PAST PLANT DIVERSITY AND CONSERVATION

EDITORS
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News

6th Open Science Meeting and 4th Young Scientists Meeting
PAGES is proud to announce the next meetings will be held in Agadir, Morocco, from 16-22 May 2021. The winning bid was submitted by a consortium from Université Ibn Zohr, Sultan Moulay Slimane University and Mohamed VI Polytechnic University. They are now being assisted by Universiapolis for the events’ organization.

Held once every four years, the OSM and YSM are the premier events in PAGES’ calendar and provide an invaluable opportunity to bring the international past global change community together to share, discuss, learn, and plan for the future.

Abstract submission and registration opens 1 July 2020. We look forward to seeing you there! All details: pages-osm.org

Rescheduling of workshops/meetings
The COVID-19 pandemic has resulted in several PAGES workshops and supported meetings being cancelled or postponed – groups affected include ACME, C-PEAT, CVAS, LandCover6k, PEOPLE 3000 and potentially more. Please check PAGES calendar for updated news and information regarding rescheduling: pastglobalchanges.org/calendar/upcoming

Working group news
Two working groups have started their second phases:
Climate Variability Across Scales (CVAS; pastglobalchanges.org/cvas)
Spleleothem Isotopes Synthesis and Analysis (SISAL; pastglobalchanges.org/sisal)

Three groups recently sunsetted:
Paleoclimate Reanalyses, Data Assimilation and Proxy System modeling (DAPS)
Ocean Circulation and Carbon Cycling (OC3)
Global Paleofire Working Group 2 (GPWG2)
Find out more about them here: pastglobalchanges.org/science/wg/former

Did you know PAGES working groups have designated mailing lists? Stay up to date and be involved in their activities. Find all the active mailing lists here: pastglobalchanges.org/science/wg/intro

PAGES Early-Career Network (ECN)
Abstract submission closes 8 May and registration opens 2 June for the joint PAGES-INQUA ECR Workshop "Past Socio-Environmental Systems" (pASES) from 9-10 September 2020. All details: pages-osm.org

PAGES Abstract submission closes 8 May and registration opens 2 June for the joint PAGES-INQUA ECR Workshop "Past Socio-Environmental Systems" (PASES) from 7-13 November 2020 in La Serena, Chile: pastglobalchanges.org/calendar/2020/127-pages/1972

All details of ECN activities can be found at: pastglobalchanges.org/ecn

Deadline for new working groups and workshop financial support
The next deadline to propose a new PAGES working group or to apply for financial support for a meeting, workshop, or conference is 16 September 2020: pastglobalchanges.org/my-pages/introduction

Thanks to financial support from the Chinese Academy of Sciences (CAS) and the Swiss Academy of Sciences (SCNAT), PAGES can assist in the running of paleoscience meetings. Applications should be received at least six months in advance of the planned event. All details: pastglobalchanges.org/my-pages/meeting-support

PAGES IPO staff update
PAGES’ International Project Office recently welcomed Monika Hofer, who replaced Alexandra Gerber as Finance and Office Manager. Her contact details can be found here: pastglobalchanges.org/about/structure/international-project-office

Help us keep PAGES People Database up to date
Have you changed institutions or are you about to move? Please check if your details are current: pastglobalchanges.org/people/people-database/edit-your-profile. If you have problems updating your account, we can help. Contact pages@pages.unibe.ch

Hard copy or email notification of Past Global Changes Magazines
In an effort to reduce our carbon footprint, PAGES would like to know if you would prefer to receive an email notification for future Past Global Changes Magazines instead of hard copies. If you would like to change, please send an email to pages@pages.unibe.ch and we will update your preferences, or you can easily update your own People Database account: pastglobalchanges.org/people/people-database/edit-your-profile

Upcoming issue of Past Global Changes Magazine
Our next magazine, guest edited by members of PAGES’ CRIAS working group, is titled “State of the Art and New Perspectives of Historical Climatology”. Although preparations are well underway, if you would like to contribute please contact our Science Officer: sarah.eggleston@pages.unibe.ch

Calendar

PALSEa workshop:
Improving understanding of ice-sheet and solid-Earth processes driving paleo se-level change
14-16 September 2020 – Palisades, NY, USA

QUIGS workshop:
Glacial Terminations: processes and feedbacks
22-24 September 2020 – Cassis, France

VICS workshop:
Moving forward by looking back
30 September-2 October 2020 – Aarhus, Denmark

LandCover6k 4th General Workshop
9-10 October 2020 – Philadelphia, PA, USA

Featured products

In December 2018, GEOTRACES and PAGES held a joint synthesis workshop in France. The first resulting paper is available in Paleoceanography and Paleoclimatology: pastglobalchanges.org/products/12969

PAGES was involved in a Nature Sustainability paper which proposes a set of four general principles that underlie high-quality knowledge co-production for sustainability research: pastglobalchanges.org/products/12966

LandCover6k
A new paper from Sandy Harrison et al. presents the overall strategy of the LandCover6k working group plus the products and their current and potential uses: pastglobalchanges.org/products/13016

Furong Li et al. released the first pollen-based REVEALS reconstruction of Holocene land-cover change in temperate China: pastglobalchanges.org/products/12977

GPWG2
Members of GPWG2, together with another 30 international colleagues, released a Policy Brief in December 2019 discussing how the integration of traditional ecological knowledge and long-term ecology could better support the evidence base for future decisions on fire policy and biodiversity conservation. Access the Policy Brief here: pastglobalchanges.org/gpwg2

Access the accompanying Future Earth article here: futureearth.org/publications/issue-briefs/global-fires/

PlioVAR
PlioVAR members published the paper “Late Pliocene Cordilleran Ice Sheet development with warm northeast Pacific sea surface temperatures” in Climate of the Past: pastglobalchanges.org/products/12979

Varves Working Group
A Quaternary special issue aims to present the diversity within the field and the state-of-the-art research on lake varves at all timescales and environments: pastglobalchanges.org/products/special-issues/13017

Cover

Fascinating tropical ecosystems:
A Baobab tree (Adansonia sp.) surrounded by herbs from the Lamiaceae family (Leucas martincens). Photo credit: Rachid Cheddadi.
Past plant diversity changes and future conservation issues

Rachid Cheddadi1, S. Báez2, S. Normand3, D. Payne4 and P. Taberlet5,6

Past climate changes have shaped the distribution of species across the planet by impacting their dispersal abilities, their migration rates, and the ongoing competition between and among them, as well as by triggering their local or total extinctions. This, in turn, has led to major changes in community composition and to ecosystem turnovers. Ongoing climate change is affecting species as well, and will keep doing so in the future. However, the challenge that we face today is that climate change is happening much faster than in the past and exceeding the migration rates of many species, driving a need for informed management to support the natural ability of species to respond. Moreover, climate change is no longer the only important factor in determining biodiversity patterns.

Changes in diversity are driven by complex interactions between climate change and increasing human pressures, whether from large-scale political and socio-economic transformations, overexploitation and unsustainable management of natural resources, or demographics. The accelerated growth of the human population, for example, is likely to dramatically reduce and fragment the range of species’ natural habitats, thereby threatening their potential for long-term persistence. In this modern context, many plant species will have to adapt locally or will become extinct because their natural dispersal to more suitable habitats will be compromised.

The evolutionary history of plants and their communities and of their response to environmental drivers started long ago and has followed various pathways. Accordingly, there are as many explanations for current patterns of diversity as there are species, and each species has a unique history among its coevals. Over geological time, certain plant species have adapted locally or migrated to more suitable habitats while others are now extinct because of their low dispersal abilities and/or reduced capacities for adaptation to new environments (e.g. the Atlas cedar in the Rif Mountains in Morocco; Fig. 1). For example, during the last glacial period, many temperate plant species persisted with low densities and/or with restricted ranges in so-called “refugia”, many of which are found in mountains. Phylogeographical studies have shown that many of these glacial refugia are hotspots of genetic and species diversity that result from a high relative abundance of endemic and ancestral genes and marked genetic differentiation at the local scale.

Relicts of the past can provide us with fascinating and useful information on how species reacted to different climatic situations, which in turn could inform management strategies to protect the remaining species on Earth now and in the future. For example, fossil records contain a wide range of proxies for reconstructing past climates, species occurrences, their range of fluctuations, and their extinction through time. Accordingly, studies combining surveys of modern DNA and fossil records allow us to trace Quaternary population and range dynamics with unprecedented detail, while studies on ancient DNA extracted from geological records are providing new insights into species adaptation to environmental changes.

Models offer another way to inform management strategies and species’ conservation policies. Modeling the ancient distribution of plant species illustrates potential migration pathways and refugia, and can increase our understanding of the demographic history and current genetic structure of species. However, numerous decisions and challenges accompany such modeling efforts, including the choice of modeling paradigm and algorithm, biases in algorithms and distribution data, variability in estimates of paleoclimate, means of model validation, genetic heterogeneity and model equilibrium, estimates of dispersal capacities, and our understanding of the liability of species’ environmental tolerance and biotic interactions.

Management strategies and species’ conservation policies can certainly be optimized through the integration of knowledge from different sources, disciplines, and timescales. However, the implementation of such strategies requires multi-scale, multi-species, and multi-disciplinary approaches to evaluate the relationship between long-term climate changes and biodiversity.

In this issue we present highlights from research on past climate changes, genetic approaches, and vegetation modeling to examine the relationship between past environmental changes and their impacts on different aspects of plant species’ diversity. We then discuss how such approaches could inform conservation and management.

The articles in this issue fall into four main sections: (1) Issues related to past climate variability in terms of amplitude, velocity, and abruptness, and how they have affected where species have persisted during climatically unsuitable time periods (Braconnot and Vimeux p. 4; Zheng et al. p. 6; Cheddadi and Bennett p. 8), (2) Impacts of long-term isolation and migration of species on their genetic diversity (Ficetola and Taberlet p. 10; Alsos et al. p. 12; Paruducci p. 14), (3) Models of past species’ distributions and biases to modeling approaches (François and Hambuckers p. 16), and finally (4) Lessons from the past to inform plant species conservation in the future (Payne et al. p. 18; Dirk and Gillson p. 20; Sarmiento et al. p. 22; Castilla-Beltrán et al. p. 24).

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Figure 1: The fossil record shows that Atlas cedar (Cedrus atlantica), which currently is endangered, formerly comprised significantly more extended forests in the Rif Mountains (Morocco). This picture shows a relict population of the Atlas cedar at ~1800 m above sea level in the Rif Mountains. Snow still occurs in winter but the increase of winter temperature and the steady decrease of annual precipitation over the past few decades is threatening the local persistence of the Atlas cedar in North Africa.
Past and future contexts for climate and water-cycle variability, and consequences for the biosphere

Pascale Braconnot\(^1\) and Françoise Vimeux\(^1,2\)

The Earth’s climate has changed in the past and is expected to continue to change in the future with implications for water resources, ecosystems, and human societies. Climate models are now reaching a level of complexity that makes them more useful to understand the linkages between climate and environment, and to connect global and regional changes.

**Temperature projections for the 21st century**

Human activity, by modifying the atmospheric composition (greenhouse gases and aerosols) and the land surface (agriculture and deforestation), affects the Earth’s energetics. Future perturbations are characterized by Representative Concentration Pathways (RCPs) indicating the amount of energetic change by 2100. Simulations with climate models able to predict the evolution of the climate system on timescales of seasons to decades and longer have become references to inform decisions regarding adaptation and mitigation. Temperature projections exhibit robust spatial patterns and consistent upward evolution over the 21st century. They all show a gradual increase in temperature everywhere, regardless of which RCPs are considered, with more pronounced changes in high latitudes than in the tropics. In all projections, the magnitude of changes after 2050 depends on the future greenhouse gas emissions and changes in land use. Note that uncertainties associated with model output for a given emission pathway are actually lower than the uncertainties concerning the evolution of human activities.

**Precipitation projections for the 21st century**

While water vapor increases in the atmosphere by about 7% for each degree Celsius of warming, following the Clausius-Clapeyron thermodynamic law, global precipitation is estimated to increase at a lower rate of about 3%/ºC (Collins et al. 2013). This precipitation trend simulated over the 21st century is associated with large spatial disparities as shown in Fig. 1a (Chavallaz et al. 2016). Globally, we observe an increasing contrast between regions of moisture convergence and divergence: additional precipitation is expected in tropical and mid-to-high latitudes (up to 50% more compared to the end of the 20th century), whereas less precipitation is simulated in subtropics. This arises from atmospheric thermodynamics and changes in the atmospheric dynamics that lead to the counterintuitive result of increasing heat export from the tropics associated with a slowdown of the Hadley-Walker circulation (Chadwick et al. 2013). Some regions also exhibit more pronounced trends, such as the drying of the Mediterranean area (by as much as 30% less precipitation). Still positive, the rate of precipitation changes throughout the 21st century clearly depends on the RCPs (Fig. 1b). Under RCP 2.6 (the strongest mitigation pathway), the rate of the precipitation change stabilizes both in drying and moistening regions, and even reduces at the end of the 21st century to return to historical values. In contrast, for RCP 8.5 (considered as the “business-as-usual” emission pathway), the rate of the precipitation change continuously increases during the 21st century (Fig. 1b), both in drying and moistening regions, leading to a doubling of the rate of change in mean precipitation by 2080. This reflects an acceleration of precipitation changes regardless of if it is in a regressing or a moistening region. As a consequence of small discrepancies between precipitation rate in moistening and drying regions (Fig. 1b), the fraction of moistening regions changes during the 21st century: under RCP 8.5, it tends to increase slightly by a few percent and to stabilize in the mid-21st century whereas under RCP 2.6, moistening region area clearly decreases by almost 10% by 2080. In addition, as stated in the 5th Assessment Report of the IPCC (IPCC 2013), there are regions and climate variables for which model projection remains particularly uncertain. This is the case for tropical precipitation projections, which are critically important for assessing the social impacts of climate change (tropical areas are among the most populated and the most vulnerable regions of the world), but for which no consensus emerges among climate models (Meehl and Washington 2012). The large spread among models illustrates important differences in climate sensitivity and biases (Dufresne and Bony 2008). Over tropical continents, not only the magnitude of the changes differs among models but also their sign (e.g. Douville et al. 2006; Sun et al. 2007; IPCC 2013).

