HOLOCENE POTENTIAL NATURAL VEGETATION IN EUROPE: EVALUATING THE MODEL SPREAD WITH THREE DYNAMICAL VEGETATION MODELS

A PREPRINT

Isabeau A. Bertrix^{*1}, Isashi Sato^{†2}, Isashi Soto^{†2}, Isa

²Research Institute for Global Change (RIGC), Japan Agency for Marine-Earth Science and Technology (JAMSTEC), Yokohama, Japan

³Department of Natural Sciences and Environmental Health, University of South-Eastern Norway, Bø, Norway ⁴Earth and Climate Cluster, Faculty of Science, Vrije Universiteit Amsterdam, Amsterdam, the Netherlands

March 5, 2025

This manuscript is a non-peer-reviewed preprint submitted to EarthArXiv. It has not yet undergone formal peer review. This manuscript has been submitted for peer review to Quaternary Science Reviews (QSR).

Highlights

- Three different DGVM models (SEIB-DGVM, ORCHIDEE-DGVM and CARAIB) were run under climate conditions from five time-slices from 8.5 k.a. BP to pre-industrial
- When comparing simulated PNV to pollen-based reconstructions, all models agree on a similar evolution highlighting the increase human pressure with time
- CARAIB is in worse agreement with the reconstructed vegetation cover than SEIB-DGVM and ORCHIDEE-DGVM, the latter two being skilled in different but complementary aspects
- Overall, the three models differ significantly, suggesting that the use of several models could be a good way to estimate errors in further studies

^{*}isabeau.bertrix@lsce.ipsl.fr

[†]hsatoscb@gmail.com

[‡]nicolas.viovy@lsce.ipsl.fr

 $^{^{\$}}$ hans.renssen@usn.no

 $[\]P$ didier.roche@lsce.ipsl.fr

ABSTRACT

The period of the early Holocene in Europe is marked by climate warming as Earth comes out of the last glacial period and is followed by the emergence of agriculture and animal husbandry in the second half of the period. Increased human influence had profound impacts on the land surface, but the Holocene climate evolution also drove some changes that are intertwined with it. Deciphering the role of each in the vegetation evolution is becoming more difficult as one progresses to the earlier parts of the Holocene here human induced impacts were fainter. Within this general context, we aim at understanding how much Dynamical Vegetation Models (DGVMs) differ in their representation of Potential Natural Vegetation (PNV) in Europe during the mid- to late Holocene (8.5 k.a. BP to preindustrial). We ran three different DGVMs, SEIB-DGVM, ORCHIDEE-DGVM and CARAIB, in Europe, for five time-slices and forced them with identical climatic imputs obtained from the iLOVECLIM Earth system model (downscaled and bias-corrected). Results are then compared with pollen-based reconstructions from the LANDCLIM II database. Overall, the three models have a similar performance in representing the pollen-derived vegetation cover at the european scale. However, their results are largely different at regional scales, particularly in mountainous areas and in boreal regions. They also show a very large spread in simulated plant diversity at the grid cell scale, highlighting the impact of each model's internal dynamics on the results.

Keywords Holocene · DGVM · paleo-vegetation · climate modeling

1 Introduction

Climate always had a strong influence on both vegetation and humanity. For example, in Europe, the drop in temperature during the last glacial maximum (LGM) resulted in a drop in human population, as shown by Tallavaara et al., 2015 by using climate envelope modeling tools and modern ethnographic datasets. Climate variation also influenced landscape, which is highly sensitive to temperature, precipitation, CO2 levels, humidity and solar radiation. Another big influencer of landscape are human activities. Indeed, if we look at the Holocene period, human activities such as deforestation, hunting and agriculture, have led to substantial changes in the european vegetation cover Nikulina et al., 2024, as well as a rise in CO2 levels more recently Petit and Raynaud, 2020. Moreover, land cover is an integral component of the climate system that can lead to two types of feedback loops : biogeophysical (albedo, thermal conductivity) and biogeochemical (carbon dioxide absorption or release during photosynthesis or decomposition). Indeed, forests have a lower albedo than crops or pasture, and this difference is even more marked in the presence of snow. Furthermore, when they grow, trees convert atmospheric carbon into carbon that they store in their trunk, branches, leaves, roots and in the soil. This carbon, as well as dust aerosols, is released into the atmosphere during deforestation due to biomass burning. Human societies impact on vegetation thus modifies the climate directly and through the mediation of the carbon cycle. Additionally, for a given set of climate and human impact conditions the vegetation cover is not ubiquitous but is also a result of the past history. Most of the studies made on the Holocene vegetation have been made on the Green Sahara Lu et al., 2018.

A good candidate to quantify the human influence on vegetation is the comparison between pollen-based reconstructions and Dynamical Global Vegetation Model (DGVMs) simulations. Indeed, pollen-based reconstructions are a good indicator of the type of plants, even sometimes species, that lived at a given period of time on a land surface. They represent the vegetation effectively affected by the combination of human activities and climate conditions. In the absence of specific land-use forcings (for example, Chini et al., 2014), DGVM simulations represent the Potential Natural Vegetation (PNV) that could have existed in the absence of large scale human intervention. DGVMs use climatic data as input to compute the carbon uptake, vegetation cover and competition and related surface conditions. As such, they are sensitive to climatic variations. Under such natural forcing conditions, the differences between pollen-based reconstructions and DGVM simulations should represent the impact of human activities.

In a previous study using this method Zapolska et al., 2023 has shown that humans had a large impact on vegetation cover even before agriculture started. Indeed, simulations with the CARAIB DGVM model forced by bias-corrected climate model outputs showed that vegetation cover in Europe differed significantly from the state of potential natural vegetation even before 6 kyrs BP and challenges the hypothesis that vegetation during the mid-Holocene was in a relatively natural state. This study shows an increase in the difference between PNV and pollen reconstruction with time. Strandberg et al., 2022 studied the difference between PNV and pollen-based reconstruction in Europe at 6 k.a. BP using LPJ-GUESS as DGVM, and showed that PNV was mostly composed of forests, while pollen-based reconstructions has a large component of open lands. In another study at global scale, Dallmeyer et al., 2022, Dallmeyer et al., 2023 also showed disagreement between numerical simulations with the coupled MPI-ESM1.2 and pollen reconstructions with forest expansion post-deglaciation occuring 4,000 years before what the pollen reconstructions

indicate in the Northern Hemisphere.

To progress one step further, we need to identify whether these discrepancies are arising from the capability of our current vegetation models to represent fundamental aspects of vegetation evolution during climate change or our ability to apprehend the time-cumulative impacts of early human societies on vegetation, or both. Indeed, vegetation models differ in sensitivities to climate and CO2 levels, which results in uncertainties in simulation results. Comparing results of different DGVM models seems a good way to minimize errors, as it has been used in several studies. For example, on the Holocene period, Li et al., 2019 runs two DGVM models, LPJ-GUESS and VECODE on a global scale, Hopcroft et al., 2017 runs three models, JULES version 4.1 SDGVM, and LPJ on the green Sahara. Our study seeks to further intercompare three different DGVMs over the european area by studying the simulated potential natural vegetation response to climate forcing during the Late Holocene period (8k5 BP to preindustrial). By comparing the different responses we seek to get a grasp at the common and diverging patterns (in space and time) to assess the common response to climate change; by comparing their responses to an appropriately clustered pollen compilation, we aim at adding a quality assessment for the different regions. We also want to compare the DGVMs in general which has rarely been done.

The first section describes the different methods used during this study, the DGVM models and the adaptations we incorporated, as well as an introduction to the pollen-based reconstruction. The second section provides a detailed description of the numerical experiment setup, the third section presents the results obtained, and the fourth section develops a discussion around the ability of DGVM models to describe PNV.

2 Methodology

In order to study the impact of long-term climate changes on the vegetation of the second-half of the Holocene in Europe, we need a climatic dataset. We choose using climate data simulated with the iLOVECLIM Earth system model for its rapidity and flexibility, required for a long term paleo climate study. iLOVECLIM results have to be bias-corrected, before being used as inputs for a DGVM. We also compared the results of the DGVM simulations with pollen-based reconstructions.

2.1 Climatic data

Three elements are needed to implement our approach: a climatic dataset for each timeslice we are investigating, the bias correction method and a reference climatic dataset to anchor the bias correction.

