carbonate', 'AirOcean' and 'Closed' a simulation was run with the GCI_ELE carbon scenario (see Table 1). Therefore seven transient cGENIE simulations were run in total.

It should be noted that we do not attempt to change the boundary conditions required by the cGENIE model dynamically 5 through the glacial-interglacial cycles, namely: orbital parameters, planetary albedo, sea-level (and ocean salinity). These are instead kept fixed at modern (following Lord et al. (2016).) Hence, changes in the sensitivity of atmospheric CO_2 to unit CO_2 input (or removal) will not be accounted for. We expect such an effect to exist due to e.g. the dependence of the Revelle factor (the sensitivity of dissolved $CO_{2(aq)}$ to changes in total dissolved inorganic carbon (Zeebe et al., 1999)) on both (ocean surface) temperature and atmospheric pCO_2 , changes in ocean circulation and the efficiency of the biological pump, and changes in

- 10 the carbonate buffering of ocean chemistry. Some of these factors could in theory be imposed (e.g. changes in ocean surface temperatures), but others would require the glacial-interglacial dynamics in both ocean circulation and marine carbon cycling to be sufficiently accurately represented in the model. The latter is far beyond what the current state of understanding of glacial-interglacial global carbon cycling allows for Kohfeld and Ridgwell (2009). Hence our assumption of fixed late Holocene boundary conditions will impart a small bias in our estimates of the atmospheric CO₂ response, but not one that would affect
- 15 our overall conclusions.

In addition, in making estimates of the mean global air surface temperature change corresponding to the projected change in atmospheric pCO_2 in cGENIE, it is important to also note that the climate sensitivity is effectively prescribed (Edwards and Marsh, 2005). In the Lord et al. (2016) configured used here, only sea-ice cover, via its associated albedo, can provide feedback on climate. In the absence of a dynamical atmosphere, glacial-interglacial changes in climate sensitivity due to

20 changes in atmospheric circulation and clouds are not possible. Nor do we account for the possible influences of changes in total land surface area (from sea-level change) or vegetation cover and distribution. However, the assumption of an effectively fixed climate sensitivity across the glacial-interglacial cycle is unlikely to impart significant bias or unduly affect our overall conclusions.

3 Results

25 3.1 Results: Vegetation Dynamics

The changes in climate over time affects the vegetation cover in the Dynamic simulations (shown in Figure 1). In general, cooling leads to an equator-ward shift in vegetation, as the high latitudes become covered in ice or otherwise inhospitable for significant quantities of vegetation. There is also exposure of continental shelves, providing potential for vegetation increases. At the last glacial maximum (LGM) at 21 ka, we can see needleleaf trees and shrubs giving way to very low productivity grasses

30 in the high latitudes. However, because of the small number of PFTs (five) in this model, the shifts may be underestimated, as each PFT represents a wide range of vegetation types. The shrubs and trees do not have a significant presence in northern Europe after 100 ka until the climate ameliorates into the Holocene. It is the vegetation changes shown in Figure 1, and their associated soil changes, that drive the climate feedbacks and other changes described hereafter.

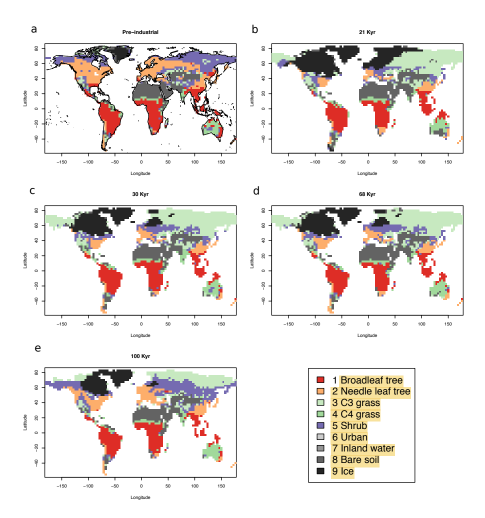


Figure 1. Dominant plant functional type (PFT) for some time periods of interest in the 120 ka covered by the simulations. a) Pre-Industrial, b) 21 ka, c) 30 ka d) 68 ka, e) 100 ka. Note that the dominant PFT is calculated as the land cover with the highest proportion of cover, compared to the other land surface types, and does not necessarily indicate the highest or a significant amount of net primary productivity (NPP).

Hoogakker et al. (2016) have shown that HadCM3 broadly reproduces the known changes in vegetation across the glacialinterglacial cycle. Hoogakker et al. (2016) uses HadCM3B-M1 (without dynamic vegetation), then uses the climate to drive BIOME4. The climate is very similar between HadCM3B-M1 and HadCM3B-M2.1a used here. In Hoogakker et al. (2016) they ran an offline vegetation model, BIOME4, driven by the climate anomalies from HadCM3. Our results from TRIFFID

5 are consistent with the relative changes although, since TRIFFID uses the actual climate from the models, the vegetation can have biases (e.g. Australia has a tendency to be too wet in HadCM3 in the present day and hence the coupled model has too much vegetation in this region). However, during glacial times there is a decrease in biomass, consistent with Hoogakker et al. (2016).