As a consequence of climate change, vegetation is also expected to evolve during the 21st century with a potential decrease of the forest cover (e.g. in the tropics) or increase (e.g. in high latitudes). As a consequence of the expected climate change, some plant species are also expected to expand northward and towards higher altitudes with a modified growth season (e.g. Harsch et al. 2009; Savage and Vellend 2015; Pecl et al. 2017). Furthermore, simulations with interactive vegetation have confirmed the observed biophysical feedbacks associated with large-scale changes in vegetation.

**Future changes in relation to past changes**

These rapid changes are part of a longer history that has shaped our modern environment, and for which the major forcing was not trace gases such as CO\(_2\), but changes in the incoming solar radiation at the top of the atmosphere related to slow changes in the Earth’s orbit. The current generation of global climate models, such as the one used for the future climate projections presented above, can now be used to investigate the interplay between the long-term evolution of the external forcing, temperature, precipitation, and vegetation in different regions, though still in a simplified way (Braconnot et al. 2019; Dallmeyer et al. 2020). Figure 2 illustrates this long-term evolution. The IPSL climate model was only forced by Earth’s history that has shaped our modern environment, and for which the major forcing was not trace gases such as CO\(_2\), but changes in the incoming solar radiation at the top of the atmosphere related to slow changes in the Earth’s orbit. The current generation of global climate models, such as the one used for the future climate projections presented above, can now be used to investigate the interplay between the long-term evolution of the external forcing, temperature, precipitation, and vegetation in different regions, though still in a simplified way (Braconnot et al. 2019; Dallmeyer et al. 2020). Figure 2 illustrates this long-term evolution. The IPSL climate model was only forced by Earth’s climate sensitivity and biases (Dufresne and Bony 2008). Over tropical continents, not only the magnitude of the changes differs among models but also their sign (e.g. Douville et al. 2006; Sun et al. 2007; IPCC 2013). As a consequence of climate change, vegetation is also expected to evolve during the 21st century with a potential decrease of the forest cover (e.g. in the tropics) or increase (e.g. in high latitudes). As a consequence of the expected climate change, some plant species are also expected to expand northward and towards higher altitudes with a modified growth season (e.g. Harsch et al. 2009; Savage and Vellend 2015; Pecl et al. 2017). Furthermore, simulations with interactive vegetation have confirmed the observed biophysical feedbacks associated with large-scale changes in vegetation.
Figure 2: Evolution of climate conditions and vegetation over Europe, as simulated over the last 6000 years with the IPSL model. The different panels from top to bottom represent (A) the evolution of the temperate and (B) boreal tree fractions, expressed as a ratio to their initial cover; (C) parallel evolution of the total area (% of total land surface) covered by tree, grass, and bare soil, expressed as a deviation from the first 100-year average of the total area covered in the initial mid-Holocene state; (D) annual mean, annual monthly minimum (driest month), and annual monthly maximum (wettest month) precipitation (mm/day), and (E) annual mean, annual monthly minimum (coldest month), and annual monthly maximum (warmest month) 2-m air temperature. All curves have been smoothed by a 100-year running mean. The gray bar shows the pre-industrial period in 1750.

However, the pace of change varies with time, and the largest trends are found between about 5000 years BP and 3000 years BP. During the last 200 years of the simulation, the rapid increase of greenhouse gases (mainly CO₂) drives rapid changes in climate variables and vegetation that counteract their long-term evolution; this represents about half of the signal for climate and, in the case of total tree cover, most of the signal for vegetation over Europe. This stresses the fact that the linkages between ecosystems and climate cannot simply be deduced from past trends behavior, and that major differences are expected in the interplay between climate and vegetation in the future, with many unknowns compared to past climate evolution.

Another important message is that periods of several decades or centuries are characterized by both large climate and vegetation variations. Even though most of the curves in Figure 2 exhibit similar large variations, there is no systematic correspondence in the relative magnitude and duration of the temperature and precipitation events, or between climate and vegetation fluctuations. One reason is that climate variations need to be large enough to cross a threshold and impact the environment, and that different types of ecosystems react to different thresholds. In this respect, the total tree fractions over Europe also mask different dynamics between the different tree plant functional types (pfts). The different simulated pfts provide a more dynamic view of changes in ecosystem diversity. In this simulation, only the temperate broad-leaved summer tree pft declines with time, whereas the other pfts show a more chaotic behavior and do not systematically exhibit synchronous variability. They respond differently to the different events, depending on their sensitivity to temperature or precipitation or to specific patterns of climate events over Europe that may or may not affect their geographical range.

Even though such long simulations are still imperfect, and model biases are large and prevent detailed explanations of reconstructed changes, they highlight the different spatio-temporal scale that need to be accounted for to properly anticipate future climate change. Introduction of land use is also an important factor to better characterize the Holocene impact of human activity in addition to the rapid increase in greenhouse gases. This additional human effect is modifying regional climate characteristics (Harrison et al. 2020). Ensembles of such long simulations will soon be available to the community, providing new opportunities to understand the evolution of both the water cycle and ecosystems.

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How the Asian subtropical area became evergreen since the Last Glacial Maximum

Zhuo Zheng1, K. Huang1, Q. Wan1 and R. Cheddadi2

The onset of the evergreen forest expansion throughout subtropical eastern Asia took place progressively from the tropical latitudes after the Younger Dryas and ended between 8000 and 6000 years BP.

Eastern subtropical Asia (ESA) is a territory about the size of Europe and corresponds to a vast ecolonal area between the tropical and the northern temperate climate systems. ESA underwent a tremendous biome turnover over the last glacial/interglacial transition (Ni et al. 2010). Today, the evergreen broad-leaved (EBL) biome prevails in ESA, and during the last glacial period it was dominated by a fragmented deciduous broad-leaved (DBL) vegetation (Yue et al. 2012) over the mountain landscapes and open vegetation at lower altitudes (Yu et al. 2017). The EBL biome (known as laurophyllous forest), currently the most widely distributed biome in eastern Asia, extends from about 22°N to 33°N under a monsoon regime.

Timing and dynamics

Although the forest was not the only dominant ecosystem during the last glacial period, the DBL forest covered a large part of southern China that is dominated today by the EBL biome (Fig. 1a and 1b). A high-resolution pollen record collected in the center of the subtropical zone shows that until the last Heinrich event (H1, dated to around 16.5 thousand years before present, kyr BP) the main forest populations in southern China were composed of deciduous rather than evergreen species (Fig. 1c). However, the complete absence of the EBL biome north of 24°N during the Last Glacial Maximum (LGM; Ni et al. 2010) seems to not be coherent with the phylogeographic studies that suggest some species composing the EBL biome survived during the LGM in situ in numerous northern refugia (Fan et al. 2016).

The postglacial expansion of the EBL biome over the subtropical zone primarily began at the Bølling/Allerød (B/A) warming (around 14 kyr BP), but more substantial replacement occurred after the Younger Dryas (YD) cooling (12.7-11.5 kyr BP). A major massive migration took place from the southern tropical belt of eastern Asia rather than from the putative scattered northern refugia, suggested by the phylogeographic studies. In the western high Tibetan plateau, within the subtropical zone, despite a different vegetation type during the LGM (sclerophyllous mixed oak and conifer forests), there is also simultaneously a profound biome change over the altitudinal belts (Wang et al. 2018; Nishimura et al. 2014; Cheng et al. 2018). The transitional turnover from DBL...
to EBL after the YD event is depicted in most records located north of the Tropic of Cancer. This post-YD large biome turnover originated in the northern tropical boundary of eastern Asia and progressively spread throughout all of subtropical China. The northernmost front of the EBL was reached around 8-7 kyr BP and corresponds to the northern limit of the East Asian Summer Monsoon. The progressive timing of the EBL expansion inferred from pollen records shows a significant delay with the speleothem oxygen isotope records from Dongge and Hulu caves from southern China.

The DBL/EBL biome turnover was quite synchronous over the same latitude, with a progressive time-lag from south (11.5 kyr BP) to north (8-6 kyr BP; Cheng et al. 2018; Zhou et al. 2016). Pollen data from the semi-arid belt in northern China under the influence of the summer monsoon also indicate a delayed expansion of the EBL biome during the early to mid-Holocene, which confirms the south-north progressive recolonization of the DBL biome over the ESA (Wang and Feng 2013).

The time-transgressive propagation of the EBL across the subtropical landscape stabilized during the mid-Holocene around 4 kyr BP. Pollen data relate an opening of the vegetation and the expansion of cultivated species towards the end of the Holocene that is probably related to a combined effect of a reduction of the annual amount of precipitation along with an increase in the regional anthropogenic disturbances. The modern range of the EBL biome in the subtropical zone seems to be constrained by the superimposition of the effects of increasing human activities (slash and burn, cultivation, etc.) and climate change.

The causes
The pattern of subtropical forest changes in response to the postglacial climate warming has drawn extensive attention (Lee and Liew 2010). The strengthening of the summer monsoon over the last post-glacial warming has been considered as the main factor driving large vegetation changes in eastern Asia (Zhou et al. 2016). Several speleothem records provide strong evidence for the strengthening of the monsoon system (Dykoski et al. 2005); however, an increase in precipitation (annual and/or seasonal) may not be sufficient to explain a complete replacement of the DBL by an EBL biome at the scale of ESA.

The temperature reconstruction based on biomarkers and fossil pollen data reveals a threshold determined by the mean annual temperature for the EBL/DBL biome transformation during the early Holocene (Wang et al. 2017; Fig. 2). Today, the mean minimum temperature seems to be a more accurate or appropriate climate variable for defining the distribution boundary of EBL forest in China. The annual and winter temperature isotherms (15-16°C and 6-8°C, respectively) delineates the northern limit of modern EBL forests in the subtropical mountains and corresponds to the transition between the DBL and EBL biomes. The temperature simulated by a transient model is consistent with the observed trend (Liu et al. 2009), and the combined data of biome transformation across the overall zone is coherent with the progressive increase in temperatures, which is controlled by winter insolation and seasonality during the last deglaciation to mid-Holocene. Thus, besides the strengthening of the monsoon, the turnover of EBL/DBL forest was probably extremely sensitive to both the seasonal amplitude and the long-term increase in winter and annual temperature during the early Holocene (Fig. 2).

The temperature increase and the seasonal change in precipitation are related to several global climate variables. The gradual increase in the sea surface temperature in the warm pool of the South China Sea and the northward cross-equatorial atmospheric heat transport during the early to mid-Holocene have played an important role in heating up the subtropical continent. Besides the impact on the monsoon system, the energy delivered to China increases de facto the inland temperature. The fact that the EBL forest recolonization was halted by the last Heinrich event and then the Younger Dryas, both of which are marked cool periods recorded in the North Atlantic, tends to suggest that global ocean circulation was also an important driving mechanism for the temperature changes over China, therefore impacting its large vegetation changes.

In conclusion, ESA underwent a large biome turnover that is clearly related to the post-glacial global climate changes, and the expected future climate changes will probably impact it again as was the case in the past.

ACKNOWLEDGEMENTS
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Past climate changes and the role of refugia in the temperate Northern Hemisphere

Rachid Cheddadi1 and Keith D. Bennett2

Glacial refugia in the temperate Northern Hemisphere proved to be havens for present ecosystems’ diversity. The modern microrefugia, where isolated populations of threatened plant species are persisting, might well play a similar role for preserving future biodiversity.

Classical and cryptic glacial refugia

Plant species’ potential range (fundamental niche) is delineated by a climate space that they often occupy only partially (realized niche). Generally, temperate plant species tend to occupy a wider geographical range when climate is warm and wet, and retract when climate is cold and dry. During the last glacial period, cold-tolerant herbs dominated the mid- to high-latitude landscape in both hemispheres and the range of most temperate tree species was reduced to discrete populations until the climate became warmer and wetter during the Holocene, which allowed them to expand.

The persistence or survival of plant species in restricted areas during past unfavorable climate time spans is a fundamental ecological pattern that has contributed to shaping the modern range of species, ecosystem composition, and species’ genetic diversity. Restricted areas in the Northern Hemisphere such as in the Mediterranean basin, known today as “classical southern refugia” (Bennett and Provan 2008), played a major role in the survival of European plant species during the last glacial period. These geographically restricted and scattered areas located at lower latitudes offered local (micro) climates that were suitable for plant species. However, other refugial spots designated as “cryptic refugia” have been identified at higher latitudes than the classical southern refugia in the Northern Hemisphere, namely in Sweden (Parducci et al. 2012), Beringia, North America (Provan and Bennett 2008), and northern China (Fan et al. 2016).

Refugial areas where plant species were present continuously over the last climatic cycle under different extreme climates (Tzedakis et al. 2002), or since just the last glacial period until today under the modern fast climate change, are of primary importance for potential future species’ conservation (Morelli et al. 2016). In southern Europe, scholars have shown that there is a strong relationship between the location of the classical glacial refugia and the modern areas where high biodiversity and endemism rates are measured (Fig. 1).

Where are the refugia?