2.1.1 iLOVECLIM model description

We used the iLOVECLIM Earth system model (here in version 1.1.5), derived from the original LOVECLIM 1.2 model Goosse et al., 2010, as revised by Caley and Roche, 2013. It belongs to the class of Earth System Models of Intermediate Complexity (EMIC), allowing much faster computation than with the more computationally intensive General Circulation models (GCMs). Efficiency is an integral part of our approach since we want to run the complete Holocene time period in the long run: to keep a consistent approach throughout, we thus keep iLOVECLIM as a base model for our timeslices. iLOVECLIM is run in an atmosphere–ocean–vegetation fully coupled configuration, thus including: the atmospheric model, ECBilt, the sea-ice ocean component, CLIO, and the reduced-form dynamic global vegetation model, VECODE.

iLOVECLIM produces results at approximately $5.625^{\circ} \times 5.625^{\circ}$ latitude-longitude spatial resolution grid (T21) over the world. In order to obtain a better spatial resolution over our study area, Europe, we make use of the online interactive downscaling method embedded in iLOVECLIM, first described by Quiquet et al., 2018. It is an online dynamical downscaling of temperature and precipitation included in iLOVECLIM, replicating the processes of surface temperature and precipitation on a refined vertically extended grid. It allows for the computation of surface temperature and precipitation at any altitude within a given subgrid. For our European context, we use a $0.25^{\circ} \times 0.25^{\circ}$ latitude-longitude spatial resolution grid over mainland Europe.

However, as in all climate simulations, iLOVECLIM presents biases and its outputs need to be corrected using observation records before being used in DGVM models.

2.1.2 Bias-correction with CDF-t

The method we used for bias-correction is the "cumulative distribution function transform" (CDF-t), a statistical method based on Quantile Mapping (QM). But contrary to the QM method, which considers that model and observation distributions keep the same shape with time, CDF-t considers that model and observational distributions can evolve and be different. To bias-correct a climatic dataset using the CDF-t method, an observationnal reference datased of the same resolution is needed. Three climatic iLOVECLIM outputs have been corrected using this method : the surface air temperature, the precipitation, and the relative humidity.

2.1.3 Reference observation dataset

The EWEMBI climatic dataset was used as the observational reference dataset during the bias-correction Lange, 2019. It is a re-analysis combination resulting in a daily temporal resolution climatic dataset from 1979 to 2016 at 0.5° horizontal resolution for the entire globe. Before using it, we bilinearly interpolated the data on the $0.25^{\circ} \times 0.25^{\circ}$ European grid. In addition to be used as the observational reference dataset for the bias-correction, we also used it as a preindustrial climatic forcing for our DGVM inputs.

2.2 DGVM models

2.2.1 SEIB-DGVM

Spatially Explicit Individual Based (SEIB-DGVM, here in version 3.03) is a dynamic vegetation model, which aims to simulate the transient impacts of climate change on the terrestrial ecosystem and land-atmosphere interactions Sato et al., 2007. It contains mechanical or empirical algorithms describing terrestrial physical processes (hydrology, radiation, air, etc.), plant physiological processes (photosynthesis, respiration, growth, etc.) and plant dynamic processes (establishment, mortality, disturbance). Unlike other existing DGVMs, SEIB-DGVM only simulates the local interactions of individual trees within a spatially explicit virtual forest; several sample plots are placed in each grid box where the growth, competition for light and decay of each individual tree within a group of trees are then calculated. While accurately representing plant dynamics is crucial for capturing the time lag in vegetation distribution responses to climate change, the specific way plant dynamics are incorporated varies significantly among existing DGVMs Fisher et al., 2018. SEIB-DGVM takes the most direct approach by modeling individual trees competing for space and light, thereby explicitly simulating vegetation dynamics.

SEIB-DGVM simulates 16 PFTs: 6 tropicals, 3 temperates, 5 boreals (with 2 specifically siberian) and 2 herbaceous PFTs. As no tropical or siberian pollen have been recorded in Europe during the Holocene, we chose to disable the 6 tropical PFTs and the 2 siberian PFTs.

As previous studies using SEIB-DGVM were mainly applied either on global scales Tong et al., 2022 or on african or siberian Sato et al., 2023 zones, there was no mediterranean PFT (contrary to the pollen records). We decided to create one by modifying the existing "Temperate broad-leaved evergreen" PFT. The parameters associated with this new PFT can be found in Table S7. We also add the drought parameter, which is the state of water satisfactory for photosynthesis (0.0-1.0), from SEIB-DGVM 3.10 version Sato et al., 2023 to the SEIB-DGVM 3.03 version. Following Sato et al., 2023, we choose to associate coniferous PFTs (temperate or boreal) with an optimal state of water satisfactory of 0.75, a minimum state of water satisfactory of 0.20, and a maximum state of water satisfactory of 0.90; deciduous PFTs (temperate or boreal) with an optimal state of water satisfactory of 0.30, and a maximum state of water satisfactory of 0.30. An other change in SEIB-DGVM is the modification of the net CO_2 assimilation rate per needle area for the coniferous. Indeed, it has a strong dependence on leaf age, as measured in Robakowski and Bielinis, 2017, which is not taken into consideration in the SEIB-DGVM. Consequently, we decreased the net CO_2 assimilation rate per needle area from 9.0 $\mu mol.m^{-2}.s^{-1}$ to 7.0 $\mu mol.m^{-2}.s^{-1}$ for both the temperate needle-leaved evergreen PFTs.

Finaly, as SEIB-DGVM was never used to study paleoclimate, it had to be adapted to consider astronomical parameters. Indeed, SEIB-DGVM calculates the solar declination and the distance between the Earth and the Sun for a given day of a year, in a way that is only applicable at present day. To make this calculation valid for paleo times, we need to obtain it only using the astronomical parameters (from Berger and Loutre, 1999) : the obliquity ϵ , the eccentricity e, and the climatic precession ϖ . The details of the calculation are described in supplementary.

2.2.2 ORCHIDEE-DGVM

ORCHIDEE is the land surface model of the IPSL (Institut Pierre Simon Laplace) Earth System Model. In this study, we used ORCHIDEE version 2.2 with DGVM activated. ORCHIDEE is based on three different existing models : SVAT SECHIBA, that describes exchanges of energy and water between the atmosphere and the biosphere, and the soil water

budget; LPJ-DGVM for the parameterizations of vegetation dynamics (fire, sapling establishment, light competition, tree mortality, and climatic criteria for the introduction or elimination of PFTs); and STOMATE, for processes such as photosynthesis, carbon allocation, litter decomposition, soil carbon dynamics, maintenance and growth respiration, and phenology. The DGVM part of ORCHIDEE simulates the growth of 10 PFTs, and have also a bare soil PFT: 2 tropical, 3 temperate, 3 boreal, and 2 herbaceous PFTs. As no tropical pollens as been recorded on Europe during the Holocene, we choosed to disabled the two tropical PFTs.

2.2.3 CARAIB

CARbon Assimilation In the Biosphere (CARAIB) model is a dynamic vegetation model originally designed to study the role of vegetation in the global carbon cycle, at present and in the past. It is composed of five modules François et al., 2011 describing respectively (1) the hydrological budget, (2) canopy photosynthesis and stomatal regulation, (3) carbon allocation and plant growth, (4) heterotrophic respiration and litter/soil carbon dynamics, and (5) plant competition and biogeography. In CARAIB, photosynthesis and plant respiration are computed every two hours, and water and carbon reservoir are updated every day. The model simulates 26 plant functional types (PFTs), which can coexist on the same grid cell: 3 tropical, 3 subtropical, 1 subdesertic, 4 mediterranean, 6 temperate, 6 boreal and 3 herbaceous PFTs. In CARAIB, herbs and shrubs are assumed to grow under the trees, leading to a two layers structure. Each layer has a maximum coverage of 1, meaning that the maximum vegetation fraction of a grid cell is 2. As no tropical pollen has been recorded in Europe during the Holocene, we chose to disable the 3 tropical PFTs and the subdesertic one. Another difference with SEIB-DGVM and ORCHIDEE-DGVM is the presence of shrub PFTs. In order to compare properly the three models, we chose to disable the shrubs PFTs, leaving 16 PFTs (3 herbaceous and 13 tree types). In the post-treatment process, we decided to reclassify CARAIB's remaining PFTs into the 8 SEIB's PFTs. The reclassification is presented below in Table (1).