Comparison with the BIOME6000 Mega-Biome maps for LGM (Pickett et al., 2004; Prentice and Jolly, 2000; Bigelow et al., 2003; Harrison et al., 2001) dataset shows general agreement. The model has considerable expansion of grasses in

5 Eurasia where BIOME6000 has grassland and dry shrubland. Broadly speaking, North America shows little change from the mid Holocene to LGM. One key weakness of the model is in western Europe, where BIOME6000 shows grassland and dry shrubland, whereas the model has shrubs and needleleaf trees. Similarly, assessment of the PI vegetation cover of HadCM3 by Valdes et al. (2017) shows good agreement with reconstructions of 1800 vegetation.

The forest extent in the tropics at the LGM is similar to PI (see SI Figure 9 for shifts in vegetation at 21 ka). This is supported by pollen and other data (Maslin et al., 2012; Anhuf et al., 2006), and modelling (Cowling et al., 2001) which find that the although there is diminished tropical forest, there is still substantial tree cover at the LGM and little sign of widespread grasslands. Because of the PFT (rather than biome) approach of TRIFFID, and the limited number of PFTs, it's difficult to be sure whether trees in the tropics are a tropical rainforest at the LGM, because they equally could be temperate forest.

3.2 Results: Biogeophysical Feedbacks

- 15 The biogeophysical impacts of vegetation are calculated by subtracting the Dynamic simulations from the corresponding Static simulations. We find that vegetation is acting as a positive feedback to the climate, enhancing the cooling (Figure 2a). Broadly, the Static and Dynamic simulations both agree with an approximation of global temperature over the whole period (the EPICA dataset halved) (Figure 2a). The Static set generally do better in 70 ka to 10 ka, whereas the Dynamic set are closer to the EPICA data in the period 110 to 70 ka. The biogeophysical differences between the Static and Dynamic sets alter global, annual mean surface temperature by as much as -0.91°C (see Figure 2b). Regionally this temperature cooling is up to 5°C
- (Figure 3).

The albedo changes are in the same location as the vegetation carbon changes, and the main temperature changes (see Figure 4 and 3). These temperature differences are mainly driven by reductions of tree cover and its replacement with bare soil or grasses, which is a result of the vegetation dynamics in the model (see Figure 1 and Figure 4). Trees have a lower albedo, and

- when they are replaced by higher albedo grasses, there is a cooling effect. The change in tree fraction between the Static and Dynamic sets is a good predictor of the temperature changes ($r^2 = 0.79$ using a linear model of the global temperature and tree anomalies). This is exacerbated by the presence of snow cover as the snow covered visible and near infrared albedo of grasses, shrubs and bare soil is higher than that of trees (Essery et al., 2001). Therefore when trees are replaced by grasses where there is snow cover for part of the year, there is a larger change in albedo than where there is no snow cover. Thus the albedo changes
- 30 can be seen mainly where a change between trees and grasses occurs in an area with snow cover (see Figure 1 and Figure 5). The exact contribution of the snow as opposed to the no-snow albedo is difficult to disentangle, but the influence of this effect is well established (Betts, 2000).

The land surface albedo changes caused by the vegetation have an even stronger correlation with these biogeophysical temperature changes ($r^2 = 0.86$). However, we can see that although the forcing is land based (the dynamic vegetation),

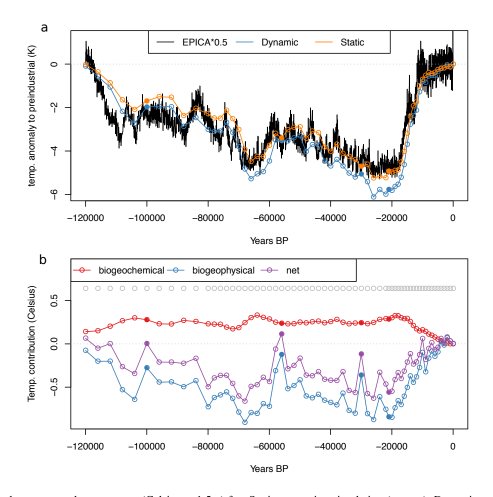


Figure 2. a) Global mean annual temperature (Celsius at 1.5m) for: Static vegetation simulation (orange); Dynamic vegetation (blue); and the EPICA core inferred temperature data (black), halved (to give an approximation of global temperature over the whole period). Time periods of particular interest are highlighted as filled points: 21, 30, 56, 68 and 100 ka. b) Temperature anomaly over time of Dynamic - Static simulations for: biogeochemical temperature effects of the vegetation change, calculated with GENIE, averaged to the same temporal resolution as the HadCM3 simulations (red); biogeophysical temperature effects of vegetation change (blue); the net (biogeophysical and biogeochemical) effect of vegetation on temperature (purple). Grey points show the time points of the HadCM3 simulations.

significant changes occur in the ocean (see Figure 4 and Figure 3) that drive the resulting temperature changes. Ocean only surface albedo anomaly as a determinant of global temperature anomaly has an r^2 of 0.95 - lower only than the r^2 of the global (land and ocean) surface albedo of 0.96. By comparison, the r^2 of the latent heat anomaly as a predictor of temperature anomaly is lower for land, ocean, and global than surface albedo (0.70, 0.93 and 0.91 respectively).