Locating past refugia is also important for evaluating species’ migration rates, hindcasting the species’ range change through time using different climate scenarios, and testing and validating process-based vegetation models for competition between species. These species and/or vegetation-type models help to evaluate species’ diversity loss under different future climate scenarios, their capacity for migrating, and the potential future suitable habitats for their conservation.

During a change in global climate, the areas where local microclimates are found to play the role of refugia for the persistence of plant species over several centuries or millennia (Tzedakis et al. 2002). This could be the case during either glacial or interglacials (Bennett and Provan 2008). Refugial areas are often located in mountainous regions (Bennett et al. 1991), which offer a wide range of landscapes, geological substrates, and exposure time to the sun. Mountains with high rugosity may provide more microrefugial areas (Rull 2009) for plant species under regionally less favorable climate than areas with smoother landscapes (Valencia et al. 2016). In addition, mountains represent a much faster option for species to compensate for global climate change by migrating easily up or down towards cooler/warmer and/or wetter/drier altitudes.
The Föhn winds can raise temperature and bring moisture from the adjacent seas to mountains, which may create local microclimates. By migrating 1000 m in altitude, a species may compensate for a temperature change between 6 and 10°C, depending on the latitudinal location of the mountain (Bennett et al. 1991). Today, under ongoing global warming, many species persist as scattered populations in mountainous areas because of the existence of offset suitable microclimates. The estimated velocity of temperature change is lower within tropical and temperate mountain biomes than in lowlands (Loarie et al. 2009). Species respond individually to climate change, and their estimated past migratory rates seem to be slower than the modern climate change. Survival of many species under the ongoing climate change may rely on their ability to persist in some modern microrefugia (Rull 2009) with favorable (micro)climatic conditions.

**Glacial refugia and species’ modern genetic diversity**

Refugial areas in the past can be identified from paleorecords and genetic data (Fig. 2). Fossil remains, such as pollen grains and macro-remains, allow for the identification of the occurrence of a species and its time of persistence at a site or the area surrounding the study site. Phylogeographers and paleoecologists have worked together over the past few decades to identify glacial refugia and to reconstruct the pathways of species expansion, based on species haplotypes using different genetic markers (Petit et al. 2003) along with extended sets of fossil pollen records.

For many species, microrefugia with stable microclimates that are offset from the global climate may represent the only alternative to extinction if their migration rate is lower than the velocity of climate change (Hannah et al. 2014). Modern microrefugia may have an important role in preserving species under the ongoing global climate change (Cheddadi et al. 2017) and may represent a reservoir of biodiversity. Genetic diversity would be maintained within these isolated populations and perhaps even improved if there is a genetic flow between different populations.

Phylogeographic studies have highlighted the role played by the glacial refugia for species survival and their legacy in shaping their modern genetic diversity (Petit et al. 2003). Depending on the range size and the time span over which plant species remain isolated in a refugium, there may be a local genetic shift of the species. The genetic drift during this time may lead to reduced genetic polymorphism in different geographically scattered refugia. Consequently, differences between DNA haplotypes could appear in the different refugia, which may be traced back by genotyping individuals of the same species collected over its modern range.

**Potential contribution of modern refugia to future biodiversity**

Genetic studies have shown that glacial refugia do not necessarily harbor the most diverse populations. However, the post-glacial crossing of different lineages of the same species during the recolonization process from multiple refugia produced the most genetically diverse populations (Petit et al. 2003). Thus, even if the genetic diversity of a population tends to decrease over time when a species remains isolated in a refugium, its persistence into multiple scattered refugia may contribute to conserving and improving future species diversity. The future persistence of different species under the ongoing global warming and the preservation of biodiversity might well depend on the identification of potential modern refugia for threatened species and the protection of these refugia from anthropogenic degradation.

Identifying and protecting microrefugial areas may represent an additional option for species conservation (Hannah et al. 2014) which is likely to be one of the most efficient scientific solutions for species conservation with a reasonable social cost, especially in areas where local public resources are low.

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**Figure 2:** Multiple glacial microrefugia of different plant species around the Alps identified from (A) fossil records on different bedrocks (blue: calcareous; red: siliceous) that have been confirmed by (B) phylogeographic analyses, which provide more details about the different types of refugia (different colors on the map) and the species diversity that they shelter (modified from Schönswetter et al. 2005).
Next generation sequencing, phylogeography, and paleoecology

Gentile F. Ficetola and Pierre Taberlet

Genome skimming and environmental DNA extracted from lake sediments are increasingly important to measure genetic diversity, and to understand how environmental changes affected species distribution through time. Here we describe how genome skimming on the current geographic distribution of species can be combined with the analysis of lake sediments, to perform unprecedented tests of phylogeographic hypotheses, and reconstruct past refugia and colonization routes.

DNA sequencing has greatly improved over the past 15 years, and next generation sequencing is transforming DNA analysis (van Dijk et al. 2014). Today, the most powerful sequencer, the NovaSeq 6000 (Illumina), is able to produce up to 40 billions of sequence reads of 250 nucleotides in a single experiment. Such sequencing power induces a complete change in assessing genetic diversity of wild species. It is now affordable to analyze a significant portion of the genomes instead of looking only at a few target regions. Here, we will focus on two approaches linked to next generation sequencing: genome skimming and environmental DNA analysis; both can greatly improve our understanding of how environmental changes have impacted the genetic diversity of species.

Genome skimming (Coissac et al. 2016; Dodsworth 2015; Straub et al. 2012) corresponds to the sequencing of random DNA fragments of the genome, at a sequencing depth that does not allow for the assembly of the whole nuclear genome, but that allows for the assembly of all repetitive DNA such as, for plants, the chloroplast genome, the nuclear ribosomal DNA, and possibly the mitochondrial genome. Genome skimming has been proposed as an extension of the current barcoding approach (Coissac et al. 2016), as it provides much more information for DNA-based species identification and for phylogenies (e.g. Malé et al. 2014).

The analysis of environmental DNA extracted from lake sediments allows us to reconstruct the variation of species distribution over the last few millennia, and is becoming a widely used approach for assessing past communities, either targeting a single species, or dealing with all species from a taxonomic group such as plants or mammals (review in Bálint et al. 2018; see also Giguet-Covex et al. 2014; Pansu et al. 2015). Although most of the studies performed so far focused on individual species or on higher-level taxonomic entities, lake sediments could also be a source of information at the intraspecific level, provided that the appropriate DNA markers are available.

Combining genome skimming and environmental DNA

By combining genome skimming on the current geographic distribution of a target species, and the analysis of lake sediments, it is now possible to test intraspecific phylogeographic hypotheses, i.e. to find past refugia and colonization routes. Figure 1 illustrates this approach, which has not yet been implemented but has a great potential for understanding past distributions of different lineages within a species.

The first step must start with a phylogeographic study of the considered species. This consists of collecting representative samples over the whole current distribution (Fig. 1a). After DNA extraction, these samples are sequenced on a next generation platform, using a genome skimming approach, to produce about one gigabase of sequence, e.g. about seven million sequence reads, each 150 base pairs long. Based on these sequences, for each sample, the whole chloroplast DNA and the whole nuclear ribosomal DNA can be assembled. The comparison of these sequences among all samples reveals the phylogeographic structure of the species, showing the geographic distribution(s) of either a single homogeneous lineage (Fig. 1b), or different lineages (Fig. 1c).

The second step is the identification of genetic markers that unambiguously characterize each lineage, with the goal of being able to recover these markers from lake sediments. This requires choosing markers that are as short as possible, with conserved flanking regions where PCR (polymerase chain reaction) primers, i.e. short DNA sequences that are necessary for DNA amplification, can be anchored. As genome skimming provides the sequences of the whole chloroplast DNA and of the ribosomal DNA, it is quite easy to find such short diagnostic markers for each lineage.

The third step consists of collecting lake sediments, within both the current geographic distribution and potential refugia. The different cores must be precisely dated, and DNA extraction is carried out from many core slices. The objective is to test for the presence or absence of the different
lineages at different times and locations. This presence is determined using the DNA markers designed in step two. These markers are amplified via PCR and sequenced on next generation sequencers. The results should allow us to reconstruct the history of each lineage very precisely, including past presence in areas where it later disappeared, and colonization routes towards the current geographic distribution.

Perspectives

All the technologies and methodologies for carrying out the above approach have already been available for more than five years. Surprisingly, not a single paper has taken the opportunity to combine them for resolving important controversies concerning potential refugia, postglacial colonization routes, and the evolution of intraspecific genetic diversity. For example, the re-establishment of flora in Scandinavia is controversial, opposing the “tabula rasa” versus the “nunatak” hypotheses (Brochmann et al. 2003). The “tabula rasa” hypothesis suggests that after the last glacial maximum, Scandinavia was only recolonized from the South. Alternatively, the “nunatak” hypothesis opens the possibility for a recolonization from non-glaciated cryptic refugia in Scandinavia such as Andøya Island (Parducci et al. 2012). The recolonization of Scandinavia could be tested by collecting lake sediments in the plain south of the ice sheet during the last glacial maximum. If for many species the same lineages are found in Scandinavia and in these plains where they are currently absent, this will be in favor of the “tabula rasa” hypothesis. On the contrary, if the lineages in these plains about 20 thousand years ago are different from the ones found in Scandinavia, this will support the possibility of northern cryptic refugia.

Climatic oscillations occurring since the Pleistocene have shaped the present day biodiversity of plant and animal species; yet, human-driven global changes are leading to unprecedented changes of species distributions and diversity. Understanding the processes that have determined the present-day genetic diversity can also help us understand the biotic response to ongoing environmental changes, and identify appropriate management strategies.

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Learning from the past for a sustainable future
Survival and spread of arctic plants in response to climate change: DNA-based evidence

Inger G. Alsos¹, M.E. Edwards¹,² and C.L. Clarke²

Preserving biodiversity requires knowledge of the processes that underlie geographical patterns of species' distribution and sustain diverse communities over long periods of time. Studies of modern and ancient DNA may provide new insight into these processes.

The fate of the NW European biota during the last glacial period has been debated for almost 150 years. While the consensus in the 1960s was that at least the arctic-alpine elements of the flora survived in small, coastal, ice-free areas or on nunataks in Greenland, Iceland, Svalbard, or Fennoscandia, a more recent review concluded that all extant species colonized post-glacially (Brochmann et al. 2003).

These opposing views of arctic plant history are based on different perspectives on understanding the frequency of long-distance dispersal, evolution rate, and species' ability to persist under a changing climate. Molecular tools have shed new light on this debate.

By analyzing genetic variation within present-day populations (modern DNA studies), the relationship among populations in different geographical areas can be established. Further, based on the amount and spatial structure of genetic diversity, modern molecular data can also be used to infer areas of glacial persistence (i.e. where there is high genetic diversity and/or distinct genetic markers) and the direction of postglacial dispersal (Avise 2000). More recently, advances in the use of ancient DNA have greatly increased our ability to detect past species' distributions and responses to climate changes.

Lessons learned from studies of modern DNA

Studies of modern DNA have greatly advanced our knowledge of past dispersal. For example, genetic fingerprinting of 17 widespread arctic plant species throughout their current distributional range showed that the main barriers to dispersal were the Urals Mountains, the Greenland Ice Sheet, and the Atlantic Ocean, plus the lowland areas between the Alps and the Scandinavian Mountains (Eidesen et al. 2013). Alsos et al. (2015) extended the dataset to 25 species and used this in combination with floristic studies to infer the main dispersal routes in the north Atlantic region (Fig. 1).

These data show that long-distance (300-1000 km) dispersal in this region was frequent. In contrast, long-term isolation and evolution, as indicated by unique genotypes and high levels of genetic diversity for some species in some areas, appears to have been uncommon. Thus, current data indicate that postglacial dispersal may have been extensive while glacial survival was rare (Alsos et al. 2015; Westergaard et al. 2019). However, studies of modern DNA only give a relative age of past events and may overlook processes such as extinction of lineages or repeated dispersal. Recent advances in ancient sedimentary DNA have great potential to fill this knowledge gap (Parducci et al. 2017).

Combining modern DNA data with species' distributions may enable an estimation of genetic consequences of climate change on arctic plants in the past (Pellissier et al. 2016) and projection of future scenarios (Alsos et al. 2012). Past changes in species' ranges were associated with a genetic bottleneck effect: a shift in species' distribution by 1000 km is associated with an average loss of 21% of the genetic diversity. All species, even common ones, are expected to lose part of their current range as a consequence of ongoing global warming. This range loss is expected to cause loss of genetic diversity, with herbs lacking berries or other adaptations for long-distance dispersal being especially at risk.

What can we learn from studies of ancient DNA?

Ancient DNA can provide direct evidence for the occurrence of a plant species at a given site and time in the past and thus greatly improve our ability to reconstruct past glacial refugia and dispersal routes. While past presence of a given species may be indicated by the presence of macrofossils (e.g. Binney et al. 2009), arctic-alpine plant species are often poorly represented due to variable preservation; furthermore, they are often under-represented in pollen records because the majority of taxa are insect-pollinated, having poor pollen productivity and transport characteristics (Birks 2008).

Calibration studies have demonstrated that sediment DNA comes from within the local catchment area (Alsos et al. 2018) and that pollen is not likely to be a source of sediment DNA (Sjögren et al. 2017). Ancient DNA extracted from sediments (sedaDNA) has proven a useful new tool for reconstructing past plant community composition, particularly with regard to arctic-alpine herbs (Willerslev et al. 2014), and it can significantly augment information derived from pollen and plant macrofossil records (Alsos et al. 2016; Parducci et al. 2017).