SEIB-DGVM PFT	CARAIB PFT
Temperate needle-leaved evergreen	NEg Te cool trees
Mediterranean	NEg sMed trees, NEg mMed trees, NEg subTr trees, BEg mMed trees, BEg
	tMed trees, BEg subTr trees, NSg subTr swamp trees
Temperate broad-leaved summergreen	Sg Te cool trees, BSg Te warm trees
Boreal needle-leaved evergreen	NEg B/Te cold trees
Boreal needle-leaved summergreen	NSg B/Te cold trees
Boreal broad-leaved summergreen	BSg B/Te cold trees
Herbaceous	C3h, C3d, C4

Table 1: CARAIB PFTs classified into SEIB-DGVM PFTs

2.3 Pollen-based reconstruction: the LANDCLIM II database

2.3.1 The REVEALS method

In order to compare the results of our European DGVM Holocene simulations, to land-cover reconstructions, we used the LANDCLIM II pollen-based reconstruction dataset. It comes from the aggregation of several raw pollen databases (1607 records, mainly from lakes and peatlands) which are interpreted quantitatively using the REVEALS (Regional Estimation of VEgetation Abundance from Large Sites) model Serge, 2023. The taxa used are only anemophilous taxa, their dispersion is assumed to be isotropic Serge et al., 2023. To limit representation biases in the model, data is weighted by the Relative Pollen Productivity (corresponding to the number of pollen grains produced per species), by the average pollen fall speed, by the diameter of the sampling zone and by climatic conditions, following the protocol of Githumbi et al., 2022.

The REVEALS model output, LANDCLIM II, is a dataset comprising a total of 378 different coordinates for 31 taxa, on a 1 °x 1° grid across 30° – 71 °N, 20° W–47°E (north-western, central Europe, Mediterranean area, and part of the East until 47°E) for 25 contiguous time slices of 100-500 years covering all of the Holocene.

In the LANDCLIM II dataset, total cover of plant taxa is always 100 %, as it is challenging to estimate the proportion of bare soil in pollen reconstruction.

2.3.2 Classification of pollen taxons into PFTs

As DGVMs simulate the growth of PFTs and not species, we classified LANDCLIM II taxa into PFTs in order to compare models and data efficiently. The LANDCLIM II taxa are given as Taxons in Tables 2 and 3.

We classified the PFTs of CARAIB into those of SEIB-DGVM in Table 1. But SEIB-DGVM and ORCHIDEE-DGVM do not exactly have the same PFTs. Instead of a mediterranean PFT, which can contain both broad-leaved and needle-leaved species, ORCHIDEE-DGVM has a temperate broad-leaved evergreen PFT. This is why Pinus is classified as mediterranean in SEIB-DGVM, and as Temperate needle-leaved evergreen in ORCHIDEE-DGVM. All the other PFTs classified as mediterranean in SEIB-DGVM are classified as Temperate broad-leaved evergreen in ORCHIDEE-DGVM. All the other PFTs classified each taxa into one or two of the SEIB-DGVM PFTs, as shown in Table 2, and into one or two of the ORCHIDEE-DGVM PFTs in Table 3. For Juniperus, Ericaceae, Pinus, and Salix, we chose to classify them into two different PFTs, as their ecology can be associated with both. For each grid cell, those taxa are classified in one of the two PFTs, according to the fraction of these two PFTs already present on the grid cell considered. At the end, we obtained the percentage of each PFT for a given box. That way, every box can be composed of the sum of the 6 PFTs, for a total of 100 %. No taxa corresponded to the boreal needle-leaved summergreen PFT. Also, the herbaceous PFT contain not only C3 herbaceous taxa, but also heather (Calluna Vulgaris) and C4 herbaceous.

SEIB-DGVM and CARAIB PFT	Taxons
Temperate needle-leaved evergreen	Abies alba, Juniperus
Mediterranean	Buxus sempervirens, Carpinus orientalis, Ericaceae, Phillyrea, Pinus, Pistacia,
	Quercus evergreen
Temperate broad-leaved summergreen	Alnus glutinosa, Carpinus betulus, Castanea, Corylus avellana, Fagus sylvatica,
	Fraximus, Quercus deciduous, Salix, Tilia, Ulmus
Boreal needle-leaved evergreen	Ericacea, Juniperus, Picea, Pinus
Boreal needle-leaved summergreen	
Boreal broad-leaved summergreen	Betula, Salix
Herbaceous	Amanranthaceae chenopodiaceae, Artemisia, Calluna vulgaris, Cerealia t,
	Cyperaceae, Filipendula, Plantago lanceolata type, Poaceae, Rumex acetosa t,
	Secale

Table 2:	LANDCLIM	II taxons	classified into	SEIB-DGVM PFTs
----------	----------	-----------	-----------------	----------------

ORCHIDEE-DGVM PFT	Taxons
Temperate needle-leaved evergreen	Abies alba, Juniperus, Pinus
Temperate broad-leaved evergreen	Buxus sempervirens, Carpinus orientalis, Ericaceae, Phillyrea, Pistacia, Quer-
	cus evergreen
Temperate broad-leaved summergreen	Alnus glutinosa, Carpinus betulus, Castanea, Corylus avellana, Fagus sylvatica,
	Fraximus, Quercus deciduous, Salix, Tilia, Ulmus
Boreal needle-leaved evergreen	Ericacea, Juniperus, Picea, Pinus
Boreal needle-leaved summergreen	
Boreal broad-leaved summergreen	Betula, Salix
Herbaceous	Amanranthaceae chenopodiaceae, Artemisia, Calluna vulgaris, Cerealia t,
	Cyperaceae, Filipendula, Plantago lanceolata type, Poaceae, Rumex acetosa t,
	Secale

Table 3: LANDCLIM II taxons classified into ORCHIDEE-DGVM PFTs

2.3.3 Distance between model and data

In order to compare model and data, we also create a set of distance values between PFTs in Tables 4 for SEIB-DGVM and CARAIB PFTs compared to LANDCLIM II PFTs, and 5 for ORCHIDEE-DGVM PFTs compared to LANDCLIM II PFTs. This distance will be used later to estimate how much each model differs with the pollen data. Basically, a distance 0 is when PFTs are exactly the same, 1 is for same climate (temperate or boreal) but different PFTs (for example, a Boreal Broad-Leaved Summergreen instead of a Boreal Needle-Leaved Evergreen), 2 is for different climate but still tree PFT (for example a Temperate Broad-Leaved Summergreen instead of a Boreal Broad-Leaved Summergreen), and 3 is for herbaceous instead of tree or vice-verça. Note that the combination TeNEg/BNEg has a

distance of 1 even if one is boreal and the other	one temperate,	because in the	pollen, the	e taxa classifi	ed as [FeNEg or
BNEg can all belong to both of those PFTs.						

	TeNEg	Med	TeBSg	BNEg	BNSg	BBSg	Herbaceous
TeNEg	0	1	1	1	2	2	3
Med	1	0	2	3	3	3	3
TeBSg	1	2	0	2	2	2	3
BNEg	1	3	2	0	1	1	3
BNSg	2	3	2	1	0	1	3
BBSg	2	3	2	1	1	0	3
C3	3	3	3	3	3	3	0
C4	3	3	3	3	3	3	0

Table 4: distance PFTs SEIB-CARAIB/LANDCLIM II

	TeNEg	TeBEg	TeBSg	BNEg	BNSg	BBSg	Herbaceous
TeNEg	0	1	1	1	2	2	3
TeBEg	1	0	2	3	3	3	3
TeBSg	1	2	0	2	2	2	3
BNEg	1	3	2	0	1	1	3
BNSg	2	3	2	1	0	1	3
BBSg	2	3	2	1	1	0	3
TeC3	3	3	3	3	3	3	0
C4	3	3	3	3	3	3	0
TrC3	3	3	3	3	3	3	0
BC3	3	3	3	3	3	3	0

Table 5: distance PFTs ORCHIDEE/LANDCLIM II

3 Numerical experiment setup

During this study, we ran five different time-slice DGVM simulations within the Holocene: preindustrial, 3 k.a., 4 k.a., 6 k.a. and 8.5 k.a. BP (Before Present). To force the different DGVMs, we create climatic inputs consistently generated by iLOVECLIM (downscaled and bias-corrected), and the EWEMBI climatic inputs (for present day as we did not have any preindustrial climatic dataset). Each model uses a set of closely related climate fields, as shown in Table 6. We also provided appropriate CO_2 concentration relative to the Holocene to each model (note that pre-industrial CO2 levels were used for the pre-industrial DGVM simulation), as well as astronomical parameters (obliquity, eccentricity and climatic precession).