The other parts of the energy balance, in particular the latent heat, sensible heat, and the net shortwave radiation, do not have such a clear relationship with the temperature change (see SI Figure 10 and compare to Figure 3).

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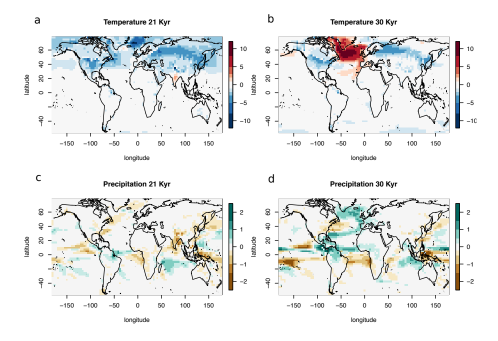


Figure 3. Anomaly of (a and b) temperature at 1.5m (Celsius); and (c and d) precipitation (mm day⁻¹); between the Dynamic Vegetation simulation and the equivalent Static Vegetation Simulation. For (a and c) 21 ka and (b and d) 30 ka. The pattern of reduced surface albedo at 30 ka is similar to the pattern at 56 ka and 100 ka.

Although the biogeophysical changes cause cooling, there are some minima of biogeophysical temperature change seen at 30 ka, 56 ka and 100 ka (Figure 2, filled symbols). These minima have an oceanic source and are caused by vegetation interacting with thermohaline circulation changes. In our new simulations we account for the net transport of water from ocean to the ice sheets by a parameterisation that instantaneously balances any net accumulation of water on ice. This parameterisation

- 5 results in fresher ocean conditions during times of precession driven N. Hemisphere summer insolation highs (less water is being used to build the ice sheets). The instantaneous nature of the parameterisation is physically unrealistic but reductions in accumulation and an increase in ablation during precession highs has been seen in fully coupled climate-icesheet EMIC simulations (e.g. Ganopolski et al, 2010). During weaker accumulation periods, the parameterisation results in a freshening of ocean surface waters and a reduction in AMOC strength from 16Sv to 10 -12Sv.
- 10 Superimposed upon this general behaviour, the addition of interactive vegetation generally does not change the AMOC strength. However, at times of weak AMOC, small changes in runoff and temperature are sufficient to cause some changes in the response. For instance, in the static vegetation simulations there is a relatively weak AMOC in the simulations for 60 ka, 58 ka, and 56 ka. In the interactive vegetation simulations, the weakened AMOC only occurs at 60 ka. Thus at 60 ka the changes in climate are fairly typical of preceding times but at 58 ka and 56 ka there is a substantial difference between the static and
- 15 dynamic vegetation simulations. The cause for this difference is associated with a combination of reduced runoff into the N. Atlantic (principally from changes in land surface in N. America) and colder temperatures, both of which act to stabilise the

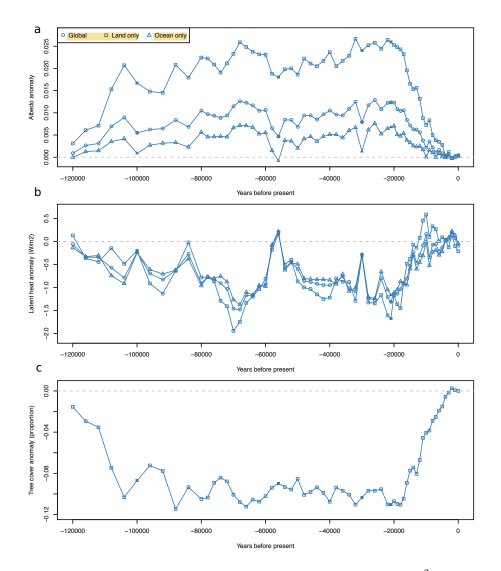


Figure 4. Mean annual anomaly Dynamic - Static simulations for: a) surface albedo b) latent heat (W/m^2) and c) tree cover (proportion of land area).

AMOC in all three periods but it is sufficient to prevent the AMOC weakening in the 58 and 56 ka simulations. This threshold like behaviour of the AMOC is almost certainly highly model dependent and hence the result is not robust.

The regional patterns of cooling also temporarily affect the precipitation regime (see Figure 3). This appears to be related to the AMOC weakening. There are some suggestions of similar relationships between the increases in precipitation and the

5 terrestrial changes to previous studies (Gedney and Valdes, 2000; Singarayer et al., 2009). Similar to the temperature changes, it is unclear how model-specific these changes are.

