Chapter 9 Climatic Inferences from Dendroecological Reconstructions

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Abstract Tree rings have long been employed by ecologists to study the localscale dynamics of forest stands and woodlands, but only recently have network approaches been applied to evaluate regional and broader-scale processes. As with dendroclimatic data (e.g., ring-width and ring-density chronologies), climatic drivers become much more evident in dendroecological data aggregated at broad spatial scales (relative to local-scale data). Study of dendroecology networks has led to new insights on climatic variability and change and their impacts on ecosystems. In addition to the power of network approaches, dendroecology has advanced in recent decades because of the ready availability of, and comparison with, high quality, independent dendroclimatic reconstructions of various hydro-climatic parameters (e.g., drought indices, precipitation and temperature) and ocean-atmosphere indices (e.g., ENSO, PDO, and AMO). Dendroecological reconstructions that have been most commonly employed in climate-related analyses are disturbance histories (e.g., fire and insect outbreaks). We review examples of these applications from our studies in the Southwestern United States. We also compile and describe here, for the first time, a regional network of ponderosa pine (Pinus ponderosa) establishment dates from the Southwest, and we show that episodic natality patterns are probably associated with decadal wet periods. Using another example of decadal variability in forest fire histories-specifically a hiatus in fire occurrence in the circa 1780-1840 time period-we make a case that regional to continental-scale dendroecological reconstructions can provide useful insights about 'ecologically-effective climate change'. We define this type of climatic variability as the patterns of climate at interannual, decadal and centennial scales that are most distinctly reflected in synchronous ecological responses at regional and broader scales. In the context of dendroclimatology, primary values of the investigation of specific climatic patterns that elicit regional and broader ecological responses

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is that these studies provide insights about climate variability that is relevant to ecosystems, and in turn, human concerns about future climate change impacts on ecosystems.

Keywords Dendroecology · Forest fires · Insect outbreaks · Tree demography

9.1 Introduction

Dendroecology—the application of tree-ring analysis to ecological questions—is a rapidly expanding subfield of dendrochronology with increasing relevance to the study of past and present ecosystems and climatic variations. Ecologists have typically focused on short-term studies (<10 years) at fine spatial scales (<1000 m²), but the importance of long-term and broadscale processes is increasingly appreciated (Ricklefs 1987; Levin 1992; Turner et al. 1993; Brown 1995). Many ecosystem processes, especially those affected by climate changes, manifest themselves only over longer time periods and broader spatial scales than encompassed in typical ecological studies. Understanding the dynamics of long-lived organisms and ecosystems—and the role of climate in controlling these dynamics—requires decadal to centennial and landscape- to regional-scale perspectives. Various types of tree-ring data can provide the depth of temporal and spatial information needed for multiscale, comparative analyses to fully evaluate climatic effects on ecosystems (Fritts and Swetnam 1989; Schweingruber 1996; Swetnam et al. 1999).

The most common types of dendroecological datasets with relevance to studies of climate effects are disturbance histories (e.g., chronologies of forest fires and insect outbreaks) and demographic histories of tree populations (chronologies of tree natality and/or mortality). For both of these types of datasets, there are—or there is the potential for development of-broad networks of data from multiple sites that allow for regional-scale analyses. Disturbance and demographic processes are driven by both internal system dynamics (e.g., species life histories and community dynamics, such as competition) as well as by external factors such as climate. Generally, evidence for climate effects emerges only at larger spatial and longer temporal scales because of the 'noise' introduced by local, internal ecological processes. There have been a variety of dendroecological studies that primarily focused on internal stand or community ecological dynamics (e.g., Spencer 1964; Clark et al. 1975; Lorimer 1985; Frelich and Graumlich 1994; Kneeshaw and Bergeron 1998; Morneau and Payette 2000). These studies were carried out at the stand or watershed scales and climatic analyses were not included, or they were limited because of the fine spatial scale of the datasets. In contrast, broader studies examining climate effects on ecosystem dynamics have used network-based approaches involving many sites distributed across mountain ranges or regions (i.e., 10^4 to $>10^6$ km²; e.g., Kitzberger et al. 1997; Villalba and Veblen 1997a; Kitzberger and Veblen 1998; Swetnam and Betancourt 1998; Veblen et al. 1999, 2000; Brown and Shepperd 2001; Brown 2006; Sibold and Veblen 2006; Kitzberger et al. 2007).

Networks of well-dated disturbance and demographic chronologies can be aggregated at multiple spatial scales to enhance common patterns, just as is done in regional networks of ring-width or ring-density chronologies. Climatic influence emerges as patterns of synchrony (or asynchrony) of events, oscillations, or trends. Very broadscale synchrony (at $>10^4$ km² scales) is typically related to climate variability affecting the co-occurrence of ecological events in many places, because most ecological disturbances or processes are not capable of physically spreading over such large areas. For example, fires generally can not burn across the enormous desert valleys and canyons that separate forested mountain ranges in the American Southwest. Therefore, synchrony of annual fire events (or absence of fire events) among these widely dispersed mountain ranges has to be related to regional-scale, fire-inducing (or -suppressing) climatic conditions, such as regional drought (wet) events (Swetnam and Baisan 1996).