It should be noted that iLOVECLIM runs already with a simplified vegetation model included (VECODE), to provide the first order response of climate in a coupled mode. Hence, our climate forcing is dependent upon the vegetation results of VECODE. This type of setup has been already used Li et al., 2019 and has shown that as long as we concentrate on the first order response (which is the case when looking at the climate impact in iLOVECLIM), it was comparable in vegetation models of different complexities. In addition, since the process here involves a further step with bias correction, the obtained climate forcing is not simply a direct response to that vegetation coupling.

Here, the daily surface air temperature amplitude, the minimum and maximum surface air temperature and the wind speed come from the EWEMBI dataset and were kept constant for each period. Each simulation was run until reaching equilibrium for the simulated vegetation (300 years for SEIB-DGVM, 150 years for ORCHIDEE-DGVM, 380 years for CARAIB).

Variable	Unit	SEIB	CARAIB	ORCHIDEE
Surface Air Temperature	°C	X	Х	Х
Surface Air Temperature daily amplitude	°C	X	Х	
Minimum surface Air Temperature	°C			Х
Maximum surface Air Temperature	°C			Х
Precipitation	mm.day $^{-1}$	X	Х	Х
1 - cloud cover	%		X	
Downward shortwave radiation at midday	$W.m^{-2}$	X		Х
Downward longwave radiation	$W.m^{-2}$	X		Х
Wind speed	$m.s^{-1}$	X	Х	Х
Relative humidity	%	X	Х	X

Table 6: Climate fields for each DGVMs

4 Results

4.1 Model intercomparison

4.1.1 Biome comparison

One classical way to intercompare results between different vegetation results is to express them in the form of biomes Haxeltine and Prentice, 1996. We chose to generate biomes according to the methodology from SEIB, which uses as input the dominant PFT and the mean annual maximum Leaf Area Index (LAImax) over the last ten years of simulation at each point. If the LAI max is higher than $2.5 m^2 . m^{-2}$, the associated biome will be a forest one ; if not but still higher than $1.5 m^2 . m^{-2}$, it will be a woodland one. If the LAI max is lower it will be a grassland/steppe/savanna biome, and if lower than $0.2 m^2 . m^{-2}$, a desert biome. Polar desert and Arctic/Alpine-tundra biomes are only defined by a threshold in Growing Degree Days with a limit at zero or five Celsius (GDD0 and GDD5) respectively. Biome will be a polar desert if GDD0 < 150, or a Arctic/Alpine-tundra if GDD5 < 370. In order to compare the results of the three DGVM models, we applied the same biome code on ORCHIDEE-DGVM and CARAIB results. As ORCHIDEE-DGVM has a different functioning, the LAImax needs to be multiplied by the maximum vegetation fraction. The dominant PFT is the PFT having the greatest mean annual Net Primary Production (NPP) over the last ten years of simulation. Again for ORCHIDEE-DGVM, the NPP needs to be multiply by the maximum vegetation fraction.



Figure 1: Biomes maps for SEIB-DGVM (left) and ORCHIDEE-DGVM (middle) and CARAIB (right) for 8.5 k.a. (top), 6 k.a. (middle) and preindustrial (bottom) climatic dataset



Figure 2: Average temperature map for a typical year in the 6 k.a. climatic dataset

As seen in Fig 1, the biome results have some common patterns in all DGVM models: boreal biomes are located at the high latitudes and temperate biomes at the mid-latitudes as expected. The spatial coverage of the different biomes is however different, highlighting potential different processes behind the distribution: SEIB-DGVM generates a biome coverage that is heterogeneous, with more variability in the biome determination within a single region. For example, where Scandinavia is covered by forests, it is a mixture of boreal forests, both deciduous and evergreen. Conversely, in ORCHIDEE-DGVM and CARAIB, zones are much more homogeneous, with a single biome over wide areas. Another obvious difference is that SEIB-DGVM has zones with woodland at the mid latitudes whereas ORCHIDEE-DGVM has none and CARAIB have few in the high latitudes.

In the Alps, all models differ : CARAIB grows a boreal deciduous forest, ORCHIDEE-DGVM a steppe, SEIB-DGVM a boreal conifer forest.

In boreal regions, CARAIB grows very few conifer forests but clearly separated steppes and deciduous forest, SEIB-DGVM grows a mix of conifer and deciduous forest (dominated by conifer forest), and ORCHIDEE-DGVM grows separate conifer and deciduous forest, as well as steppes on the Atlantic coast of Norway.

Also, the limit between boreal and temperate biomes is shaped differently : SEIB-DGVM seems to have a limit determined by latitude, whereas ORCHIDEE-DGVM and CARAIB seems to have a northwest-southeast limit, resulting in regions like southern Sweden and Belarus, Estonia, Lituania and Latvia with temperate biomes in ORCHIDEE-DGVM and CARAIB, but boreal in SEIB-DGVM. One probable reason for this is that boreal and temperate PFTs geographical repartition in SEIB-DGVM is less sensitive to temperature than in ORCHIDEE-DGVM and CARAIB and more to incoming solar radiation. Indeed, the shape of this limit in ORCHIDEE-DGVM and CARAIB is clearly similar to the average temperature gradient for a typical year (taken here as an example from the 6 k.a. dataset in Fig 2) while the incoming radiation is the one forcing that has a clear latitudinal component.

Now having a look at the differences across the different time windows in Fig. 1, we can clearly see that all models have very similar simulated biomes maps between 8.5 k.a. and 6 k.a. (with a clearer difference between those and preindustrial) and that all models see Europe as being temperate deciduous forest dominated. Similarly, all models grow less boreal forests at preindustrial than at 6 k.a. or 8.5 k.a. In particular, SEIB-DGVM and ORCHIDEE-DGVM grow more herbaceous in northern Scandinavia at preindustrial than at 6 k.a. or 8.5 k.a., and CARAIB grow steppes at all time scale. ORCHIDEE-DGVM and CARAIB grow less temperate broad-leaved evergreen forest at preindustrial than at 6 k.a. or 8.5 k.a., and SEIB-DGVM just a little bit less.

One big difference is the boreal deciduous forest, which is present in all models at all time scales, but with very different evolution with time : SEIB-DGVM grows more boreal deciduous forest with time, ORCHIDEE-DGVM less, and CARAIB quite the same.

4.1.2 Analysis of the differences

As biomes are computed from LAImax and dominant PFT, we hereafter investigate how these variables are simulated in the different models. In Fig S14, the LAImax per number of point for each model, at each timescale is represented on histograms. Those LAImax values are very different: LAImax is much higher in ORCHIDEE-DGVM and in CARAIB than in SEIB-DGVM, especially at preindustrial, which can explain why there is no woodland biome in ORCHIDEE-DGVM, and a few in CARAIB (located in areas surrounded by steppes : nordic regions and south-east of Turkey). Indeed, trees never have a low enough LAImax for that. In the case of CARAIB, approximately half of the points have a very high LAImax, higher than $5m^2/m^2$. The high variability in the dominant PFT in certain geographical regions in SEIB-DGVM questions the concept of dominance. In the following, we investigate how dominant a PFT is with respect to the others.

In Fig 3, the number of PFTs which represent more than 10% of the parameter used to determine the dominant PFT (the mean annual NPP for SEIB-DGVM and CARAIB or the mean annual maximum vegetation fraction for ORCHIDEE-DGVM) is represented on maps for each model and each time window, representing the number of PFTs sharing the same gridcell.

The three models give very different results. ORCHIDEE-DGVM is the model which has the largest number of PFTs per grid cell, hence the highest PFT diversity, followed by CARAIB and then SEIB-DGVM. ORCHIDEE-DGVM has globally more diversity than the other two everywhere, but especially in boreal regions, where between 6 to 8 PFTs are represented. In other words, for some regions, all possible PFTs are able to grow in the same gridcell in ORCHIDEE-DGVM. Those regions are mainly transition regions from temperate to boreal, and contain both temperate and boreal PFTs. The only parts of lower PFT diversity are the Atlantic coast of Norway, the Alps and Scotland, where only 1 to 2 PFTs are present.

SEIB-DGVM has again a substantial variability of PFT distribution between adjacent cells in boreal regions, which is not the case in ORCHIDEE-DGVM and CARAIB. This high spatial heterogeity can be attributable to its being an individual-based model. In opposite, SEIB-DGVM depict a very homogeneous distribution in temperate regions, with only 1 to 3 PFTs present in the same gridcell.