In a recent study of a long (24 m) sediment core from Lake Bolsheyoye Shuchyhe in the Polar Ural Mountains of Arctic Russia, Clarke et al. (2019) assessed how the diversity of the arctic-alpine flora fared through large-magnitude climate changes over the past 24,000 years using sedaDNA analysis. The lake catchment has supported a typical arctic-alpine community over the last 24,000 years, while there has also been diversification through time to include shrub-tundra and boreal-forest taxa (Fig. 2).

The sedaDNA record provides robust empirical evidence of the persistence of the arctic-alpine flora through a long period of climate change. This global warming, this range loss is expected to cause loss of genetic diversity, with herbs lacking berries or other adaptations for long-distance dispersal being especially at risk.

Figure 1: (A) Post-glacial dispersal routes of arctic plants within the North Atlantic region based on amplified fragment length polymorphism fingerprinting studies of 25 species (Alsos et al. 2015). Red dots indicate potential glacial survival of single species (Alsos et al. 2015; Westergaard et al. 2019), whereas green shading indicates the main focus regions. Note that the dispersal routes in most cases are not along the shortest distance, which would be expected if dispersal was a random process. Thus, dispersal agents such as sea-surface circulation (B, blue = cold water, red = warm water) or birds (C, migration routes for geese species (blue) and snow bunting (red)) may have aided dispersal. Red dots indicate potential glacial survival of single species (Alsos et al. 2015; Westergaard et al. 2019), whereas green shading represents the main focus areas.
of environmental change, including the postglacial expansion of woody taxa and a period of forested conditions between 9,000 and 4,000 cal. years BP, when summer temperatures were likely warmer than present. The diversity and abundance of arctic-alpines did, however, decline as soon as shrubs expanded, suggesting that in a future warming scenario, loss of competition-sensitive arctic-alpine species may occur long before the establishment of boreal forest. Nevertheless, most of the plants that were present during the last glacial period, including all of the arctic-alpine plants, are still found in the region today, suggesting resilience to climate changes.

The Lake Bolshoye Shchuchye record highlights the conservation significance of spatially heterogeneous mountain landscapes such as the Polar Ural. Plants adapted to extreme conditions can be at high risk from climate change; arctic-alpine plants, in particular, could “run out of space” as they are out-competed by expansion of woody vegetation. However, heterogenous landscapes may provide a range of habitats for different taxa and a variety of communities. This in turn confers resilience in the face of climate change.

Future perspectives
As more sites are analyzed for ancient DNA, we can greatly increase our knowledge of the past distribution of arctic-alpine plants. This will allow us to more effectively combine studies of modern DNA with ancient DNA records to determine past species’ distributions and model likely future ones. Direct analyses of genotypes through time and space are now possible, allowing for the study of the genetic consequences of climate change (e.g. as applied to macrofossils of silver fir, Abies alba; Schmid et al. 2017). In the near future, similar studies based on ancient sedimentary DNA alone should be possible and potentially more widely applicable.

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Figure 2: The ancient sediment DNA record from Lake Bolshoye Shchuchye shows the survival of arctic-alpine plants during a period when the climate was warmer, and shrubs (light green shaded bar) and trees (dark green shaded bar) expanded (modified from Clarke et al. 2019). Each square represents one occurrence of the plant taxon, with a blue color ramp used to indicate how many of the eight replicates per sample the taxon was detected within.
Ancient DNA from lakes: Insights into plant response to past climate changes
Laura Parducci

Considerable advances during the last decade in DNA sequencing technology and bioinformatics have enabled in-depth study of the genome from a larger number of ancient samples derived from a larger variety of substrates and environments. In particular, extraction and analysis of ancient DNA from lake sediments has provided important clues regarding the evolution of past environments in relation to climate change. This information is essential for understanding the importance of climate and plants’ dispersal ability in determining their distributional ranges, and for predicting the response of plants to ongoing climate change.

Ancient lakes
Ancient lake sediments store biological, chemical, and physical information that allow for a detailed reconstruction of past environmental and climatic changes. Biological micro- and macrofossil remains, combined with adequate chronologies, provide not only information on changes in patterns, structure, and diversity of past ecosystems, but can also be used to reconstruct past temperature and precipitation. Despite a wealth of information on past environmental and climatic changes that has been obtained from lake sedimentary records during the past decades, paleoecologists are aware that fossil assemblages represent only a tiny fraction of all the organisms that existed at a certain time in the past (due to e.g. post-mortem processes, differential decomposition, or depositional changes). In addition, each paleoecological method has its specific limitations (e.g. diatoms may partly be dissolved; high pollen producers obscure the abundance of other plant species; some species are better preserved than others in sediments).

Large-scale sequencing data from lakes
Recent advances in ancient DNA (aDNA) analyses now make use of large-scale sequencing technologies, which parallelize the sequencing process and produce millions of sequencing reads simultaneously. In this way, DNA sequenced from soils offer an unprecedented opportunity to unravel a complementary and larger spectrum of plant taxa compared to those identified by micro- or macrofossil analyses alone (Taberlet et al. 2018). They also allow identification of plants at lower taxonomic levels than conventional paleoecological proxies (Willerslev et al. 2014; Pedersen et al. 2016; Parducci et al. 2019). However, DNA from soil mainly represents taxa growing a few meters from the sampling point (Edwards et al. 2018).

Lakes, on the other hand, have large catchment areas and recent studies have suggested that plant aDNA from lacustrine sediments is a more powerful tool for paleo floristic reconstruction (Parducci et al. 2017; Giguet-Covex et al. 2019). In particular, sediment records from small lakes are excellent archives for molecular studies, as the effects of disturbances are low and seem to represent the surrounding terrestrial environment well (Fig. 1). As such, aDNA studies of lake sediments have the potential to revise and complement our understanding of how plant species and communities responded in the past to long- and short-term environmental changes. At the species level, this allows us to assess the effect of climate change over hundreds of generations, while at the community level it permits the investigation of the structure of ancient communities (dominant versus rare species) and their correlation with temperature, precipitation, and biotic interactions. This information is essential for understanding the importance of climate and dispersal ability of plants in determining past ranges and for predicting the response of ranges to ongoing and future climate changes.

Plants’ responses to climate
There are several examples of questions that can be addressed by using aDNA analyses from lakes to tackle issues related to plants’ responses to climate and that can help us understand what we can do in the face of future changes. One in particular is the question concerning the Reid’s paradox of rapid migration of plants (Reid et al. 1899). This is the observation that the speed of forest recolonization process after the last Quaternary glaciation (from ca. 12 cal. kyr BP) is faster (115–550 m/yr; Feurdean et al. 2013) than it would be expected, based on life histories and ability to transport seeds and fruits of the species involved (Clark et al. 1998). Fast recolonization can be explained by rare long-distance dispersal events mediated by birds and wind and promise well for future populations suggesting that plants can do so again in response to global warming, but alternative explanations need to be investigated thoroughly.

One alternative explanation, though controversial (Tzedakis et al. 2013), is that cold-tolerant tree species survived at high latitudes in small isolated microrefugia and recolonized locally once the ice retreated (Stewart et al. 2010). Recently, several lines of evidence based on fossil analysis, species distribution modeling, and phylegeographical surveys have been used to propose the existence of these small microrefugia at northern latitudes in the Northern Hemisphere (e.g. Napier et al. 2019). In Scandinavia, in central Sweden, several megafossils of mountain birch (Betula pubescens Ehrh. ssp. tortuosa (Ledeb.) Nyman), Norway spruce (Picea abies (L.) Karst.), and Scots pine (Pinus sylvestris (L.)) have been recovered earlier than expected (14, 11, and 11.7 kyr BP, respectively; Kullman 2002), and in the same region today, hundreds of clonal P. abies trees are growing...
with plant material under their root systems dated up to 9.5 kyr BP (Fig. 2). The early presence of tree species at high latitudes suggests either an early Holocene arrival or survival during the last glaciation. These hypotheses were tested using aDNA extracted from Norwegian lakes, and the genetic findings showed that spruce trees were present in Norway during the late glacial and the early Holocene (Parducci et al. 2012), leaving open, however, the question of their origin.

Plant survival under changing climates

These findings question traditional views on survival, resilience, and spread of cold-tolerant trees as a response to climate changes, and although restricted to specific regions and taxa, they are relevant to paleoecological studies worldwide. The ability of trees to survive at high latitudes has important implications for the response of forests to future climate changes and is particularly important for conservation and forest breeding programs interested in identifying genetic resources and provenances with traits related to survival under changing climates.

A first implication is the limited dispersal ability of certain forest tree species, which in turn implies that factors like geographical accessibility from glacial refugia and history are important constraints in species distribution (Svenning et al. 2008), and may act strongly as diversity predictors together with climate. It also implies that we cannot expect forests to simply track the forecasted 21st century climatic changes as suggested by several ecological niche modeling studies.

A second important implication is that postglacial migration rates based on pollen records require revision and should not be used alone to predict tree-range shifts in future modeling-based scenarios. Lower postglacial migration rates imply also that trees tolerate climatic changes better than previously assumed.

The scale of climate changes predicted for the next century is comparable to the scale of warming during the Holocene (several degrees Celsius depending on the region), but the rate of change is forecasted to be much faster (IPCC 2014). Recent modeling studies based on general circulation and vegetation models have suggested that rates of future climate changes require plant migration rates even faster than those observed in the fossil record, and hence may drastically reduce biodiversity by selecting for highly mobile and opportunistic plant species. However, the ability of certain species to tolerate changing climates and survive in northern refugia may mean that some catastrophic projections based on these models (Thuiller et al. 2005) might be overstated.

A final important implication for conservation is that northern populations may have preserved locally genotypes and genes best adapted to changing climatic conditions that are worth protecting. If trees only migrated from the south, the adaptation we see today to changes in temperature and photoperiod must have occurred over a relatively short time (the Holocene or the last ~13,000 years), and is based on “standing genetic variation” present in the southern macrorefugia, since this is a faster process than adaptation based on new mutations.

New directions of plant aDNA research

Despite early challenges, plant aDNA research now offers the opportunity to detect hidden plant diversity and a complementary spectrum of fossil remains than those identified by traditional paleoecological analyses alone. In addition, new sequencing technologies have recently significantly changed the direction of plant aDNA research and we have moved from the analysis of a few chloroplast loci to the study of entire nuclear and organelle genomes. This has already provided important information for plant evolution, adaptation, conservation, crop breeding, and food security (Estrada et al. 2018). However, the aDNA field cannot work alone. Collaborative studies between researchers with different backgrounds (geologists, paleoecologists, archeologists, and molecular ecologists), as well as acknowledgment and discussion of the results from complementary disciplines, is not only a recommendation, but a requirement in this rapidly growing research field.

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Modeling past plant species' distributions in mountainous areas: A way to improve our knowledge of future climate change impacts?

Louis François¹ and Alain Hambuckers²

Paleovegetation modeling, particularly in mountainous areas, could be an important method to understand and produce more reliable forecasts of the impacts of climate change on vegetation and plant species’ distribution. The two main tools used for such studies are dynamic vegetation models and species-distribution models.

Greenhouse gas emissions and climate change are, without doubt, affecting plant species’ distributions. The diversity of individual responses, the complexity of biotic interactions, and the intensity of human perturbations (fragmentation, pollution, hunting, etc.) make predictions for the future challenging. High-altitude mountain vegetation is particularly at risk and therefore interesting to study because, in the mountains, the climate gradients are steep and it is impossible for species growing near the summits to migrate to higher altitudes.

One of the keys to understand mountain species dynamics and assess future changes in their distributions, productivity and competitiveness, may be searched for in the past. Recent studies (e.g. Cheddadi et al. 2017; Pinaya et al. 2019) have shown that mountain tree species in Africa and South America have responded strongly to late Pleistocene and Holocene climate fluctuations. Here, we briefly examine whether current vegetation and plant-species models are appropriate tools to study those problems, and what the necessary requirements are to model mountain tree species’ turnover observed in connection with paleoclimatic changes.

The models and their data
The most popular tools are the species-distribution models (SDMs; Fig. 1). They establish empirical relationships between species’ presence and a set of predictors (climate variables, soil properties, land use, etc.; Thuiller et al. 2009). Vegetation can also be modeled with dynamic vegetation models (DVMs), which are process-based models describing the establishment, growth, and mortality of objects representing various plant types, competition among these objects, and processes such as photosynthesis, and respiration (Snell et al. 2014). Most often these objects are plant functional types corresponding to broad vegetation units, but it is also possible to simulate taxa, like species or genus. SDMs and DVMs require large amounts of climatic and environmental data with high enough resolution and accuracy. DVMs used with species require validation data sets like net primary productivity, which are more difficult to obtain.

In mountains, the accuracy of the forcing and validation data represents a challenge. Climate is strongly variable in space and time and it has been ascertained that growth and establishment of species are strongly dependent on local microclimatic and topographic features. The first challenge is thus to identify those conditions, by developing gridded climatic datasets at high spatial resolution over the studied areas. This is not an easy task, since not only temperature is affected by elevation and complex topography, but also precipitation and cloudiness, solar radiation, air humidity, and wind speed. All these variables must be reconstructed in a consistent way despite the scarcity of meteorological stations in high elevation.

Figure 1: The two main modeling approaches available to reconstruct paleovegetation distribution, with features needing very substantial (red-red), substantial (orange-red), intermediate (orange-orange), small (green-orange) or very small (green-green) efforts.
areas. Temperature inversions are also frequent, especially in winter, and the use of an average vertical temperature gradient may be problematic (Yang et al. 2011). Ideally, dynamic downscaling with high-resolution regional climate models should be used, but it is still rarely the case in ecological studies (Maclean 2019), especially where paleoclimatic reconstructions are concerned.