In general, synchrony of disturbance events and demographic processes at regional or broader scales can be inferred to reflect climatic influences. An apt analogy is crossdating of tree-ring measurements among trees and sites. In the case of tree-ring widths or densities, it is understood that numerous unique or local factors (e.g., soils, competition, genetics, human land uses, etc.) at the scales of trees and forest stands may affect cambial growth and thereby introduce nonsynchronous, non-climatic signals in tree-ring time series (Cook 1990). However, when sites and trees are carefully selected to maximize climatic sensitivity and ring series from many locations are combined, the existence of broadscale crossdating (i.e., synchrony) of ring patterns across regions is logically and demonstrably relatedboth statistically and mechanistically—to climatic variations (e.g., Fritts 1976). Likewise, disturbance and demographic events are affected by numerous local factors, but if common, synchronous patterns emerge across broadscale networks, those ecological patterns are most probably climatic in origin. Interpretations of these causal relations are most robust when coupled with mechanistic understanding of climatic/ecological processes, such as the effects of water balance on fuel productivity, fuel moisture content and fire occurrence (e.g., Swetnam and Betancourt 1998; Westerling et al. 2006).

Assembly of regional- to continental-scale networks of disturbance and tree demographic histories is just beginning (see the International Multiproxy Paleofire Database at: http://www.ncdc.noaa.gov/paleo/impd/). Analogous development of tree-ring width networks in North America and South America began in the 1950s with Edmund Schulman's work (Schulman 1956), and continued through the 1970s and 1980s with Harold Fritts' efforts (Fritts 1976, 1991). Many other dendrochronologists have assembled tree-ring data networks and contributed these to publicly accessible databases (see the International Tree-Ring Data Bank at http://www.ncdc.noaa.gov/paleo/treering.html). These datasets have now been applied in continental to global-scale dendroclimatic reconstructions (e.g., Fritts 1991; Briffa et al. 1994; Mann et al. 1998, 1999; Cook et al. 1999, 2004). In addition to providing an example for dendroecology of the power of the network approach, dendroclimatic networks provide a tremendous source of comparative data. Replicated time series of reconstructed hydroclimate variables or global

circulation indices are often used to evaluate the effects of annual to multicentennial climate variability and change on ecosystems. The easy availability of these climate reconstructions is now stimulating a surge of dendroecological research with a climatic focus.

The fact that synchronous ecological processes occur at very broad spatial scales (e.g., Hawkins and Holyoak 1998; Koenig and Knops 1998), and that this synchrony is driven in part by climatic variability (e.g., Swetnam 1993; Villalba and Veblen 1997a; Swetnam and Betancourt 1998; Brown and Wu 2005), raises important questions for both ecology and climatology:

- How can chronologies and networks of ecological patterns be used to study past climate variability and change?
- How does climate entrain ecological disturbance events and population dynamics (natality and mortality) at regional and broader scales?
- When and in what manner do climate-forced effects override internal or local factors (e.g., competition, predation, random variations) in determining the dynamics of ecosystems?
- How can we use our understanding of past climate influences on ecosystems at long-term and broad scales to predict impacts of future climate change on ecosystem structure, function, and disturbance dynamics?

In this chapter we describe several examples of ecological disturbances and population dynamics that are influenced by climatic variations. In particular, we illustrate how spatial networks are useful for evaluating climatic and ecological relationships by drawing upon our own fire history, insect outbreak, and tree population studies in the western United States. In addition to describing and discussing the ecological insights and implications of these tree-ring studies, we also focus on their relevance for climatological investigations. We compile both regional tree recruitment and fire occurrence data to highlight an unusual hiatus of forest fire occurrence in the western United States and southern South America during the early 1800s as an example of what we term here as an 'ecologically effective climate change.'

9.2 Examples of Dendroecological-Climate Reconstructions

9.2.1 Fire History and Fire Climatology

Fire history studies using crossdated fire-scarred trees are the most common type of dendroecological analysis involving assessments of climatic effects. There are numerous examples of subregional to continental (and even intercontinental) studies of fire climatology using fire-scar chronologies and various hydroclimatic reconstructions (e.g., see summaries in Veblen et al. 2003). Fire-scarred trees are a fortuitous ecological and physical phenomenon, whereby past fire events (primarily low-intensity surface fires) create very distinctive lesions within tree-ring series

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Fig. 9.1 Photograph of a cross section of a fire-scarred ponderosa pine (*left*), and map of locations of 120 fire-scar collection sites in the southwestern United States (*right*). The arrows on the cross section point to fire scars created when surface fires burned near the base of this tree. After each fire left a scar, subsequent tree rings grew over the dead tissue. The last fire scar in 1899, and subsequent century of growth with no fire scars, reflects the effects of fire suppression and the exclusion of widespread surface fires

(Fig. 9.1). The ubiquitous presence of fire-scarred trees in many forests worldwide has provided evidence to reconstruct detailed chronologies of fire history in hundreds of forest stands (e.g., Kitzberger et al. 2007). Fire history networks are now the most developed among dendroecological data networks, with more than 400 fire-scar time series included in the International Multiproxy Paleofire Database (http://www.ncdc.noaa.gov/paleo/impd/).