Whereas SEIB-DGVM and ORCHIDEE-DGVM have low PFT diversity in temperate regions, CARAIB has, on the contrary a large diversity with between 3 to 5 PFTs growing in the same gridcell in temperate regions, but seems to have a very clear latitudinal border at around 60°N: beyond that limit to the north, there are only 1 or 2 PFTs growing. We can notice that, except for the preindustrial, SEIB-DGVM simulates more PFT diversity in boreal regions than temperate regions, which is a common feature shared with ORCHIDEE-DGVM, but not with CARAIB, which has a very poor diversity in Scandinavia and Russia.



Figure 3: number of PFTs with a NPP greater than 10% of the mean annual sum for all PFTs for SEIB-DGVM (left), ORCHIDEE-DGVM (middle) and CARAIB (right), at 8.5 k.a. (top), 6 k.a. (middle) and preindustrial (bottom)

Those results suggest that the concept of PFT dominance is largely model dependent. In SEIB-DGVM, in general one PFT dominate all the others whereas in ORCHIDEE-DGVM several PFTs can be present with close PFT fractions.



Another way to look at this relative dominance could be via the magnitude of NPP fraction that is represented by the first dominant PFT as shown in Fig. 4.

Figure 4: percentage of total NPP of the dominant PFT for SEIB-DGVM (left), ORCHIDEE-DGVM (middle) and CARAIB (right) at 8.5 k.a. (top), 6 k.a. (middle) and preindustrial (bottom)

Here again, the three models yield very different results. CARAIB has low overall values, consistent with a high number of dominant PFTs. In ORCHIDEE-DGVM, in contrast, there is still a high value of % of NPP for the dominant PFT, meaning that even in the region where there are 7 PFTs that represent at least 10 % of the total NPP, the dominant PFT can represent half of the total NPP. Whereas in CARAIB, most of the map has a NPP proportion of the first PFT of only 20-40%. SEIB-DGVM has very few regions with less than 50% of NPP dominance. If we consider that a true dominance is more than half the NPP or maximum vegetation fraction, this means that the dominant PFT is actually a dominant PFT for SEIB, some parts of ORCHIDEE and only a few zones in CARAIB.

To have a better view on how mixed the type of plants are in a same grid cell, we further simplified the PFTs in three classes: broad-leaved forest, needle-leaved forest, and herbaceous. Fig 5 shows the proportion of each of those three groups of PFTs for each pixel for SEIB-DGVM, ORCHIDEE-DGVM and CARAIB and for the three time periods.

Again, we can see that there is not much difference between 8.5 k.a. and 6 k.a. for all models. From 8.5 k.a. to preindustrial period, ORCHIDEE-DGVM progressively gains needle-leaved trees, while SEIB-DGVM has a quite constant distribution, again showing a less mixed forest per point than the other models. There are more herbaceous in Scandinavia at preindustrial than at 6 k.a. and 8.5 k.a. in both SEIB-DGVM and ORCHIDEE-DGVM but not in CARAIB, which gets more needle-leaved. CARAIB has herbaceous everywhere, unlike ORCHIDEE-DGVM and SEIB-DGVM which have them mainly in Scandinavia and the Alps, especially at preindustrial. Needle-leaved are more abundant in SEIB-DGVM.



Figure 5: color map (red = herbaceous [%], green = broad-leaved trees [%], blue = needle-leaved trees for SEIB-DGVM (left), ORCHIDEE-DGVM (middle) and CARAIB (right), for 8.5 k.a. (top), 6 k.a. (middle) and preindustrial (bottom) climatic dataset

4.2 Model/data comparison

To evaluate how model simulations compare with data, we used the LANDCLIM II pollen database. As a first step, we expressed the LANDCLIM II dataset in PFTs fraction for each gridcells.

Figs 6, 7 and 8 show respectively the results and analysis for 8.5 k.a., 6k.a., and preindustrial. On each figures, SEIB-DGVM is on the left, ORCHIDEE-DGVM on the middle, CARAIB on the right, pollen map reconstruction on the top, dominant PFTs map for each model on the middle, and distance map between pollen data and models on the bottom (see Tables 4 and 5).



Figure 6: Dominant PFT map for pollen-based reconstruction (top), dominant PFT map for simulation (middle), and distance between model and data (bottom), for SEIB-DGVM (left), ORCHIDEE-DGVM (middle) and CARAIB (right), for 8.5 k.a.

The dominant PFT distribution for each of our three models is hereafter computed following the same methodology as before, using averages over the last ten years. To quantify the model data disagreement, we use the distance matrix introduced in 2.3.3, Tables 4 and 5. The first main globalresult from such a comparison is that the distance between model and data became higher with time for all models. This is coherent with the increase of land use activities Zapolska et al., 2023. Indeed, the pollen reconstruction maps contain more herbaceous PFTs at preindustrial than at 6 k.a. and 8.5 k.a.

The 8.5 k.a. simulations are overall in good agreement with the data, except for Iberian Peninsula and south-east of Europe, dominated by herbaceous in the pollen data. ORCHIDEE-DGVM has a better score than SEIB-DGVM and CARAIB, and SEIB-DGVM doesn't represent the temperate PFT in southern Sweden, again probably because of its distinction between boreal and temperate PFTs which may not be temperature dependent enough (unlike ORCHIDEE-DGVM and CARAIB).

At 6 k.a., the same result is found regarding the limit between boreal and temperate trees, again with ORCHIDEE-DGVM and CARAIB in better agreement than SEIB-DGVM. Also, there are a lot of scores with a value of one in the distance map (orange color), which correspond to needle-leaved versus broad-leaved disagreement. As seen before, this is due to the mixed forest in ORCHIDEE-DGVM. We consider that difference to be a rather small one. Globally, 6 k.a. results and distance maps are very similar to 8.5 k.a..

Concerning the Alps, all models are in disagreement with the pollen reconstructions: LANDCLIM-II indicates a dominance of temperate needle-leaved trees, whereas SEIB-DGVM gives boreal needle-leaved trees and the output of ORCHIDEE-DGVM and CARAIB yield a dominant herbaceous cover. The classification between temperate and



Figure 7: Dominant PFT map for pollen-based reconstruction (top), dominant PFT map for simulation (middle), and distance between model and data (bottom), for SEIB-DGVM (left), ORCHIDEE-DGVM (middle) and CARAIB (right), for 6 k.a.

boreal needle-leaved is somewhat arbitrary; taking that aspect into consideration, SEIB-DGVM is probably the model closest to the reconstructions in the alpine realm.

As expected, the preindustrial map is not data compliant at all for all three models, because of the large proportion of herbaceous dominance in numerous parts of Europe (caused by crop-land and urbanisation), except for the boreal regions where pollen-based reconstruction show a broad forest dominance.

In order to have a better perspective at the time evolution of the data to model agreement, we introduce a generic matching score between the two as follow:

$$M_s = 100 - \sum \frac{distance}{maxdistance} \times 100$$

where distance is represented by the values arising from the bottom panels in figures 6, 7 and 8 and maxdistance is three.

In addition to 8.5 & 6 ka B.P. and preindustrial, we also performed similar computations for the 3 k.a. and the 4 k.a. time windows, and reported all distance values as a matching score in Fig 9.

As already mentioned, the first order signal is a decrease in the matching score between model and data with time. At 8.5 k.a. and 6 k.a., the results are not significantly different in each of the three models. From 6 k.a. BP onwards, all models agree on an accelerating decrease of the matching score towards the preindustrial. The difference between models is about 6-8%. For the pre-industrial period, the results show a matching score of $\approx 20 - 27\%$.

Since the three models are yielding quite similar and comparatively high matching results at 8.5 k.a. BP and 6 k.a. BP, we will as following step analyse where the models give a coherent spatial response for those two time windows. Fig 10 shows the number of models having a distance to pollen data below or equal to one, which corresponds to a perfect agreement with the dominant PFT, or a small error, at 8.5 k.a. and 6 k.a. .

This map defines robust areas for which all models agree and are consistent with pollen data. To a first order, the results are spatially coherent in temperate areas (western and central Europe) and most coastal areas but highlight strong divergences in the Iberian Peninsula, the Alps, Scandinavia and Eastern Europe.



Figure 8: Dominant PFT map for pollen-based reconstruction (top), dominant PFT map for simulation (middle), and distance between model and data (bottom), for SEIB-DGVM (left), ORCHIDEE-DGVM (middle) and CARAIB (right) for preindustrial

Since the largest human impact within this time period is the expansion of agriculture, we expect the difference between trees and herbaceous PFTs to be more telling than the difference between different tree PFTs. It is therefore useful to further simplify our comparison using a simplified weighting matrix, only counting a distance of 1 between herbaceous and trees and 0 in other cases. It is interesting to see (Fig 11) that for 8.5 k.a. and 6 k.a., the difference between the highest and the lowest score is still around 10% as in Fig 9 and that the three models have a reduced spread at the preindustrial.