Complex topography also impacts soil hydrological conditions because of water transport along slopes. The water flow establishes hydrological gradients between mountain summits, and valleys, which induces changes in water-table depth. The presence of springs and the access to or capillary rise of groundwater may be critical for species survival during droughts; this has been found to be essential in explaining the distribution of vegetation at high spatial resolution (at scales smaller than a few kilometers) on any type of terrain. For that reason, SDMs and DVMs should take into account the water transport among grid cells and possibly the dynamics of the water table, when they target a very high spatial resolution. This approach, however, requires detailed and accurate information on soil depth, soil texture, and underlying rock porosity (Wolf 2011), generally not available over the entire distribution area of typical plant species.

**CO₂ and plant traits**

The atmospheric CO₂ level is another environmental variable that controls plant growth and distribution. The mixing ratio of CO₂ in the atmosphere (~410 ppmv in 2019) has shown very large fluctuations in the past: 1000 to 2000 ppmv in the Eocene, 190 ppmv during the Last Glacial Maximum, and 280 ppmv in the pre-industrial era. It has been established that elevated CO₂ stimulates plant growth and resistance to hydric stress at least in the short term. Only DVMs are able to integrate these effects. While DVMs seem more appropriate than SDMs to simulate past plant distribution, particularly in view of their sensitivity to CO₂, their use is not straightforward, because this requires information on the morphophysiological properties of the taxa, the so-called plant traits. Examples of important traits are leaf and wood nitrogen contents, specific leaf area (i.e. area-to-weight ratio), and leaf longevity. Today, trait data are implemented in international databases, such as TRY (Kattge et al. 2020). However, they are still lacking for many species and the collection of such data is time consuming. In addition, many traits are sensitive to climate and environmental conditions (Fig. 2), reflecting the acclimation capacity of the organisms.

Thus, there may have been variations in these traits in the past, modifying the species’ sensitivity to climate variables. The involved processes are complex and consequently large uncertainties remain. For instance, this is the case with the long-term impacts of high CO₂ levels (Van Der Sleen et al. 2015). CO₂ enters the leaf through the stomata. This inflow increases under elevated CO₂, which stimulates photosynthesis. But carbon fixation by photosynthesis also depends on leaf nitrogen, and it has been observed that, in the long term, the plant acclimates to elevated CO₂ by allocating less nitrogen to leaves, which reduces the productivity enhancement. Also, stomatal density declines under elevated CO₂, a phenomenon used to evaluate paleo-CO₂ from fossil leaves (Royer 2001), which finally increases plant resistance to water stress and impacts the distribution.

**The biotic interactions**

Beyond the problems of physiology, additional complications in predicting the past or the future are associated with biotic interactions (competition-facilitation, predation-parasitism) and migration limitations. The ranges of environmental conditions in which the species are able to thrive (fundamental niche) are modified by these biotic interactions. Species could also be absent in suitable conditions owing to dispersal limitations. SDMs include both, since they straightforwardly and statistically reflect the species-distribution sample used for their computation. This explains their ability to reproduce the present distribution, but also the challenge in predicting the past or the future. Indeed, SDMs rely on the hypothesis that biotic interactions and dispersal limitations are temporally constant. With species, DVMs reflect more the fundamental niche because the environmental thresholds limiting plant growth are derived from extreme supported values (Raghunathan et al. 2019). Obtaining the realized niche distributions requires long computation times with other plants to allow for competition for light and water to equilibrate.

Moreover, when making past or future projections, both modeling approaches produce new areas of climatic suitability but are rarely able to determine whether these areas could be reached by the organisms from their initial range. A first approach consists of using cellular automaton constraining cell occupancy of the new suitability areas with dispersal factors (Engler and Guisan 2009). The second approach is the coupling, with the plant models, of mechanistic models of seed dispersal by wind or animals. The main difficulty may be the discrepancy between the scales of the processes, since the dispersal model works for individual plants, while the resolution of plant models is typically on the order of at least several kilometers. Both methods could be used with DVMs or SDMs but data are lacking particularly for validation.

In conclusion, for achieving reliable reconstructions of paleovegetation, current models could be improved. However, it is mandatory to develop spatio-temporal series of data to allow for robust validation of these models.

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Past changes in species diversity: A view from the mountains

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Mountains host a spectacular diversity of species. Yet current species distribution is only a snapshot in time that bears the memory of past geomorphological and climatic changes. Understanding past changes is key to interpret current patterns and inform mountain biodiversity conservation and management in the future.

The present: mountains are (bio)diverse
Terrestrial mountains sensu Körner et al. (2011; 2017) cover approximately one-tenth of the total land surface. These mountains vary in many ways, from their geological history to their appearance and ecosystems. The Chaîne des Puys in France or Mount Fujiyama in Japan are volcanic, whereas the Rocky Mountains of North America are so-called fold mountains that resulted from the thickening of the Earth’s crust. In the Black Hills of South Dakota, USA, mountains are dome-shaped, whereas the table-top mountains of Venezuela are plateaus.

Within the same African continent, some mountains like Mount Kilimanjaro are high and capped with ice, while others such as the Eastern Arc Mountains are lower and mostly covered with dense tropical forest. Some, like the Kohala Mountains of Hawaii, USA, cover a small geographic area, while others like the Himalayas are immense and stretch across several countries and thousands of kilometers. And yet as diverse as they are (Gordon 2018), all these mountains share a common characteristic: they host a spectacular diversity of species, of which many are endemic, and contribute disproportionately to the terrestrial biodiversity on Earth (Rahbek et al. 2019a). The Cape Fold belt of South Africa alone has 68% endemism across the 9000 plant species known to science (IPBES 2018), the Andes are home to an approximate 45,000 plant species (Myers et al. 2000), and more than half of the world’s “biodiversity hotspots” are entirely or partly in mountainous areas (Körner and Ohsawa 2005).

The past: geomorphology, climate, and evolution
The current distribution of mountain species and their high levels of richness and endemicity just represent a snapshot in time that bears the collective memory of the millions of years of geomorphological and climatic changes that drove species along individual evolutionary trajectories. These environmental drivers often left almost intangible traces, making the reconstruction of the historical biogeography of mountains and their biodiversity an intriguing puzzle. Yet, the general consensus is that uplift events (Hoorn et al. 2018) set the stage for species’ evolution by providing novel ecological opportunities along newly arising surfaces (Antonelli et al. 2018a).

The uplift of the tropical Andes, for example, was instrumental in the development of biodiversity and humidity entrapment in the Neotropics, including the Amazon basin. It created both dispersal barriers and opportunities for colonization and diversification through novel high-elevation environments and greater topographic complexity, as well as for dispersal along newly opened routes. Uplift and other geomorphological processes created diverse mountain morphologies (Elsen and Tingley 2015), as well as complex and unique landscapes of ridges, valleys, peaks, and plateaus (Molnar 2018). With recent methodological advances in thermochronology and paleoaltimetry (e.g. Antonelli et al. 2018a; Perrigo et al. 2019), estimations of the timing and rate of these processes are becoming increasingly precise, robust links can be drawn between spatial biogeographic patterns, and geological scenarios can be tested. Such techniques revealed, for example, that some mountain systems such as the Southern Alps in New Zealand uplifted in the past 15 million years (Myr) whereas others such as the European Alps appeared longer ago.

The interaction between novel topographies, tectonic and erosional processes, regional climates partly dictated by the geographic location of mountains and their orientation in relation to global atmospheric circulation patterns (Antonelli et al. 2018a), and shifts in climate patterns subsequently served to increase environmental heterogeneity and achieve the extraordinary diversity of species and life histories in mountains.

Over the last three or more Myr, successive climatic cycles modified patterns of biodiversity by stimulating geographic-range shifts within mountain regions and between mountains and adjacent lowlands, driving rapid altitudinal migrations upslope during interglacial and downslope during glacial periods, temporarily opening pathways or creating barriers for dispersal to new environments, or dividing and isolating populations.

Accordingly, species richness can be envisioned as the result of multiple episodes of diversification triggered by the consecutive connection and isolation of mountains and species, a hypothesis described in the so-called “flickering connectivity system” framework (Flantua and Hooghiemstra 2018; Flantua et al. 2019). This hypothesis builds on the premise that changes in historical connectivity are important – and possibly as important or even more so than mere isolation – in determining present biodiversity, with historically connected areas characterized by higher species richness (Flantua and Hooghiemstra 2018).

An important fragmentation mechanism associated with climatic cyclicity that played an important role in the diversification of species, such as birds in the Southern Alps of New Zealand (see Flantua and Hooghiemstra 2018) or plants in the Tibeto-Himalayan region (Mosbrugger et al. 2018; see Mueller-Riehl 2019), comes from ice caps on mountaintops and ridges. Yet, while fragmenting habitats, ice also created islands of ice-free habitats, microrefugia from where species such as Saxifraga florulenta in the Maritime Alps could recolonize downhill habitats after episodes of warming (see Patsiou et al. 2014).

Accordingly, the glacial and interglacial periods resulting from climatic fluctuations also played an important role in triggering the evolutionary mechanisms driving present-day species diversity (e.g. Theodoridis et al. 2017), including fragmentation, hybridization, speciation, and extinction (Rahbek et al. 2019b), which all varied in time and space among species and mountain ranges, leaving unique “mountain fingerprints” (Flantua and Hooghiemstra 2018).

The future: why and how to learn from the past
In view of expected changes in present climatic conditions, a better understanding of past species’ community dynamics (see Koutsoudendris et al. 2019) and of the relationship between environmental transitions and temporal dynamics in morphological and functional traits (Papadopoulou and Knowles 2016; Huang et al. 2019; Mueller-Riehl 2019) can extend the lead-time for adopting sustainable management and adaptation measures. This is particularly important in high elevation and arctic regions.
exposed to rapid warming (Pepin et al. 2015) and likely to experience particularly severe biodiversity losses (Dullinger et al. 2012). Episodes in Earth’s history characterized by temperatures and atmospheric carbon dioxide concentrations comparable to those expected in the near future are therefore particularly informative.

Episodes that can serve as an analogue for a future anthropogenic world are the Pliocene (Burke et al. 2018) and the early Eocene (Hollis et al. 2019), when global temperatures were some degrees warmer than today, atmospheric carbon dioxide concentration was slightly higher, and major shifts in plant communities occurred. Examples of such shifts include the strong expansion of Artemisia at the expense of Chenopodiaceae and other drought-tolerant taxa in the Chinese Loess Plateau (Koutsodendris et al. 2019).

Unraveling the biogeographic history of mountain species, determining the relative role and importance of geomorphological processes and climate on the evolutionary trajectories of species, and learning from the past to inform the future requires interdisciplinary approaches and novel methodologies. Interdisciplinary approaches are needed that integrate expertise across fields and temporal scales from paleobiology and -geology to contemporary conservation, genomics, and remote sensing (see Antonelli et al. 2018b on transdisciplinary biogeography).

Novel methods, in turn, are needed to accurately measure the timing of mountain formation and disentangle the effects of mountain uplift from environmental and climatic changes (Perrigo et al. 2019). Together, approaches from thermochronology or paleoaltimetry have considerably improved models and temporal estimates of mountain uplift (Perrigo et al. 2019). Fitting increasingly robust models with data ranging from fossil records to genome sequences is an effective approach to address the correlative nature of prevalent analyses and the space-for-time substitution (Perrigo et al. 2019). Paleo-niche reconstructions and modeling combined with phylogenetics and phylogeography are in turn powerful approaches to reconstruct mountain biome dynamics and improve our understanding of the role of these dynamics in shaping current biodiversity patterns (Flantua and Hooghiemstra 2018).

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Figure 1 Linking the evolution of mountains and biodiversity. (A & D) Climatic (blue bar) and geological (green bar) variables influence species richness through speciation, immigration, and extinction (lower bars) through different orogenic phases. Darker/warmer and lighter/colder colors in the bars indicate high and low levels, respectively. As the arrows indicate, climatic and geological processes exert the strongest effects on species richness, but some feedbacks exist (such as plants colonizing bare rocks and increasing erosion, and animals affecting soils through burrowing). Figure courtesy of I. Voet, Muséum National d'Histoire Naturelle, Sorbonne Université, Paris; first published in Antonelli et al. (2018b).
Using paleoecology to inform restoration and conservation of endangered heathlands

Cherié J. Dirk and Lindsey Gillson

When restoring damaged or degraded ecosystems, land-use managers usually have a restoration target in mind. But how do they decide this target, and how do they know if it is realistic? This question is especially important when dealing with Renosterveld, a highly endangered component of the Fynbos biome in South Africa.

Restoring a biodiversity-rich, but endangered, ecosystem

Renosterveld is an evergreen, fire-prone Mediterranean-type shrubland, which is restricted to fertile, fine-grained soils of the Fynbos biome. The composition of Renosterveld generally includes grasses and a high diversity of endemic geophytes (bulbs), for example, of the genera Amaryllidaceae, Asparagaceae, Iridaceae, Hyacinthaceae, Orchidaceae, Oxalidaceae, and Geraniaceae, many of which are endemic, rare, or threatened. For example, of the Iridaceae, about 48% of the known species are listed as rare or threatened and about 76% are endemic to Renosterveld. Some bulb species (such as Freesia, Ixia, Ornithogalum, and Pelargonium) are economically important world cut-flowers (Rebelo et al. 2006). Unfortunately, 91-97% of Renosterveld has been transformed for agriculture, so the remaining fragments are vital to the survival of this megadiverse biome. The species that currently dominates Renosterveld fragments is an evergreen, unpalatable shrub called Elytropappus rhinocerotis (Renosterbos). Little is known about the historic abundance and distribution of Renosterbos, leaving managers and conservationists to speculate on whether the current composition is typical and therefore a worthy conservation target, or whether restoration efforts are needed.