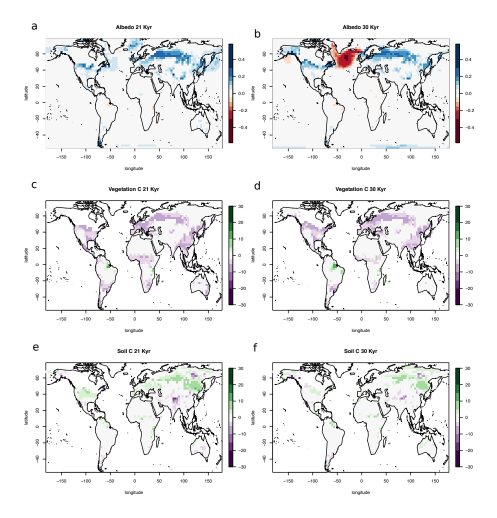


Figure 5. Mean annual anomaly of: a) and b), surface albedo (unitless); c) and d), vegetation carbon (kg C m⁻²); and e) and f), soil carbon (kg C m⁻²), between the Dynamic Vegetation simulation and the equivalent Static Vegetation simulation. For 21 ka (a, c, and e) and 30 ka (b, d, and f). The pattern of reduced surface albedo at 30 ka is similar to the pattern at 56 ka and 100 ka. Since in the Static simulations the carbon remains at PI levels, figures c - f also represent the anomaly to PI. The ice-sheets are excluded from these plots. For the carbon under ice-sheets, see Figure 11.

3.3 Results: Biogeochemistry

We now calculate the total change in terrestrial carbon stores in the HadCM3 simulations. We consider scenarios of terrestrial carbon change with combinations of including or excluding uncertain aspects of the carbon cycle, specifically depending on the fate of soil carbon under ice and the changes related to the expansion of land. Zeng (2003) suggested that the soil and

5 vegetation carbon formed during the warm last interglacial could simply get covered by ice and is stored there, rather than being released into the rest of the system as is typically assumed in past estimates. Similarly, the amount of carbon stored on

Name	Carbon storage Scenarios	Soil C change (PgC)	Vegetation C change (PgC)	Total C change (PgC)
GCI_ELE	Carbon under ice sheets re-	-145	-295	-440
	leased to atmosphere. No car-			
	bon on expanded land area.			
GCE_ELE	Carbon under ice sheets stored	+77	-222	-146
	under the ice. No carbon on ex-			
	panded land area.			
GCE_ELI	Carbon under ice sheets stored	+211	-173	+37
	under the ice. Modelled carbon			
	storage on new land included.			
GCI_ELI	Carbon under ice sheets re-	-11	-246	-257
	leased to atmosphere. Modelled			
	carbon storage on new land in-			
	cluded.			

Table 1. Terrestrial carbon changes from PI to LGM. For storage values at the LGM, see Appendix Table 1.

newly emerged land is also uncertain as it depends on both the area of emergent land and the surface properties. Therefore we calculate the changes in soil and vegetation carbon from these various sources. In Table 1 we focus on the changes between pre-industrial and LGM, which corresponds to the largest overall change through the glacial-interglacial cycle.

In the model, 222 PgC of soil carbon and 73 PgC of vegetation carbon is associated with areas covered with ice at the LGM (see Figure 8). Similarly, 134 PgC of soil carbon and 49 PgC of vegetation carbon is associated with new land. The resulting range of total carbon storage is large, from a loss of 440 PgC at the LGM (no carbon stored under new ice sheets with all being released to ocean-atmosphere, and no build-up of carbon on new land surface) to a possible small increase of carbon (if carbon is stored under the new ice sheets and there is no carbon storage on new land).

In reality, glacial systems are known to export carbon in a highly labile form (Lawson et al., 2014), erode soil and bedrock creating major landscape changes, and release large amounts of methane when they retreat (Wadham et al., 2012). Although the conversion of this terrestrial carbon to atmospheric carbon may be through riverine or oceanic systems, it seems likely it would return to the atmosphere within the time periods we consider. We therefore use this largest scenario as a conservative option for our main analysis.

The other major change to soil carbon in the model is newly exposed land, which is revealed when the water in the ice-sheets causes lower sea levels (see Figures 1 and 5). For the new land we use a nearest-neighbour interpolation of basic soil properties (e.g. water holding capacity etc.) and the model is run for sufficient length of time for the soil and vegetation carbon to reach equilibrium.

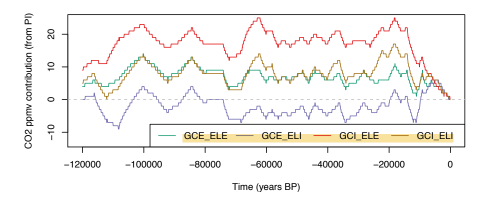


Figure 6. Contribution of terrestrial biosphere carbon emissions to atmospheric CO_2 . Run with the cGENIE 'Full' configuration and normalised to pre-industrial CO_2 levels. The four scenarios are as detailed in Table 1.