5 Discussion

In this study, we assumed that the DGVM results are relevant to represent potential natural vegetation. However, as those models are based on empirical relationships between vegetation and climate Levavasseur et al., 2013, they do not take into account the role of non climatic factors such as megafauna on vegetation. It is important to note that megafauna may have had an impact on the opening up of environments, as suggested by Feurdean et al., 2018 who studied actual big mammals and their role in land cover in central Europe, and Zhu et al., 2018, who shows how taking herbivores into account modifies productivity in ORCHIDEE. That said, we assumed here that climate and human activities are the main driving factors of vegetation cover variability during the Holocene in Europe.

Another caveat is arising from the climate forcing used. The Holocene period has an overall weak climate signal and is notoriously difficult to consistently capture in climate models as has been shown previously for example in Hargreaves et al., 2013. The iLOVECLIM model results used here provide one climate forcing that is specific to that model. While our approach correct the large scale model biases through the bias correction, the climate anomalies between present-day and the difference past periods are conserved and are characteristic of the iLOVECLIM model simulations. These have been analyzed elsewhere Arthur et al., 2023. To further analyse the impact of model specific simulations of climate change, the use of a range of climate models results could be used in a future study. A period such as the mid-Holocene Brierley et al., 2020 for which these simulations are available could be a nice target to do so.



data/model matching score as a function of time

Figure 9: Data/model matching score as a function of time for SEIB (blue), ORCHIDEE (green), and CARAIB (red)



Figure 10: Number of models with a distance lower or equal to one with the pollen data for 8.5 k.a. (left), and 6 k.a. (right)



Figure 11: Data/model matching score as a function of time for SEIB (blue), ORCHIDEE (green), and CARAIB (red) for simplified pollen vs model distance

In terms of comparison with data, the DGVM simulations are statistically similar for all models since they have overall similar percentage of correlation to the pollen data. Matching score decrease with time, which is expected as human pressure increase, consistently with what as been shown in Zapolska et al., 2023 on similar time period. The matching score indicates that 74 to 80 % of the studied land cover differs from the PNV at preindustrial time according to Fig 9. Again, those results are consistent with the results found in Zapolska et al., 2023. Looking at Fig 9, even at 8.5 k.a, the maximum matching score is about 54% (SEIB-DGVM), which either means that the human pressure was already important, or the models are not reliable at reconstructing the vegetation cover. Looking back at the spatial results, the distance between pollen reconstructions and SEIB-DGVM in Fig 6 shows that the strongest divergences (score of 3) are located in Turkey and Spain – locations where agriculture was likely already present Gronenborn et al., 2023–, part of Ireland, Scandinavia and a few sporadic zones in temperate Europe. Those results could suggest that the presence of herbaceous taxons in Turkey and Spain have an anthropic origin, but ORCHIDEE-DGVM grows mainly herbaceous in Turkey as PNV. Looking at the LANDCLIM II database, the anthropic origin of the herbaceous PFT is confirmed : herbaceous points in France, Ireland or Scandinavia are not dominated by human related herbaceous species (such as Castanea, Cerealia, Plantago lanceolata, Secale), and could be considered as natural, whereas Spain and Turkey are yet dominated by human related species. Please note that here, we did not count the taxon Poaceae, in either of the herbaceous category (natural or human-related), as Poaceae are a vey big group contening both human-related species such as cerealia and natural species. Concerning Scandinavia, the climate-vegetation models do not account for the long-lasting impact of the ice cover, including soil development and succession of vegetation, since the models calculate the potential vegetation that is in equilibrium with the climate at that moment. In reality the vegetation could have lagged the climate response by up to a few thousand years and this lag effect can be expected to be present in the LANDCLIM II pollen database but not in the models. So part of the difference between the models and pollen data could maybe be explained by this lag-effect Moen and Lillethun, 1999. We can highlight that none of the DGVM models is able to reproduce the herbaceous dominance in the center of France (taken as a local example) and in Spain (Fig 10). As discussed above, the presence of herbaceous dominance in Spain at 8.5 k.a. BP could be explained by human land use. For the few points in the center of France, it could be due to the nature of the soil not being taken into account in the models - or to specific eco-environnements.

The three models outputs are very different. SEIB-DGVM have a very low diversity of PFTs, whereas CARAIB and ORCHIDEE-DGVM have a higher diversity. The high diversity of PFTs of CARAIB and the low importance of its dominant PFT could be explained by its two layers structure, leading to an absence of competition between the herbs



Figure 12: number of PFTs with a fraction of the cells greater than 10% in the REVEALS dataset (top), and percentage occupied by the dominant PFT in the REVEALS dataset (bottom) at 8.5 k.a. (left), 6 k.a. (middle) and preindustrial (right)



Figure 13: openness of pollen data at 8.5 k.a. (left), 6 k.a. (middle), and preindustrial (right).

and the trees. There will be only herbs or trees if the other cannot grow.

In Fig 3 and Fig 4, the three models differ a lot concerning both the number of PFTs per grid cells and the relative importance of the dominant PFT. Like for Fig 3 and Fig 4, in Fig 12 we show the number of PFTs and percentage of the dominant one but estimated from the pollen dataset. Indeed, all the taxons are given as a percentage, so as the reclassified PFTs we created. Those results suggest that the pollen-based reconstruction favors a dominance of between 1 and 4 PFTs per grid cell, mainly 2 or 3. For this particular aspect, SEIB-DGVM is in better agreement with the data than CARAIB and above all ORCHIDEE-DGVM, which grows many more. However, ORCHIDEE-DGVM is better at reproducing the relative importance (in terms of NPP) of a dominant PFT. Given those results, we can notice that the best model on an aspect is not necessarily the best for all the studied parameters ; using two or more models can thus add confidence into the simulated fields.

On a regional basis, pollen data suggest a higher number of PFTs in boreal zones, a tendency well reproduced by SEIB-DGVM and ORCHIDEE-DGVM; CARAIB shows exactly the opposite. CARAIB is also the only model which does not correspond to either the number of PFTs, the geographical distribution, and the importance of the dominant PFT. From this perspective, CARAIB is more of an outlier.

One classical framework to look at vegetation changes over time, especially in relation to the human occupation Nikulina et al., 2024 is to express the vegetation in terms of openeness.

Fig 13 represents the percentage of herbaceous PFT in the LANDCLIM II dataset, taken here as an approximation for vegetation openness. The mean value in Europe is 23.7% at 8.5 k.a. (which is the period with the lowest openness), and 40.2% for the preindustrial. The only model approaching the 8.5 k.a. openness is ORCHIDEE-DGVM. The vegetation fraction of CARAIB cannot be used, as it is half tree half herbaceous covered everywhere, the openness is always equal to 50%, as a consequence of its two layers structure.

8.5 k.a. and 6 k.a. DGVM simulations are very similar for all models, which should be excepted, as the climate was very similar in those period in the northern hemisphere Borgatti and Soldati, 2013. Temperate zones seems to be well represented by all models but, models show big differences in boreal zones. In particular, CARAIB shows almost no trees northward from a latitude of about 60° N.

Another aspect to consider is the differences in representation of the PFTs which greatly conditions the differences between models. Indeed, in ORCHIDEE-DGVM, the PFTs are represented by a surface covered by each and spatially separated (at least partially because in reality, in the DGVM there is a competition for light which therefore supposes an overlap) and by internally representing the functioning of an average plant. This is why the NPP of ORCHIDEE-DGVM needs to be multiplied by the maximum vegetation fraction to be compared with the NPP of SEIB-DGVM and CARAIB. On the other hand, SEIB-DGVM has a totally different way of representing vegetation since it does not have explicit an VEGETMAX the PFTs being intermingled. For CARAIB, there is no VEGETMAX either, and there is also as we saw a two-layer structure : one with herbs and shrubs, and one with trees.

6 Summary and perspectives

In our study, we ran three different DGVMs forced by simulated climates for five different time-slices, from 8.5 k.a. B.P to preindustrial, and compared their results to reclassified pollen-based reconstructions. The three models have statistically similar results compared to the pollen-based reconstructions, but they also differ a lot on certain regions. The Alps and Scandinavia are poorly described by the models, suggesting a problem in the description of cold and mountainous regions.