A long-term understanding of Renosterveld

Very little is known about the history of Renosterveld, and whether what remains today is typical of its once much more extensive past. Though fire and herbivores are integral to Renosterveld ecology, it is difficult for land-use managers to know whether their prescribed grazing and burning regimes are appropriate to maintain such an extraordinary and unique species diversity. Furthermore, there are conflicting ideas of what Renosterveld vegetation should consist of – less or more grasses and shrubs (Cowling et al. 1986; Curtis 2013; Curtis and Bond 2013).

One way to resolve this dilemma is to look at long-term data to find out how Renosterveld looked and functioned ecologically prior to intensive management and land transformation, which began in the mid-17th century with the arrival of European settlers and further increased in the 20th century due to agricultural intensification. As there is little documented information from this period, long-term paleoecological data can be used to reconstruct how ecosystems looked in the past. Fossil pollen, coprophilous fungal spores, and charcoal from sediment cores can be used to reconstruct changes in the surrounding vegetation, herbivory (grazing) and fire regime. This multiproxy approach can be used to assess the relative importance of changing land use (specifically fire and herbivory) and climate in driving vegetation change.

Elandsberg Private Nature Reserve as a case study

Elandsberg Private Nature Reserve (PNR) is located in the Wellington District of the Western Cape, South Africa (Fig. 1), and contains one of the few and largest remaining fragments of West Coast Renosterveld. It is especially important because it is home to the critically endangered geometric tortoise (Psammobates geometricus), which is endemic to this region and one of the rarest tortoises in the world (Boycott and Bourquin 1988). The area has a long history of land use, with people managing fire and herbivory for more than 2000 years. Hunter-gatherers were in the region from 10,000-2000 yr BP (Deacon 1992) and land use further intensified with the arrival of the Khoikhoi pastoralists, who introduced livestock approximately 2000 yr BP (Cooke 1965; Elphick 1977; Boonzaier et al. 1996). With the arrival of European-settler agriculture from the mid-17th century, the region saw an increase in grazing pressure and grain crop cultivation, and combinations of fire suppression (to protect agriculture and property) in some areas and regular burning to promote pasture in others. In the 20th century, technology and industrialization allowed agricultural intensification and further transformation of natural vegetation.

A paleoecological study has now shed light on the history of the vegetation of Elandsberg PNR, providing important pointers for management and conservation in the future (Forbes et al. 2018). Pollen, charcoal, and fungal-spore data showed that the landscape of Elandsberg PNR today is still heavily influenced by human use, despite the area being a nature reserve since 1973. The data showed that Elandsberg PNR has been heavily transformed by farming since about the 1950s, when domestic livestock were abundant.
Farmers started burning renosterveld much dramatically at this time, suggesting that charcoal abundance increased the 1950s, when herbivores were fewer, fire frequencies could help managers to find ways of reducing renosterveld abundance and restoring vegetation to a more healthy state. Maintaining a diverse array of vegetation types in the reserve is good for renosterveld biodiversity and also might help build resilience to future climate change. Therefore a flexible management approach and long-term monitoring at the patch and landscape level is recommended. Furthermore, future research using system dynamics modeling will provide decision-making support, allowing managers to explore the effects of future scenarios of changes in fire and grazing (Dirk et al. in prep).

Paleoecology provides a long-term perspective needed for restoration
Projects such as these show how short-term perceptions can mislead managers, to the detriment of biodiversity conservation. Most landscapes have been heavily transformed by people over the past century, and many have also been impacted by other stressors, such as climate change, and rising CO₂ and Nitrogen deposition. Using 21st century baselines for conservation and restoration targets could therefore be inappropriate. Restoration is an important component of conservation biology as it seeks to actively reverse degradation of an ecosystem, but accurate information is needed to inform managers about the historical range of variability of biodiverse landscapes, and the effects of human management in recent decades. Paleoecological research can provide context for interpreting recent changes and in developing restoration targets that conserve biodiversity and ecosystem services for future generations. Paleoecology can inform restoration targets by providing a window into how ecosystems functioned before heavy impact by people in the 20th century, and therefore guide future restoration. In the case of renosterveld, managers can use paleoecological data to decide whether landscapes contain less or more, for example, grass, bulbs, grazing, or fire, to be at a healthy state that is resilient and sustainable in the future.

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Mountain science poised to help ecotourism in Peruvian cloud forests

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We seek to highlight how paleoecology, archaeology, and geoecology can add to the repertoires of ecotourism guides in Peru’s Chachapoya region, providing informed portraits of the history of cloud forest ecology in Peru’s northeastern Andes and raising concerns about the future conservation of these mountainscapes under human impact.

The eastern slopes of the Andes are underexplored and underexploited from the perspective of ecotourism. In the northern region of Chachapoyas (Fig. 1), where we have focused recent research, phenomenal biodiversity and pre-Hispanic cultural diversity are present in such proximity to each other that an ecotourist spending seven to 10 days on a single visit might encounter everything from mummies to marsupials. Biological and cultural diversity in these tropical montane cloud forests (TMCFs) have been, and continue to be, shaped by climate change; but scientists have largely missed the opportunity to engage guides and tourists, and to enrich their narratives and experiences while informing them on matters of montane conservation concern.

New research initiatives are improving our understanding of linkages between climate oscillations and how past human activities influenced modern trajectories of succession and community composition in Chachapoyas. Mountain geographers are now collaborating with paleoecologists and archaeologists to explore the origins of Anthropocene landscapes in the tropical Andes (Bush et al. 2015; Guengerich and Berquist 2020).

Cloud forests as a transformed landscape

Until recently, the Andean-Amazon slopes were considered remote and inaccessible, a prordial wilderness. We now know that humans arriving by the end of the Pleistocene probably contributed to the extinction of megafauna (Raczka et al. 2018), and initiated megadroughts (yellow bars from the Laguna de los Cóndores pollen record): 12,000 years of landscape domestication (Schjellerup et al. 2005). Between ca. 1200 CE and 1400 CE, a cultural tradition with shared attributes, including some elements of architectural and ceramic style, reached its peak across a settled area of ca. 30,000 km² (Schjellerup 2019). Termed “Chachapoyas culture” by archaeologists, the Inka first referred to the region as the “land of the people of the clouds” or sachapuyu, and this Kichwa name appears on the earliest colonial documents. A combination of demographic collapse, social disintegration, and lack of interest in the native cultures shown by Spanish colonists rapidly eliminated cultural knowledge, leaving the task of reconstruction to historical geographers, landscape archaeologists, historical ecologists and paleoecologists.

This is where new data on climate change are revealing unexpected patterns. Intense droughts immediately conjure images of hardship – possibly even landscape abandonment. But in these very wet settings, those droughts appear to have fueled agricultural incursions into cloud forest (Åkesson et al. 2020). Portions of the landscape were cleared for maize cultivation until wet conditions returned. Thus, the more accessible the cloud forest, the more likely it was to have supported past human activities. In large part, the factors that make a modern valley or ridge accessible to ecotourists also made it a usable space for past cultivators. Consequently, the areas that tourists can readily access today are very likely to have long histories of intensive land use, dating to pre-Columbian times (Sarmiento 2012) constituting de facto biocultural microrefugia.

Cross-cutting science for montological studies

Montology is the transdisciplinary science of mountains, where current geocological knowledge, paleoecological insights, and archaeological studies in the region’s TMCFs now complement one another. Pollen data obtained from lakes tell stories of climate extremes beyond our modern experience, both wet and dry, against a backdrop of long-term change. Pollen records collected from Lake Pomacochas (Bush et al. 2015) and Laguna de los Cóndores indicate millennia of maize cultivation, demographic and subsistence shifts, and ecological turnover before the Spanish invasion of 1532-35 CE (Matthews-Bird et al. 2017). Within the last 1000 years, as regional climates became wetter, some locations show forest recovery apparently at the expense of cultivated lands, perhaps as land-use strategies have shifted toward exploiting drier slopes, the net effect of which was widespread forest recovery. Within the last 500 years, the Inka incorporation followed by the Spanish invasion and subsequent colonization, including the effects of epidemic diseases, decimated local cultures and concentrated inhabitants in readily controlled centers of occupation, further emptying the forests of inhabitants. During the 20th century, accelerating population growth and enhanced accessibility have ignited new efforts to conquer the “wilderness” and resume landscape domestication (Schjellerup et al. 2005).

Another realization is that many TMCF Andean habitats that were long thought to be “natural” are now seen as having long histories of human modification or “engineered” landscapes, and that a reconsideration of their resilience to climate change is needed (Sarmiento 2016). Because fire was part of past land management, it would have influenced survival rates and increased edge effects on unburned areas. The hunting
pressure from cultivation-based communities would have also reduced the population of game animals and disproportionately removed large seed-dispersers (Aubad et al. 2010). Favorled plants might have been tended to, planted, or even accidentally introduced, even at some distance from villages. Consequently, when the forests regrew, they would have been on a different successional trajectory than in an undisturbed setting (Levis et al. 2012). While species compositions changed, so too did the physical landscape, as hill slopes were modified by either earthen terracing or stream canalization.

Modern trade has accelerated landscape transformation and introduced many exotic species to the region, including African grasses, Australian Eucalyptus, and Monterey pine. Consequently, these forests, which exist as a full gradient from "novel" communities to nearly natural ones, will respond differently to climate stimuli than their natural counterparts. Stimuli also come from the socio-ecological systems in need of applied montology (Schjellerup et al. 2019) and better conservation strategies, such as those emphasizing ecotourism ventures, such as "the route of petroglyphs", "the route of mum-mies", and "waterfall circuits" featuring one of Peru’s tallest waterfalls (Goceta), a spectacular hilltop citadel (Kuelap), and impressive mortuary sites such as the sarcophagi of Karajía and the mausoleums at Revash. Together, they create a vibrant touristic destination of “natural” and “cultural” sites that may one day may rival the Sacred Valley near Cuzco, if properly encouraged.

We argue that the only way to maintain the unique characteristics that draw ecotourists to an area is to curate the biocultural heritage and to seek to apply regenerative development strategies that confer additional resilience to landscapes in the face of the expansion of cattle grazing. We could start with restoring degraded slopes and reforesting large areas that could bring back the former TMCF, including iconic animals that attract ecotourists, such as Peru’s national bird, the cock-of-the-rock (Rupicola peruviana), the spectacular bear (Tremarctos ornatus), the spatula-tailed hummingbird (Loddigesia mirabilis), the mountain lion (Puma concolor), the critically endangered yellow-tailed woolly monkey (Oreonax flavicauda) and the long-whiskered owlet (Xenoglaux loweryi). Such protections require not only trained guides but also changes in management strategies by owners, as well as an about-face in popular consciousness to curve destruction of the “Ceja de Selva”.

Effective management can be informed by natural fire periodicities of >1000 years elicited from paleoecological records and lists of taxa that should form the basis for replanting in both natural and civic settings. It is imperative that the modernization efforts encourage tourists to spend time in the city of Chachapoyas, where old native trees of the plazas have been replaced by newly planted exotic species. But perhaps one of the most valuable roles that mountain science can play is by informing the narratives and anecdotes that provide local insights and environmental appeal to the descriptions given by science-trained guides. They must provide information from montology that is accurate, enriching, and deepens the understanding of how our history shapes what we perceive to be natural in the biocultural microrefugia of the tropical Andes.

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**PAGES**

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Columbus’ footprint: Land-use change before and after European incursion in Hispaniola

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Shortly after Columbus’ landing in the New World in 1492 CE, indigenous land use diminished and cattle grazing was introduced. Large-scale agriculture in the Cibao Valley, northern Dominican Republic, did not develop before the 17th century. Here, we read the environmental history of the Cibao Valley from two sediment cores.

In 1492 CE Columbus landed in the “New World”. In 1494 CE he founded La Isabella, the first colonial town in the Americas, located at the northern coast of the current Dominican Republic (Hayti island; Fig. 1; Ortega 1988). Five hundred years of colonial exploitation followed, which had far-reaching impacts for the indigenous people and local landscapes (Hofman et al. 2014; Siegel et al. 2015). Paleoeccological analyses of sediment cores collected in sediment-filled meanders of the Yaque del Norte River (Cibao Valley) show the regional environmental history. Pre-Columbian indigenous land use had an impact on the forests of the valley. Depopulation due to post-contact epidemics and warfare was a major driver of dramatic landscape change.

Sources of information
Changes in land use were studied from various threads of paleoecological evidence. Organic matter was radiocarbon-dated and provided a time scale for the sediment column. The records of fossil arboreal and herbaceous pollen and phytoliths (silica structures occurring in plant cells characteristic of specific groups of plants) show terrestrial vegetation changes. Non-pollen palynomorphs (e.g. spores of coprophilous fungi) are indicative of land-use changes, as they provide evidence of the presence of big mammals, and these changes in abundance are indicative of the introduction of European cattle. The record of charcoal documents the fire history (increased fire occurrence indicates deforestation to develop agriculture) and the plant macrofossils show the composition of local wetland vegetation. Grain size analysis shows the changing energy levels of the river (coarse-grained silt and sand reflects intense flooding and fine-grained clay standing water) and the proportion of organic matter in the sediments reflects the abundance of aquatic vegetation. Fossil pollen of food plants and phytoliths show which food plants have been grown through time. All these threads of evidence come together to reveal a detailed account of how people have interacted with the landscape during the last 1800 years (Fig. 2).

The sediment core from site Biajaca reflects the last 1100 years (see Castilla-Beltrán et al. 2018). Biajaca is centrally located in the Cibao Valley ca. one kilometer away from the Columbus Route (Ruta de Colón), connecting the northern coast with the mountainous interior. This first informal “road” in the New World allowed for the transportation of gold and other products. At a kilometer distance from Biajaca, there is an archaeological site of an indigenous settlement. The sediment core from site Los Indios, some 40 km to the west, shows the last 1800 years (see Hooghiemstra et al. 2018). Both records together show a fascinating view of human activities before and after European incursion in the Americas.