This estimate of carbon on expanded lands also has uncertainties. We have reasonable confidence in the sea level estimates and consequent change in land area, but it is much more uncertain the state of the land and whether carbon could accumulate on it. For instance, in our simulations the East Siberia ice-sheet is absent (see Figure 1), whereas many other ice-sheet reconstructions include it (e.g., Niessen et al., 2013). The area of the ice sheet alone accounts for an average of 56 PgC soil carbon in these simulations when it is absent. But soil carbon takes a long time to accumulate, especially with low NPP and vegetation carbon storage averages just 0.5 PgC over all the expanded lands.

If exposed land carbon was included and glacial land soil carbon excluded, the terrestrial carbon is a gain from PI to LGM of +37 PgC (see Table 1). However, as discussed above, we would argue that excluding glacial land soil carbon change is probably unreasonable. Most previous studies have also assumed that all carbon under ice is removed. If we include the loss of carbon, then the range in total amount of terrestrial carbon lost in this model between pre-industrial (PI) and the LGM at 21 ka is -440 to -257 PgC.

The change in terrestrial carbon found in our simulations contributes to atmospheric carbon dioxide change. Using the cGENIE model to approximate the carbon uptake by the ocean we therefore calculate the atmospheric carbon dioxide change (see methods and Figure 6).

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Selecting the largest change in carbon storage (-440 PgC at the LGM, including glacial soil carbon changes and excluding expanded lands) the results suggest a peak contribution compared to pre-industrial CO_2 of 25 ppm CO_2 (Figure 6). In all scenarios except GCE_ELI, the terrestrial carbon contribution to atmospheric CO_2 acts as a negative feedback to the climate, dampening the effect of other climate forcings, including the net contribution of the terrestrial biosphere (Figure 2b).

Within cGENIE, the change in atmospheric CO₂ produces a warming at the LGM of 0.29°C (equivalent to a climate sensitivity of around 2 Wm-² °C⁻¹, see Figure 2b). This is much smaller than the biogeophysical contribution of -0.84 °C. It is also much less variable. For most of the glacial period, from 100 ka to 20 ka, the implied biogeochemical warming is around 0.26 °C (Figure 2b). This results in the dominance of the biogeophysical impacts over biogeochemical feedbacks.

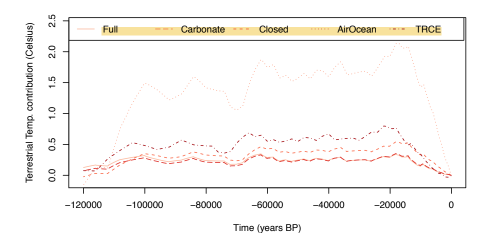


Figure 7. Temperature changes resulting from the same terrestrial carbon emissions scenario (GCI_ELE) with different model set-up for cGENIE and for the TRCE of HadCM3. cGENIE simulations were: 'Full' with silicate weathering feedback and just carbonate compensation, as Lord et al. (2016); 'Carbonate' with just carbonate compensation only, as Ridgwell and Hargreaves (2007); 'Closed' with no weathering or sediment response, and hence is just ocean-atmosphere repartitioning; 'AirOcean' where the carbon remains in the atmosphere. 'TRCE' is the simple calculation of the TRCE of HadCM3 (taken from Matthews et al. (2009)) for the same carbon inputs into the atmosphere as used for the cGENIE simulations. Note, we include the TRCE for completeness, but it is not a cGENIE simulation (see methods).

However, with different earth system processes included, the biogeochemical effects vary substantially (see Figure 7). In the simulations discussed above, silicate and carbonate weathering are both included and this results in the lowest temperature change from the same carbon emissions. The temperature contribution at the LGM increases (from Full, 0.29°C); as the silicate weathering is excluded (Carbonate, 0.30°C); all weathering is excluded (Closed, 0.47°C); a decadal to millennial scale carbon uptake is used (TRCE, 0.86°C); and if all carbon remains in the atmosphere (AirOcean, 1.92°C). Note that the TRCE as shown above includes the terrestrial biosphere as a sink, so will slightly overestimate how much carbon will be removed from the atmosphere when the source is the natural vegetation. Comparing these values to the biogeophysical terrestrial effect in Figure 2b, we can see that the shorter the timescale, the more likely biogeochemical terrestrial processes will dominate as it weakens over time. On longer timescales the biogeophysics dominates because the scale of the effect doesn't diminish over time relative

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4 Discussion

The biogeophysical results found here broadly concur with comparable model studies of past vegetation biogeophysics. Claussen et al. (2006) found the biogeophysical contribution of vegetation to LGM cooling of around 1°C in the northern hemisphere, whereas Jahn et al. (2005) found around -0.6°C, and up to 2 °C locally. Our result of -0.84°C is in the middle of the other LGM studies.