All those results suggest that SEIB-DGVM and ORCHIDEE-DGVM are complementary and that there is a benefit of them being used together in order to have a better description of the natural vegetation cover of the Holocene. Overall, our study suggests that CARAIB performs less well for simulating the PNV over the Holocene, as, by and large, it reproduces neither the PFT diversity nor the PFTs dominance geographical distribution. An interesting perspective would be to make the same study under the first part of the Holocene (from 12 k.a. BP to 9 k.a. BP), to see if the pathway is the same for the matching score between pollen-based reconstructions and DGVM models.

To improve on the latter, an objective methodology to build a continuous spatial reconstruction from the LANDCLIM II database, which will be the subject of a future study.

With such differences between DGVM model results, we can ask the impact such differences could have on climate models. Indeed, results such as snow layer, leading to a change in albedo, and canopy height, impacting the wind, could be different depending on the DGVM used in climate model. On further interesting aspect could be to work on adaptating SEIB-DGVM and/or ORCHIDEE-DGVM to run online within the same climatic model to assess the importance of climate vegetation coupling.

The already mentioned impact of the climate forcing could be also worth investigating by targeting one particular time window such as the mid-Holocene (similar to PMIP) and perform a large ensemble of multi-climate / multi-vegetation model combinations to assess in full the uncertainty arising from the modeling components on the simulated vegetation cover.

References

- Arthur, F., Roche, D. M., Fyfe, R., Quiquet, A., & Renssen, H. (2023). Simulations of the holocene climate in europe using an interactive downscaling within the iloveclim model (version 1.1). *Climate of the Past*, 19(1), 87–106. https://doi.org/10.5194/cp-19-87-2023
- Berger, A., & Loutre, M.-F. (1999). Parameters of the Earths orbit for the last 5 Million years in 1 kyr resolution [Supplement to: Berger, A; Loutre, M-F (1991): Insolation values for the climate of the last 10 million of years. Quaternary Science Reviews, 10(4), 297-317, https://doi.org/10.1016/0277-3791(91)90033-Q]. https://doi.org/10.1594/PANGAEA.56040
- Borgatti, L., & Soldati, M. (2013). Hillslope processes and climate change. *Treatise on Geomorphology*, 7, 306–319. https://doi.org/10.1016/B978-0-12-374739-6.00180-9
- Brierley, C. M., Zhao, A., Harrison, S. P., Braconnot, P., Williams, C. J. R., Thornalley, D. J. R., Shi, X., Peterschmitt, J.-Y., Ohgaito, R., Kaufman, D. S., Kageyama, M., Hargreaves, J. C., Erb, M. P., Emile-Geay, J., D'Agostino, R., Chandan, D., Carré, M., Bartlein, P. J., Zheng, W., ... Abe-Ouchi, A. (2020). Large-scale features and evaluation of the pmip4-cmip6 *midHolocene* simulations. *Climate of the Past*, *16*(5), 1847–1872. https://doi.org/10.5194/cp-16-1847-2020
- Caley, T., & Roche, D. M. (2013). δ¹⁸O water isotope in the iloveclim model (version 1.0) part 3: A palaeo-perspective based on present-day data–model comparison for oxygen stable isotopes in carbonates. *Geoscientific Model Development*, 6(5), 1505–1516. https://doi.org/10.5194/gmd-6-1505-2013

- Chini, L., Hurtt, G., & Frolking, S. (2014). Luh1: Harmonized global land use for years 1500-2100, v1. https://doi.org/10.3334/ORNLDAAC/1248
- Dallmeyer, A., Kleinen, T., Weitzel, N., & Cornou, M. E. (2022). The deglacial forest conundrum. https://doi.org/10. 5194/egusphere-egu22-6979
- Dallmeyer, A., Poska, A., Marquer, L., Seim, A., & Gaillard, M.-J. (2023). The challenge of comparing pollen-based quantitative vegetation reconstructions with outputs from vegetation models a european perspective. *Climate of the Past*, *19*(7), 1531–1557. https://doi.org/10.5194/cp-19-1531-2023
- Feurdean, A., Ruprecht, E., Molnár, Z., Hutchinson, S. M., & Hickler, T. (2018). Biodiversity-rich european grasslands: Ancient, forgotten ecosystems. *Biological Conservation*, 228, 224–232. https://doi.org/https://doi.org/10.1016/ j.biocon.2018.09.022
- Fisher, R. A., Koven, C. D., Anderegg, W. R. L., Christoffersen, B. O., Dietze, M. C., Farrior, C. E., Holm, J. A., Hurtt, G. C., Knox, R. G., Lawrence, P. J., Lichstein, J. W., Longo, M., Matheny, A. M., Medvigy, D., Muller-Landau, H. C., Powell, T. L., Serbin, S. P., Sato, H., Shuman, J. K., ... Moorcroft, P. R. (2018). Vegetation demographics in earth system models: A review of progress and priorities. *Global Change Biology*, 24(1), 35–54. https://doi.org/10.1111/gcb.13910
- François, L., Utescher, T., Favre, E., Henrot, A.-J., Warnant, P., Micheels, A., Erdei, B., Suc, J.-P., Cheddadi, R., & Mosbrugger, V. (2011). Modelling late miocene vegetation in europe: Results of the caraib model and comparison with palaeovegetation data [The Neogene of Eurasia: Spatial gradients and temporal trends -The second synthesis of NECLIME]. *Palaeogeography, Palaeoclimatology, Palaeoecology, 304*(3), 359–378. https://doi.org/https://doi.org/10.1016/j.palaeo.2011.01.012
- Githumbi, E., Fyfe, R., Gaillard, M.-J., Trondman, A.-K., Mazier, F., Nielsen, A.-B., Poska, A., Sugita, S., Woodbridge, J., Azuara, J., Feurdean, A., Grindean, R., Lebreton, V., Marquer, L., Nebout-Combourieu, N., Stančikaite, M., Tanţău, I., Tonkov, S., Shumilovskikh, L., & data contributors, L. (2022). European pollen-based reveals land-cover reconstructions for the holocene: Methodology, mapping and potentials. *Earth System Science Data*, 14(4), 1581–1619. https://doi.org/10.5194/essd-14-1581-2022
- Goosse, H., Brovkin, V., Fichefet, T., Haarsma, R., Huybrechts, P., Jongma, J., Mouchet, A., Selten, F., Barriat, P.-Y., Campin, J.-M., Deleersnijder, E., Driesschaert, E., Goelzer, H., Janssens, I., Loutre, M.-F., Morales Maqueda, M.-A., Opsteegh, T., Mathieu, P.-P., Munhoven, G., ... Weber, S. L. (2010). Description of the earth system model of intermediate complexity loveclim version 1.2. *Geoscientific Model Development*, 3(2), 603–633. https://doi.org/10.5194/gmd-3-603-2010
- Gronenborn, D., Horejs, B., Börner, M., & Ober, M. (2023). Expansion of farming in western eurasia, 9600 4000 cal bc (update vers. 2023.1). https://doi.org/10.5281/zenodo.5903164.
- Hargreaves, J. C., Annan, J. D., Ohgaito, R., Paul, A., & Abe-Ouchi, A. (2013). Skill and reliability of climate model ensembles at the last glacial maximum and mid-holocene. *Climate of the Past*, 9(2), 811–823. https: //doi.org/10.5194/cp-9-811-2013
- Haxeltine, A., & Prentice, I. C. (1996). BIOME3: An equilibrium terrestrial biosphere model based on ecophysiological constraints, resource availability, and competition among plant functional types. *Global Biogeochemical Cycles*, *10*(4), 693–709. https://doi.org/10.1029/96GB02344
- Hopcroft, P. O., Valdes, P. J., Harper, A. B., & Beerling, D. J. (2017). Multi vegetation model evaluation of the green sahara climate regime. *Geophysical Research Letters*, 44(13), 6804–6813. https://doi.org/https: //doi.org/10.1002/2017GL073740
- Lange, S. (2019). Earth2observe, wfdei and era-interim data merged and bias-corrected for isimip (ewembi). v. 1.1. gfz data services. https://doi.org/https://doi.org/10.5880/pik.2019.004
- Levavasseur, G., Vrac, M., Roche, D. M., Paillard, D., & Guiot, J. (2013). An objective methodology for potential vegetation reconstruction constrained by climate. *Global and Planetary Change*, 104, 7–22. https://doi.org/ https://doi.org/10.1016/j.gloplacha.2013.01.008
- Li, H., Renssen, H., & Roche, D. M. (2019). Global vegetation distribution driving factors in two dynamic global vegetation models of contrasting complexities. *Global and Planetary Change*, 180, 51–65. https://doi.org/https: //doi.org/10.1016/j.gloplacha.2019.05.009
- Lu, Z., Miller, P. A., Zhang, Q., Zhang, Q., Wârlind, D., Nieradzik, L., Sjolte, J., & Smith, B. (2018). Dynamic Vegetation Simulations of the Mid-Holocene Green Sahara. *Geosphysical Research Letters*, 45(16), 8294– 8303. https://doi.org/10.1029/2018GL079195
- Moen, A., & Lillethun, A. (1999). National atlas of norway : Vegetation. Norwegian Mapping Authority.
- Nikulina, A., MacDonald, K., Zapolska, A., Serge, M. A., Roche, D. M., Mazier, F., Davoli, M., Svenning, J.-C., van Wees, D., Pearce, E. A., Fyfe, R., Roebroeks, W., & Scherjon, F. (2024). Hunter-gatherer impact on european interglacial vegetation: A modelling approach. *Quaternary Science Reviews*, 324, 108439. https://doi.org/https: //doi.org/10.1016/j.quascirev.2023.108439
- Petit, J.-R., & Raynaud, D. (2020). Forty years of ice-core records of CO₂. *Nature*, 579(7800), 505–506. https: //doi.org/10.1038/d41586-020-00809-8