Pre-Columbian landscape changes
The translocation of continental domestic plants, such as maize, chili pepper, and sweet potato was a process that started from the first peopling of the Caribbean, around 7800 years ago (Pagán-Jiménez et al. 2015). In two later main pulses of migration, around 3500 and 500 BCE, the islands of the archipelago were populated by indigenous communities (Hofman et al. 2018). Until ca. 1400 CE, human impact on the environment of the studied area consisted of slash-and-burn practices to open areas for the cultivation of crops such as maize (Zea mays) and squash. This is evidenced by sustained charcoal deposition and the presence of pollen and phytoliths of these cultivars in the Biajaca record. Settlements were probably distributed in a mosaic structure, leaving relatively little disturbed forest on the floor of the Yaque River.

The impacts of European colonization
The period from 1450 CE to ca. 1650 CE shows increasing human impact on the landscape. Columbus encountered a relatively densely populated valley with agricultural production on mounds and river terraces. Introduced European diseases decimated the indigenous population (Koch et al. 2019). This is evidenced by the absence of charcoal deposition at the Biajaca record and in the forest regrowth in both records. First colonists introduced cattle from the Iberian Peninsula; this is shown by more abundant spores from coprophilous fungi in a more open landscape. The island was becoming a hub for further exploration and occupation of the Caribbean and Central America, and chronicles report an intense conflict between Spanish colonizers and indigenous people (Hofman et al. 2018). At the same time, enslaved people from Africa were brought to the island. Ruderal plants (Artemisia, Plantago, and representatives of the Boraginaceae, Polygalaceae, and Brassicaceae families) are indicative of ecological succession in abandoned agricultural fields and cleared forest plots. Introduced taxa from the Old World, such as Rubus, also appear in the pollen record.

Near the middle of the 16th and 17th centuries, the charcoal record shows an increase in fire, reflecting forest clearance in the surroundings of the Los Indios site (western Cibao Valley) and near the site Biajaca (along the Columbus Road)
respectively. Aquatic vegetation started to develop, indicating that the meanders were isolated from the drainage system. More abundant organic matter accumulated in the oxbow lakes. The establishment of Typha, Sparganium, and Cyperaceae plants reflects local terrestrialization at the sampling site. A plethora of changes, such as increasing fire, deforestation, loss of arboreal taxa, and the appearance of cultivated plants (banana, and solanaceous plants, potentially including chili pepper and tobacco, which we currently cannot identify to the species level) reflect the introduction of monocultures on a large scale in the Cibao Valley. After independence in 1844 CE, forest remnants were burned and charcoal deposition gradually decreased. Today, plantations of banana and rice dominate in the Cibao Valley. Some 150 years later, abundant use of fire deforested the Cibao Valley, and large-scale agriculture was developed to serve European and local markets. From the moment of independence of the Dominican Republic, this valley became an agricultural center and was largely cleared of forests. Proxy-specific reconstructions can uncover many details, and a multi-proxy synthesis shows how the landscapes of the island developed into the ones we see today. Understanding these historical developments is essential to better appreciate successive steps in human-driven environmental degradation in this Caribbean island from pre-Columbian times.

Significance

"Reading" the environmental information deposited in two sediment-filled meanders near the Columbus Route yielded new information about how indigenous people used the landscape in pre-Columbian times. Indigenous land use changed dramatically after 1492 CE. Colonizers immediately introduced cattle farming, and depopulation caused forest regrowth.

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Figure 2: Paleoenvironmental change, Cibao Valley (Dominican Republic): multiple lines of evidence from the Biajaca and Los Indios sediment cores. Measured values have been normalized (varying between 1 and 0). The records show the last ca. 1800 years of landscape and land-use changes in the Cibao Valley.
Relaunching the African Pollen Database: A abrupt change in climate and ecosystems

Sarah Ivory1, A.-M. Lézine2, E. Grimm3 and J.W. Williams4

Paris, France, 13-16 October 2019

African ecosystems hold enormous ecological and economic value due to high biodiversity (Myers et al. 2000) and valuable ecosystem services provided to urban and agrarian populations (Wangai et al. 2016). However, these services are vulnerable to land use and climate change (Niang et al. 2014). Long paleoecological records from Africa provide iconic examples of abrupt environmental change, offering critical evidence for tipping points in the Earth system. Datasets in the region are notoriously difficult to access with the African Pollen Database (APD) largely unsupported for the last decade. Poor data accessibility has been a community complaint.

Hosted by the Institut de Recherche pour le Développement in Paris, France, 32 participants from 12 countries met to revive efforts by the African paleoecology community and relaunch the APD as a community curated data resource (Williams et al. 2018). This workshop was organized by the LOCEAN and funded by the Belmont Forum for Science-driven e-Infrastructure Innovation project “Abrupt Change in Climate and Ecosystems: Where are the Tipping Points?” The Belmont e-infrastructure initiative seeks to enhance open, FAIR, scientific cyberinfrastructure (Wilkinson et al. 2016) to meet pressing, policy-relevant challenges.

African Pollen Database: History and plan

The APD was first developed in 1996. The initial workshop and subsequent work, funded by the French CNRS, the EU (INCO, ENRICH) and UNESCO (PICG), established methods of collating pollen data, developed a standardized pollen nomenclature (Vincens et al. 2007), generated updated age models, collated images of pollen grains, and created a searchable web interface. Currently, the APD contains 288 fossil sites and 1985 modern samples. Due to a lack of funding and the closure of the French data center, Medias-France, the APD lapsed in 2007, making its extant data holdings largely inaccessible and leaving no home for newer paleoecological records.

Workshop participants identified three critical elements of the APD: (1) Pollen samples with associated chronological controls and other metadata; (2) Standardized taxonomy for pollen morphotypes; and (3) Pollen grain images. Participants developed a two-pronged plan for rebuilding these APD components. The APD webpage will be relaunched, in collaboration with LSCE (IPSL, France) to meet region-specific needs, including modern and paleo data, taxonomy development, and hosting of pollen images. Further, APD data will become a constituent database of the Neotoma Paleoecology Database (Goring et al. 2015), an international community-curated data resource that meets FAIR standards and is registered with ICSU-WDS and COPDESS. Planning began for a data steward training workshop in Amsterdam in January 2020 to begin uploading pollen data to APD/Neotoma. The participants saw value in these combined efforts for assuring the longevity of a living, community-curated APD data resource.

Building a community

Africa was divided into six sub-regions (Central, Southern, East, West, Arabia, Madagascar/Indian Ocean Islands, and marine), and regional experts identified records produced post-2007 and legacy datasets not included in the APD. The increase in research in Madagascar and Central Africa was notable, given prior underrepresentation of these regions. Specialists reported 365 known sites not in the APD, ranging in age from 1000 years to the Plio-Pleistocene. Modern samples (1985 entries) are also available, with more to be added. Some modern samples were collected as early as 1954 and should be used with caution to represent modern vegetation. In sum, these sites include a wealth of new information: marine and terrestrial, from diverse archives, many of which are well-dated with high temporal resolution. This will facilitate the burgeoning number of new studies and spatio-temporal meta-analysis of ecosystem-climate linkages, closing the gap with other regions.

Further, a governance body was launched, co-led by Sarah Ivory and Anne-Marie Lézine, with multiple scientists volunteering to serve as representatives and data stewards from each region. As the APD council is launched, an effort is being made to balance the numbers of African and non-African scientists as well as early-career scientists. Through representation within the APD and improved access to open data resources and software, we hope to build opportunities for young researchers from Africa.

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Figure 1: Paleocological sites within the APO (green), datasets obtained but not yet within the APD (yellow), and known datasets not currently contributed to the APD (red dots). Northern Africa is in the European Pollen Database.
Exploiting the SISALv2 database for evaluating climate processes

Yassine Ait Brahim\textsuperscript{1}, J. Hu\textsuperscript{2}, J. Baker\textsuperscript{1}, C. Perez-Mejias\textsuperscript{1}, H. Zhang\textsuperscript{1} and L. Comas-Bru\textsuperscript{3}

4th SISAL workshop, Xi’an, China, 14-18 October 2019

The SISAL working group (Speleothem Isotope Synthesis and Analysis; pastglobalchanges.org/sisal) dedicated its initial phase to creating and maintaining a database of worldwide speleothem δ\textsuperscript{18}O and δ\textsuperscript{13}C records. The first version of the database (Atsawawaranunt et al. 2018a) contained 381 speleothem records and was published in 2018 (Atsawawaranunt et al. 2018b). An expansion of the database (SISALv1b; Atsawawaranunt et al. 2019) increased that number to 456 records from 211 sites (Comas-Bru et al. 2019). Thanks to the efforts of SISAL’s committed members, the second version of the database (SISALv2), containing 691 records, will be made available in 2020.

Twenty-seven SISAL members (including 22 early-career researchers and 13 female researchers) from 12 countries gathered at Jiaotong University in Xi’an, China, for the fourth SISAL workshop (pastglobalchanges.org/calendar/2019/127-pages/1942). The main goals of this meeting were to finalize ongoing tasks related to the construction of SISALv2, attempt to resolve research questions by improving data-model comparisons through SISALv2, host a training session on database utilization and analysis, and finally to discuss and organize prospective projects under SISAL.

The workshop kicked off with a visit to the archaeological site of the Terracotta Warriors and Horses and the Lantian loess field, 35 km east of Xi’an. During the evening, invited speakers gave presentations about multiscale climate variability in Iberia (Dr. Ana Moreno), multiscale climate variability in South America (Prof. Francisco W. Cruz) and advancements on the application of fluid inclusions in speleothems (Dr. Hubert Vonhof).

On the morning of day two, the lead coordinator of SISAL (Dr. Laia Comas-Bru) provided an overview on the progress of SISAL and its ongoing activities. Afterwards, selected participants presented their work on regional climate patterns inferred from speleothems, as published in our Quaternary special issue on “Speleothem Records and Climate” (Harrison and Comas-Bru 2019). During the afternoon session, presentations focused on isotope-enabled climate models and data-model comparisons. The day closed with local researchers presenting their own speleothem-based research.

On days three and four, the workshop was structured as a mixture of training and analysis sessions. The analysis stream took advantage of the balanced representation of speleothem researchers and climate modelers present at the workshop to design new hypotheses, which could be addressed using the new version of the SISAL database and currently available isotope-enabled climate models. Participants divided into three breakout groups focused on individual time periods and/or climate events to identify analyses that could be of interest to the Palaeoclimate Modelling Intercomparison Project (PMIP) community. Additionally, some attendees worked on a handbook for reporting speleothem isotope records, drawing from experience gathered by the SISAL initiative. The training stream was open to anyone interested, with the aim to teach researchers how to use the SISAL database. The examples included plotting multiple speleothem timeseries with age error bars, plotting a map of pre-filtered sites, applying low-pass filtering, and conducting correlation and spectral analysis. The training stream is among the new initiatives taken by SISAL, and the group intends to run similar sessions at other conferences such as K9 (uibk.ac.at/congress/kr9/scientific-programme/workshops.html).

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Figure 1: The workflow of the training sessions. Three main tasks are data entry and quality checks of SISAL workbooks; querying data from SISAL database; and plotting/conducting analyses using the database.
Is the El Niño-Southern Oscillation changing? Lessons from the past

Mary Elliot¹, S.Y. Cahyarini², N. Abram¹,4, T. Felis⁵ and H. McGregor⁶

Belitung, Indonesia, 25-30 August 2019

A group of 50 scientists met in Belitung, Indonesia, in August 2019 to discuss past changes in the El Niño-Southern Oscillation (ENSO). ENSO is the main source of inter-annual climate variability that affects seasonal temperature and precipitation patterns across the Pacific Ocean with global teleconnections and severe societal and economic impacts. Research presented during this PAGES-supported meeting (pastglobalchanges.org/calendar/2019/127-pages/1805) aimed to provide critical information required to better understand short- and long-term changes in ENSO. Recent findings were reviewed and our present understanding of the factors controlling past ENSO variance were assessed. Areas lacking information were identified, and a common and integrated strategy for future interdisciplinary studies was proposed.

Paleoclimate data, recently reviewed by Emile-Geay et al. (2020), provide unique information on the full range of natural variability of ENSO and enable assessments of rainfall and temperature anomalies and their impacts on marine and terrestrial ecosystems. Anomalies of sea surface temperature and salinity associated with past ENSO variability are primarily derived from seasonally resolved marine fossils such as corals and bivalves. New techniques, such as single foraminifera analyses within discreet layers of marine sediment (e.g. Rustic et al. 2015), provide important avenues to expand the temporal and spatial coverage of paleo-ENSO information. Terrestrial archives such as speleothems, tree rings, and lake sediments record past changes in precipitation. Combining these approaches requires inter-calibration studies, which could expand the geographical coverage of ENSO reconstructions. These data, which are now integrated in a global database of past-interannual climate variability in the tropical oceans, facilitate intercomparisons of paleoclimate data and reconstructions with forced and unforced climate simulations. The combination of paleoclimate reconstructions and model simulations thus provides a framework to assess the mechanisms and forcing factors associated with past changes in tropical climate variability.

The achievements of this workshop were the identification of a list of key questions that urgently need to be addressed (see Box 1 and Fig. 1). Most recent studies still differ on the factors that control trends in ENSO during the Holocene: stochastic processes, solar insolation, response to the mean climate conditions, or a combination of these factors (Cobb et al. 2013; McGregor et al. 2013). Temporal trends in ENSO variance differ and are site specific, highlighting the importance of mapping geographical patterns as well as producing single-site records. Similarly, projected future changes in ENSO are model dependent (Collins et al. 2010). Advances in analytical precision and novel geochemical tools that have recently been developed provide a new impetus to address these research questions. We propose a road map to make progress on these research questions.