The dominance of the biogeophysical effects found here is contrary to the results found for short time scale problems, which find that biogeochemistry tends to be comparable in magnitude to biogeophysical effects (e.g., Davies-Barnard et al., 2014b; Pongratz et al., 2010). This is because the centennial simulations have a stronger biogeochemical effect since the transient response to cumulative emissions is stronger than the equilibrium response. In climate simulations up to around a century long,

- 5 more carbon tends to remain in the atmosphere. This makes a strong warming effect that is approximately linearly related to the amount of greenhouse gas emissions (Matthews et al., 2009; Gillett et al., 2013). The transient response to cumulative emissions (TRCE) accounts for the uptake of atmospheric carbon by the ocean and terrestrial biosphere, but only on short timescales. The uptake of atmospheric carbon by the ocean requires hundreds or thousands of years, and is slower when the increase of carbon into the system is small and staggered (Lord et al., 2016). However, the simulations we use are on a millennial timescale,
- 10 allowing much of the carbon to be taken up by the ocean (Lord et al., 2016). From a climate sensitivity point of view, this means that on shorter timescales, the effects of dynamic vegetation can cancel each other out. This provides some rationale for the fact that dynamic vegetation has been generally not included in the majority of state-of-the-art earth system models used in CMIP5, as it doesn't significantly affect the climate sensitivity. At longer time scales, it is more important to include dynamic vegetation, as without the positive feedback of the biogeophysical effects, the climate sensitivity would be under estimated.
- For the biogeochemical effects of the terrestrial biosphere, previous estimates of carbon stocks on exposed continental shelves based on models are between 112 to 323 PgC at the LGM (Montenegro et al., 2006). The comparable number in this simulation is 183 PgC, which is on the lower end of the wide range of other models. However, it has good agreement with the vegetation reconstruction (not model) values by Montenegro et al. (2006) of 182 to 220 PgC.
- The LGM terrestrial carbon change here is -440 to +37 PgC, including a zero contribution of terrestrial carbon. This is smaller than the values of -900 to -400 PgC range reviewed by Kohfeld and Ridgwell (2009). More recent modelling studies are also somewhat larger than our estimate range, such as -500 PgC (Brovkin et al., 2012), -597 PgC (O'ishi and Abe-Ouchi, 2013), and (Prentice et al., 2011) -550 to -694 PgC. However, recent inventory and isotope approaches are closer to our range of values, e.g. -378 ± 88 PgC (Menviel et al., 2017) and -330 PgC (Ciais et al., 2012).
- For present day, Hugelius et al. (2014) shows around 75 100 kg C m² far north Siberia, 20 40 further south. Far northern
 Canada is much more heterogeneous, with values from 20 150 kg C m². The modelled PI values are on the low side, and much more homogeneous, around 15 20 kg C m², but is similar to Hugelius et al. (2014) in that it shows far north America to be less consistent, with some higher areas of 35 40kg C m² in the far north. (See Supplementary Figure 11 of the loss of soil carbon under ice-sheets at LGM.) What this suggests is that while on the correct order of magnitude, the model has a very modest amount of soil carbon that could be considered permafrost. Therefore, we think it's reasonable to include this low
- 30 estimate of soil carbon in the uncertainties.

The soil carbon change under ice-sheets between PI and LGM is modelled as ~220 PgC. Extrapolating from a comparison with Hugelius et al. (2014), this might be a third too little. If the true value were ~330 PgC, this would make the total C change PI to LGM 550 PgC, more in line with some previous model estimates. It would affect the global mean annual biogeochemical contribution by ~0.1 K. This would mean the net effect of vegetation was closer to zero, but the biogeophysical effect would still deminente

³⁵ still dominate.

However, the exact size of the terrestrial carbon emissions is uncertain. Other carbon stores not accounted for here are potentially important, for example methane during sea level rises or changes to the wetlands in the tropics. Modelling studies that look at wetlands at the LGM suggest that although the wetland area is larger, but the methane emissions are lower compared to modern day (Kaplan, 2002). However, paleohydrological data indicates a drying in the African tropics (Gasse, 2000).

- 5 Our model does not have a process based permafrost or wetlands component, and therefore the changes in methane are not accounted for. This is a particular limitation when considering the carbon stored in deep permafrost soils in Northern peatlands. Saito et al. (2013) show that, based on the temperature changes, there is a substantial expansion of permafrost area during glacial times but cannot estimate any changes in carbon storage. Zimov et al. (2006, 2009) have argued that permafrost storage could be a major source of carbon through the deglaciation, and Ciais et al. (2012) argue that there was a large extra pool of
- 10 inert carbon at the LGM. Similarly, Köhler et al. (2014) have argued that large amounts of carbon were locked into permafrost which were then released rapidly at the Bolling-Allerod.