- Quiquet, A., Roche, D. M., Dumas, C., & Paillard, D. (2018). Online dynamical downscaling of temperature and precipitation within the *i*loveclim model (version 1.1). *Geoscientific Model Development*, *11*(1), 453–466. https://doi.org/10.5194/gmd-11-453-2018
- Robakowski, P., & Bielinis, E. (2017). Needle age dependence of photosynthesis along a light gradient within an abies alba crown. Acta Physiologiae Plantarum, 39(3). https://doi.org/10.1007/s11738-017-2376-y
- Sato, H., Itoh, A., & Kohyama, T. (2007). Seib–dgvm: A new dynamic global vegetation model using a spatially explicit individual-based approach. *Ecological Modelling*, 200(3), 279–307. https://doi.org/https://doi.org/10.1016/j. ecolmodel.2006.09.006
- Sato, H., Shibuya, M., & Hiura, T. (2023). Reconstructing spatiotemporal dynamics of mixed conifer and broadleaved forests with a spatially explicit individual-based dynamic vegetation model. *Ecological Research*. https://doi.org/10.1111/1440-1703.12385
- Serge, M.-A. (2023). Spatially extensive and temporally continuous three REVEALS pollen-based vegetation reconstructions in Europe over the Holocene. https://doi.org/10.48579/PRO/J5GZUO
- Serge, M.-A., Mazier, F., Fyfe, R., Gaillard, M.-J., Klein, T., Lagnoux, A., Galop, D., Githumbi, E., Mindrescu, M., Nielsen, A., Trondman, A.-K., Poska, A., Sugita, S., Woodbridge, J., Abel-Schaad, D., Åkesson, C., Alenius, T., Ammann, B., Andersen, S., ... Zernitskaya, V. (2023). Testing the effect of relative pollen productivity on the reveals model: A validated reconstruction of europe-wide holocene vegetation. *Land*, 12(5). https://doi.org/10.3390/land12050986
- Strandberg, G., Lindström, J., Poska, A., Zhang, Q., Fyfe, R., Githumbi, E., Kjellström, E., Mazier, F., Nielsen, A. B., Sugita, S., Trondman, A.-K., Woodbridge, J., & Gaillard, M.-J. (2022). Mid-holocene european climate revisited: New high-resolution regional climate model simulations using pollen-based land-cover. *Quaternary Science Reviews*, 281, 107431. https://doi.org/https://doi.org/10.1016/j.quascirev.2022.107431
- Tallavaara, M., Luoto, M., Korhonen, N., Järvinen, H., & Seppä, H. (2015). Human population dynamics in europe over the last glacial maximum. *Proceedings of the National Academy of Sciences of the United States of America*, 112(27), 8232–8237. https://doi.org/10.1073/pnas.1503784112
- Tong, S., Wang, W., Chen, X., Jie, Chong-Yu, Sato, H., & Wang, G. (2022). Impact of changes in climate and co₂ on the carbon storage potential of vegetation under limited water availability using seib-dgvm version 3.02. *Geoscientific Model Development*, 15(18), 7075–7098. https://doi.org/10.5194/gmd-15-7075-2022
- Zapolska, A., Serge, M. A., Mazier, F., Quiquet, A., Renssen, H., Vrac, M., Fyfe, R., & Roche, D. M. (2023). More than agriculture: Analysing time-cumulative human impact on european land-cover of second half of the holocene. *Quaternary Science Reviews*, 314, 108227. https://doi.org/10.1016/j.quascirev.2023.108227
- Zhu, D., Ciais, P., Chang, J., Krinner, G., Peng, S., Viovy, N., Peñuelas, J., & Zimov, S. (2018). The large mean body size of mammalian herbivores explains the productivity paradox during the Last Glacial Maximum. *Nature Ecology & Evolution*, 640–649. https://doi.org/10.1038/s41559-018-0481-y

7 Supplementary

7.1 Mediterranean PFT for SEIB-DGVM

Parameters for the new mediterranean PFT of SEIB-DGVM are listed in Table 7.

Tmin	2.0
Tmax	48.0
Topt	25.5
Stat water min	0.05
Stat water max	0.75
Stat water opt	0.6
TCmin	3.0
TCmax	45.0
GDDmin	2200
GDDmax	20000

Table 7: Parameters of the new mediterranean PFT added to SEIB-DGVM

7.2 Calculation for astrometric parameters added to SEIB-DGVM

We started by using the Kepler equation :

$$E - e\sin E = M \tag{1}$$

With E the eccentric anomaly, and M the mean anomaly : $M = \frac{2\pi}{T}(t - t_0)$ with T the orbital period and t_0 the moment where the Earth is at the perihelion (minimal distance to the Sun). The Kepler equation can be resolved usind the following iteration :

$$E_0 = M \tag{2}$$

$$E_i = M + e \sin E_{i-1} \tag{3}$$

For a terrestrial eccentricity, which remains under 0.06 for the whole Holocene, only a few iterations are necessary (error $< 5.10e^{-11}$ for 7 iterations; $< 10e^{-14}$ for 10 iterations). As

$$\tan\frac{E}{2} = \sqrt{\frac{1-e}{1+e}} \tan\frac{v}{2},\tag{4}$$

we can obtain the real anomaly, which is the position relative to the perihelion on the orbit, v, from E. $v = \lambda - \pi - \frac{\pi}{180}\omega$, with λ the real longitude. That way we can get the solar declination :

$$\sin s l_{dec} = \sin \lambda \sin \epsilon \tag{5}$$

But in order to use the equation 2, we need to know t_0 . At the equinox (defined as the 81st day of the calendar) :

$$v_{equinoxe} = -\pi - \frac{\pi}{180}\varpi\tag{6}$$

Using 4 we obtain the eccentric anomaly at the equinox, E_0 , and thus, using 2, the mean anomaly at the equinox M_0 . Finally, as we know that the equinox is define as the 81st day of the year, we obtain the day of the perihelion t_0 .

Concerning the Earth-Sun distance on a given day of a year, it can be obtain using the first Kepler law :

$$r = a \frac{1 - e^2}{1 + e \cos \theta} \tag{7}$$

where r is the distance between the Earth and the Sun, a is the length of semi major axis of the Earth's orbit and θ is the angle subtended at the Sun between the semi major axis line and the current position. For our low eccentricity orbit, days can be used instead of θ , so long as we divide by the number of days in a sidereal year, and multiplied by 360 as we need degrees. Hence, $\theta = \frac{360}{365.256363}(doy - t_0) = 0.9856(doy - t_0)$. As $\frac{1}{1+x} \approx 1 - x$ for small values of x, and $a(1 - e^2) = 1$ in astronomical unit, the final formula is :

$$r = 1 - e\cos 0.9856(doy - t_0) \tag{8}$$

7.3 LAImax repartitions for all three models



Figure 14: LAImax values repartition for all point for SEIB-DGVM (left), ORCHIDEE-DGVM (middle), and CARAIB (right) for 8.5 k.a. (top), 6 k.a. (middle) and preindustrial (bottom) climatic dataset