- Improve our understanding of the interactions between ENSO and other modes of variability, including the Indian Ocean Dipole and the East Asian Monsoon.
- Increase the temporal coverage to fully explore the response of ENSO to different mean background states (Last Glacial Maximum, Felis et al. 2012; previous interglacials; Pliocene).
- Reconstruct changes in the spatial patterns of ENSO (Fig. 1), which requires a denser coverage of proxy dataset (Freund et al. 2019).
- Build a global multi-archive database of interannual climate variability using a common metric. For this we need to homogenize methods for quantifying ENSO variance.

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Box 1: Key questions discussed at the workshop.

(1) How has ENSO changed over the geologic past?
(2) Most likely causes of the observed changes?
(3) Is ENSO changing now in response to anthropogenic forcing of climate?

Figure 1: December sea surface temperature anomalies associated with 1982, 1997 and 2015 El Niño events. This figure shows the evolution of the spatial distribution of temperature anomalies across the Pacific Ocean over the past decades. Data from: indi.ledso.columbia.edu/SOURCES/I N NOAA/NCEP/EMC/CMB/GLOBAL/Reyn_SmithOls2/ (Reynolds et al. 2002).
Tropical ocean hydroclimate and temperature from coral archives

Jessica Hargreaves1,2, K. DeLong3, T. Felis4, N. Abram1,2, K. Cobb5 and H. Sayani5

1st CoralHydro2k workshop, Sydney, Australia, 1 September 2019

The CoralHydro2k project of the PAGES 2k Network met at the University of New South Wales before the 13th International Conference on Paleoceanography in Sydney, Australia, for our first in-person meeting (pastglobalchanges.org/calendar/2019/127-pages/1744). The goal of CoralHydro2k is to build upon Ocean2k temperature reconstructions (Tierney et al. 2015) by using new and existing coral-based reconstructions to examine temperature and salinity changes (i.e. hydroclimate) through paired records of coral strontium to calcium (Sr/Ca) and oxygen isotopic ratios (δ18O; e.g. Felis et al. 2018). The 24 scientists, with three participating remotely, included working group members, coral experts, and experts in other fields. The workshop started with a review of the aims of the project followed by updates of current temperature and salinity reconstructions, and concluded with further discussions about the best practices for coral-based reconstructions.

One aim of CoralHydro2k is the generation and inclusion of new paired coral Sr/Ca and δ18O records that allow for investigations of seasonal-to-multidecadal variability of hydroclimate. The project incorporates these new records, along with those included in the Ocean2k dataset (PAGES2k Consortium 2017; Tierney et al. 2015), with a focus on coral records that allow not only for hydroclimate reconstruction but also temperature. The new CoralHydro2k database will compile a comprehensive reconstruction of the δ18O of seawater for the tropical oceans. This requires an in-depth understanding of each coral record, including the intricacies of how the coral grew and the sampling methods. As such, CoralHydro2k is also investigating the impact of these methodological influences on the reconstructions using the pseudo-proxy approach. During the workshop, various experts presented new coral records to be included in the database and additional approaches that have not been explored by CoralHydro2k. Subsequently, we have targeted a number of primary projects to be completed with a paper as the final deliverable:

1. Build a CoralHydro2k database
   Only a handful of century-long coral-based 818O seawater reconstructions currently exist. Based on these records and a number of emerging records since the Ocean2k reconstruction effort, the group has developed a database of the tiered records that fulfill criteria deemed important. The criteria for Tier 1 include time coverage (most of the 20th century), resolution (monthly to seasonal), and paired coral Sr/Ca and δ18O to reconstruct salinity. The other tiers relax these requirements. The database will incorporate best practices learned from previous 2k efforts including Iso2k (Konecky et al. 2020) and PaCCTS 1.0 (Khider et al. 2019).

2. Develop a δ18O of seawater reconstruction
   With the CoralHydro2k database, we are developing a spatial-temporal salinity and temperature reconstruction for the tropical oceans to identify the dynamical drivers of seawater 818O changes. This reconstruction of tropical hydroclimate will distinguish natural variability from anthropogenic trends and linkages between marine and terrestrial hydrocycles by comparing them with the observational datasets (Hakim et al. 2016) and simulations from climate models that include water-isotope tracers.

3. Pseudo-proxy assessment of coral reconstruction methods
   Use the pseudo-proxy approach to better understand how the various methods used in coral-based reconstructions impact proxy-climate calibrations and ultimately the hydroclimate interpretation. Parameters to be tested include skeletal thickening, annual average versus monthly resolution, proxy uncertainties, sea surface temperature data products, and methods used for micro-sampling, regression, and age-model development.

CoralHydro2k holds bi-weekly teleconferences that are open to anyone interested in joining the project; please contact the group leaders to get involved (pastglobalchanges.org/science/wg/2k-network/projects/coral-hydro/people).

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Figure 1: Extracting a core from a coral colony at Christmas Island (Indian Ocean). Photo credit: Jason Turl.
Integrating documentary evidence into climate reconstruction and impact studies

Martin Bauch¹, S. White², C. Camenisch³, Q. Pei⁴ and H. Huhtamaa⁵

2nd CRIAS workshop, Leipzig, Germany, 7-8 October 2019

PAGES’ Climate Reconstruction and Impacts from the Archives of Societies working group (CRIAS; pastglobalchanges.org/crias) held its second workshop at the Leibniz Institute for the History and Culture of Eastern Europe (GWZO; pastglobalchanges.org/calendar/2019/127-pages/1945). This was the second phase of Phase 1 of the working group and demonstrated progress on the goals defined during the first workshop held in Bern, Switzerland, in October 2018 (pastglobalchanges.org/calendar/2018/127-pages/1807).

These goals were defined as:
1. the development of methods and international comparisons in historical temperature and precipitation indices;
2. the development of Chinese historical climatology through calibration and verification with phenological and early instrumental data; and
3. the further integration of historical climatology results from beyond Europe, especially results from evidence not previously used for climate reconstruction.

Finally, (4) the group decided after the Bern meeting to work towards an online exhibition of historical climatological objects, in order to reach a wider public.

At the conclusion of the Leipzig meeting, participants discussed the organization of the results and ongoing research into articles for a journal special issue, which was subsequently submitted as a proposal to Climate of the Past.

Regarding the first goal, lead author D. Nash presented an advanced draft of the first global review of methods for producing temperature and precipitation indices based on written evidence. This CRIAS product will provide a foundation for comparing results across world regions and improving practices in this method for quantifying qualitative documentary evidence.

G. Adamson presented trials in the creation of indices by teams of students and experts, in order to establish the consistency of results and appropriate number required to produce valid results. S. White explained a Bayesian approach to indexing, which could draw on the wealth of historical evidence and probabilistic inferences that have necessarily been left out in current indices, and C. Camenisch examined its reconstruction skill in a sample of historical data.

Q. Pei discussed ways to examine indices, particularly the analysis of autoregression, to determine whether the statistics of index values resemble the statistics of actual seasonal and annual climate. These methodological investigations underlie research in progress and intended articles on cross-regional comparisons of extreme years for the proposed Climate of the Past special issue.

Significant progress was also made on our second goal. This CRIAS workshop represented our first cooperation with the Atmospheric Circulation Reconstructions over the Earth project (ACRE; met-acre.net) for the recovery of early instrumental data and production of high-resolution climate reanalysis.

F. Williamson from ACRE presented early instrumental data from East and Southeast Asia suitable for calibration and verification with Chinese seasonal temperature indices, and K. Lin discussed results of the Reconstructed East Asian Climate Historical Encoded Series project (REACHES; ndc.noaa.gov/paleo-search/study/23410) on databases of Chinese historical weather records, including methods for producing regional temperature and precipitation indices. Together, these re-examination of these Chinese databases and indices alongside early instrumental observations enables calibration and verification procedures that would bolster the data and enable comparisons with results from Europe and other world regions.

Regarding our third goal, S. Brönnimann provided a worldwide overview of early instrumental records in his keynote, and A.M. Burgdorf presented on a global inventory of climate reconstructions from documentary records since 1300.

In keeping with our global and comparative focus, the meeting included further presentations of descriptive, phenological, and early instrumental evidence from China, South Asia, and Africa, as well as European documentary data (e.g. histrhone.cerege.fr). New approaches to evaluate documentary data by means of text-mining word clouds (see Fig. 1) were presented as well, analyzing semantic shifts in written records over decades and centuries.

Finally, in preparation for the workshop, a “call for objects” was issued to encourage proposals from different academic communities for objects to form a virtual “Weathered History” exhibition, which curates objects related to historical climate change for a wider public. The exhibition focuses on the themes “Remembering Disaster”, “Adaption and Prevention”, and “Famine and Weather”.

The next CRIAS meeting will take place in Hong Kong in September 2020 (pastglobalchanges.org/calendar/2020/127-pages/2021).

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Figure 1: A word cloud created with Voyant Tools based on all reports on weather extremes and food prices/crisis in Giovanni Villani’s Nuova Cronica published in the 14th century.
Floods in a warmer world: Insights from paleohydrology

Juan Antonio Ballesteros Canovas¹, L. Schulte², J.P. Corella³ and B. Wilhelm³

Geneva, Switzerland, 11-13 November 2019

The second phase of PAGES' Floods Working Group (FWG; pastglobalchanges.org/floods) started in January 2019 and will run for three years. The main goal of this second phase is to develop the core activities jointly identified during the first phase (Wilhelm et al. 2017). This workshop in Geneva (pastglobalchanges.org/calendar/2019/127-pages/1943) allowed us to discuss and advance towards the realization of these objectives. Thirty-two researchers from 12 countries attended, of which more than 30% were early-career scientists. The workshop also benefited from contributions by Dr. Karen Sudmeier-Rieux, representative of the United Nations Environment Program, and Dr. Marie-France Loutre, Executive Director of PAGES.

The workshop was divided into two sessions. The aim of the first session was to discuss a range of methodologies that integrate multiple natural and historical archives to provide comprehensive paleoflood records for Disaster Risk Reduction (DRR) schemes, while the second session was mostly dedicated to discussing the occurrence and magnitude of floods during past warm and cold periods.

During the first session, the benefits of paleoflood information in DRR schemes were shown throughout different examples. A particular focus has been placed on the dimensioning of key infrastructures such as spillways in hydropower dams, or flood hazard assessments in regions with scarce instrumental data. The workshop participants agreed that advances in the inclusion of historical and paleoflood records in flood hazard assessment were achieved during recent years as well as in its incorporation to derive flood frequency (England et al. 2018). Participants also highlighted that to improve the applicability of paleofloods in DRR, the engagement of stakeholders is crucial.

Participants concluded that an integrated methodological and multidisciplinary approach is imperative to improve the potential replicability of paleofloods research worldwide. This effort, initiated during the FWG session at the PAGES Open Science Meeting 2017 in Zaragoza, is also documented by the 18 research papers compiled in the Special Issue “Pluridisciplinary analysis and multi-archive reconstruction of paleofloods”, which was published in the journal Global and Planetary Change (Schulte et al. 2019).

In addition, we noted the importance of enhancing the communication between paleohydrologists, hydrologists, flood engineers, and decision-makers is essential to anticipate impacts of floods. In particular, participants discussed the possibility of developing an interdisciplinary audiovisual project as a way to transmit the benefits of including paleofloods in DRR to a broader community. At the same time, participants have started to collect ideas for writing a joint paper on paleofloods and DRR.

During the second session, several reconstructions of past flood activity during cold and warm periods were shown worldwide. The workshop attendees discussed the importance of establishing the relationship between changes in the atmospheric variability and flood occurrence. Time-space analysis of regional flood-rich and flood-poor periods allows us to anticipate future changes through our understanding of the forces that control flood occurrence. According to the different methodological approaches and regional studies presented at the workshop, which mostly covered the last centuries, higher flood activity in many catchments was linked not only to warmer periods but also to colder climate pulses. Yet, flood magnitude and timing evolved differently in different records and regions.

Participants recognized that developing the global database could be challenging due to difficulties in data integration from different archives and the significant amount of time required for this project. However, we came to a consensus that completing the interactive database of metadata of existing records remains a priority (pastglobalchanges.org/science/wg/floods/wp1/data).

Finally, different upcoming FWG-related activities were presented, such as the 6th International Paleoflood Conference in New Zealand (January 2020); a FWG-related session at the EGU General Assembly (3-8 May 2020); and the possibility to organize the 7th International Paleoflood Conference in Zamora, Spain (2022). More information about the workshop, upcoming projects and activities is available on the FWG website.

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Wilhelm B et al. (2017) White Paper: For an improvement of our flood knowledge through paleodata. PAGES, Switzerland, 15 pp
## Announcements

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3 Past plant diversity changes and future conservation issues  
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Pascale Braconnot and Françoise Vimeux

6 How the Asian subtropical area became evergreen since the Last Glacial Maximum  
Zhuo Zheng, K. Huang, Q. Wan and R. Cheddadi

8 Past climate changes and the role of refugia in the temperate Northern Hemisphere  
Rachid Cheddadi and Keith D. Bennett

10 Next generation sequencing, phycogeography, and paleoecology  
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12 Survival and spread of arctic plants in response to climate change: DNA-based evidence  
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14 Ancient DNA from lakes: Insights into plant response to past climate changes  
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16 Modeling past plant species’ distributions in mountainous areas: A way to improve our knowledge of future climate change impacts?  
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18 Past changes in species diversity: A view from the mountains  
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