Research has also suggested that waterlogging and flooding as sea level rises during the Holocene could cause rapid anaerobic decomposition of vegetation, causing methane emissions (Ridgwell et al., 2012). This could account for emissions of as much as 25 PgC for 10 meters sea level rise (ibid). Since our simulations do not account for methane or this effect of inunda-

15 tion, it is likely it there is a slight underestimation of equivalent CO_2 effect of the carbon emissions (as methane is a stronger greenhouse gas than carbon dioxide).

The impacts are mainly determined by the vegetation shifts the DGVM simulates. Each grid-box has the potential for 5 PFTs, but generally the Lotka-Volterra equations used in TRIFFID mean that the grid-box is dominated by one PFT. The small number that means the range within each PFT is relatively large. Therefore the model probably underestimates the effects of

- 20 small perturbations in climate, as the large definition of the PFTs allows the PFT to remain the same. Conversely, it makes an abrupt change more likely as the climate tips a grid-box from being predominantly one PFT to being predominantly another. Overall, the model could be slightly underestimating the amount of change in vegetation. However, because of the ratio of the biogeophysical to biogeochemical changes, if the vegetation change is underestimated, the sign of the net effect of the terrestrial biosphere is unlikely to change. Similarly, because on the long time periods involved much of the released carbon is
- 25 taken up by the ocean, the changes in carbon densities of the vegetation would need to be wrong by a lot to change the overall signal.

Our approach here assumes that there is no non-linear interaction between the biogeochemical and biogeophysical effects. Since the biogeochemistry acts as a negative feedback and reduces over time, and the biogeophysics acts as a positive feedback and stays the same over time, there's no strong reason to believe that in equilibrium there would be any significant synergy. However, on shorter timescales and on a regional rather than global scale, it is quite possible that there could be some synergies.

5 Conclusions

Using a fully coupled atmosphere-ocean-vegetation model with static and dynamic vegetation, we find that over the last 120 ka the net effect of vegetation feedbacks on global, annual mean 1.5m air temperature is a cooling, which can be as much

as -0.66°C (Figure 2 b). For the vast majority of the last glacial-interglacial cycle, cooling associated with biogeophysical feedbacks dominate over the biogeochemical warming associated with reduced terrestrial carbon storage. The biogeophysical cooling effect is mainly due to the role that vegetation plays in changing surface albedo and particularly related to snow cover and the taiga/tundra transition (Gallimore and Kutzbach, 1996; de Noblet et al., 1996) and we believe is relatively robust.

- 5 The biogeochemical contribution to atmospheric carbon dioxide is small (~20ppmv) and hence the temperature contribution is small (on average 0.26°C with a maximum of 0.33°C). There are significant uncertainties in this calculation which would further diminish the net temperature impact of the terrestrial biosphere by cancelling out the biogeophysical impact. In this analysis, the only time periods where the effects are comparable are at times when additional mechanisms operate, such as changes in ocean circulation, but these mechanisms may be model specific.
- 10 The key uncertainties in this study originate in the biogeochemistry, especially the soil carbon build-up in newly exposed land, the fate of soil carbon in glacial systems, and the amount of carbon in permafrost (not calculated in this study). Further research is needed to fully understand the functioning of these systems and how they can be best incorporated into climate models. In addition, the technique we use for inferring the biogeochemical effects of terrestrial carbon changes has limitations and is potentially model dependent. However, the smaller estimate of terrestrial carbon emissions may make the low LGM 15 atmospheric carbon dioxide somewhat easier to reconcile (Montenegro et al., 2006).

Our work confirms previous results using EMICs Brovkin et al. (2012) that found the net terrestrial biosphere effect to be primarily biogeophysical and that the terrestrial carbon contribution to atmospheric carbon is comparatively small. Our findings also represent a clear illustration of the net climatic effect of vegetation is highly dependent on the timescale, with the biogeophysical response dominating in the longer term in contrast to century-scale future changes.

20 6 Code availability

The model code is currently available to view at http://cms.ncas.ac.uk/code_browsers/UM4.5/UMbrowser/index.html.

7 Data availability

The GCM simulation data is available at http://www.paleo.bris.ac.uk/ummodel/scripts/papers/Davies-Barnard_et_al_2017. html.

25 Appendix A

Author contributions. PJV and JSS ran the climate model simulations. TDB did the analysis and wrote the manuscript. AR ran the cGENIE model simulations. All the authors provided comments and contributed to the manuscript.

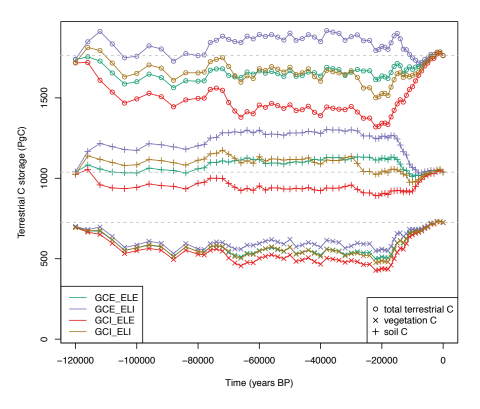


Figure 8. Absolute changes of carbon stores (vegetation and soil) over time. The four scenarios are as detailed in Table 1.

 Table A1. Terrestrial carbon storage at the LGM.

Carbon storage Scenarios	Soil C (PgC)	Vegetation	С	Total C (PgC)
		(PgC)		
GCI_ELE Carbon under ice sheets released to	893	430		1323
atmosphere. No carbon on expanded land area.				
GCE_ELE Carbon under ice sheets remains	1114	502		1617
stored under the ice. No carbon on expanded				
land area.				
GCE_ELI Carbon under ice sheets remains	1249	552		1800
stored under the ice. Modelled carbon storage				
on new land included.				
GCI_ELI Carbon under ice sheets released to	1027	479		1506
atmosphere. Modelled carbon storage on new				
land included.				

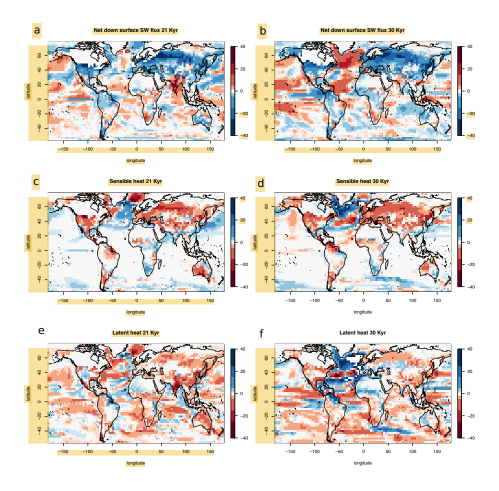


Figure 9. Maps of mean annual anomalies between Dynamic and Static simulations for net down short wave flux, sensible heat, and latent heat, for the 21 kyr and 30 kyr simulations.

Competing interests. The authors have no competing interests to declare.

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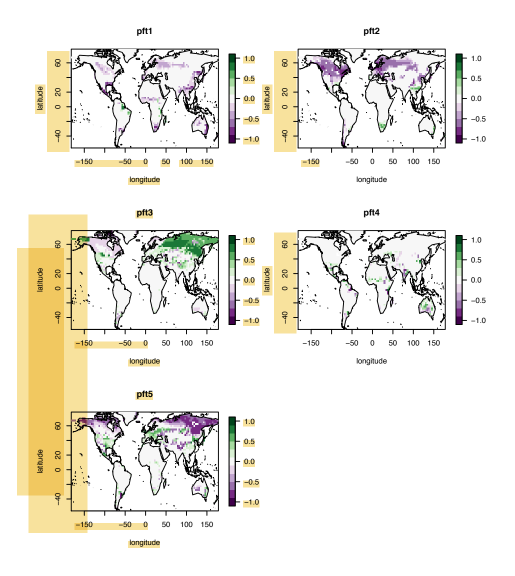


Figure 10. Maps of mean annual anomalies of vegetation cover between LGM and PI in the Dynamic simulations, for the five PFTs. PFT1 is broadleaf trees; PFT2 is needleleaf trees; PFT3 is C3 grasses; PFT4 is C4 grasses; PFT5 is shrubs.

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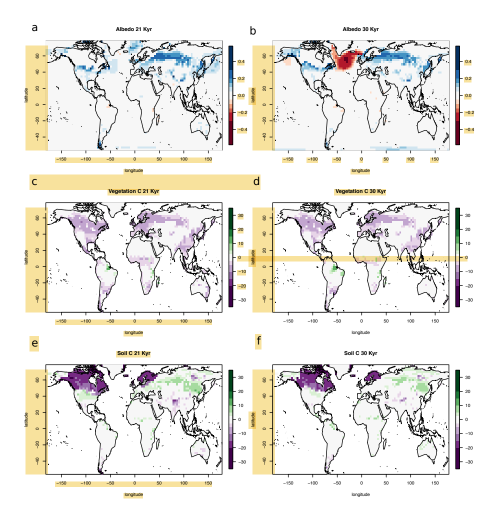


Figure 11. Mean annual anomaly of: a) and b), surface albedo (unitless); c) and d), vegetation carbon (kg C m⁻²); and e) and f), soil carbon (kg C m⁻²), between the Dynamic Vegetation simulation and the equivalent Static Vegetation simulation. For 21 ka (a, c, and e) and 30 ka (b, d, and f). The pattern of reduced surface albedo at 30 ka is similar to the pattern at 56 ka and 100 ka. Since in the Static simulations the carbon remains at PI levels, figures c - f also represent the anomaly to PI. This figure includes the carbon under ice-sheets.

{LGM}, Palaeogeography, Palaeoclimatology, Palaeoecology, 239, 510 - 527, doi:https://doi.org/10.1016/j.palaeo.2006.01.017, http://www.sciencedirect.com/science/article/pii/S0031018206000472, 2006.

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