The power of a dendroecological network for assessing fire/climate relationships is illustrated here with an example from the southwestern United States. At present, there are a total of 120 crossdated fire-scar chronologies that have been compiled from sites in Arizona and New Mexico (Fig. 9.1). Site chronologies typically are composed of sets of 20 or more fire-scarred trees sampled in forest stands of about 10–100 hectares in size (see Swetnam and Baisan 1996, 2003, and references therein for details of site selection and other details). Compilation of fire dates recorded among multiple sites results in a regional fire chronology, which documents years when both highly synchronous fires were burning in many sites across the region as well as years when few fires occurred (Fig. 9.2).

The Southwestern regional fire chronology documents strong relationships both with hydroclimatic variables and with ocean-atmosphere indices, such as the El Niño/Southern Oscillation (ENSO; Figs. 9.3 and 9.4). Regional fire years—defined as years when fire scars were recorded on trees at more than 20 sites—mainly



Fig. 9.2 The number of sites recording fire scars in the southwestern United States from 1600 to 2000. This chronology of regional fire occurrence is based on the 120-site network shown in Fig. 9.1. The largest regional fires are labeled in the *upper* plot, and the *dashed lines* show the threshold of >20 sites and <4 sites used to identify 'large' and 'small' regional fire years, respectively. The *lower* plot shows the smallest fire years. Note that the values in the *upper* plot are shown as *black vertical bars* on a white background, and in the *lower* plot the values are *white vertical bars* on a black background. Because these were mainly surface fires burning through grass and herbaceous fuels, the widespread introduction of livestock grazing that accompanied Euro-American settlement in the late 1800s led to the decline of fires in virtually all sites

occurred during relatively dry years of a set of gridded, independently derived reconstructions of summer Palmer Drought Severity Indices (PDSIs; Cook et al. 2004; Fig. 9.3, top panel). In contrast, years during which almost no fires were recorded around the region tended to occur during relatively wet years. Regional fire years also tended to occur during years of cool sea surface temperature (SST) conditions (low Niño-3, an index of ENSO, which are La Niña years), while all but one of the small fire years occurred during warm sea surface temperature conditions (Fig. 9.3, bottom panel; El Niño years).

These patterns are supported by superposed epoch analysis (SEAs; Fig. 9.4). Superposed epoch analysis is used to compare average annual climate anomalies for the set of regional fire years to climate for the entire period of the climate reconstructions (Swetnam 1993). Superposed epoch analysis also is used to compare climate



Fig. 9.3 (*Upper plot*) Tree-ring-reconstructed summer (July–August) Palmer Drought Severity Index (PDSI; *solid line*) (from Cook et al. 2004), shown with the largest and smallest regional fire years identified from the 120-site fire-scar network from the southwestern United States (*lower plot*). Tree-ring reconstructed Niño-3 index of sea surface temperatures (SSTs; Cook 2000), shown with the same set of largest and smallest regional fire years

during years prior to fire years to assess antecedent conditions that may have been important for fire occurrence. Significant climate anomalies are assessed by using bootstrapped confidence intervals based on distributions of annual climate values. Results of SEA from the Southwest regional data confirm that, on average, the larger fire years occurred during drought years and La Niña events, and that the small fire years occurred during the opposite patterns of pluvial years and El Niño events (Fig. 9.4). Interestingly, SEA also often shows that there were significant lagging relationships in climate/ecosystem dynamics, with fire years typically following



Fig. 9.4 Superposed epoch analysis (SEA) of large and small regional fire years in relation to summer Palmer Drought Severity Indices (PDSIs) in the southwestern United States and Niño-3 sea surface temperature (SST). Plots **A** and **B** show results for largest (38) and smallest (36) fire years during the period 1600–1900, using tree-ring width reconstructions of southwestern PDSI (Cook et al. 2004). Plots **C** and **D** show results from SEA using the same sets of fire years against a reconstruction of Niño-3, an index of the El Niño/Southern Oscillation (ENSO) (Cook 2000). Plots **E** and **F** show SEA results using modern PDSI based on instrumental data and the largest (19) and smallest (23) fire years from area burned records from all federal, state, and private lands in Arizona and New Mexico during the period 1905–2004. Lines in each plot represent the 0.05, 0.01, and 0.001 confidence intervals based on Monte Carlo simulations of random distributions of annual climate conditions for PDSI and Niño-3

1–3 years of wet conditions. These fires were mainly surface fires, burning in grasses and herbaceous fuels, and wet conditions would have resulted in more continuous and denser fuels that would have burned more readily and have been widespread during subsequent drought years (e.g., Brown and Wu 2005).

Overall, the Southwest climate and fire comparisons (Figs. 9.3 and 9.4) not only illustrate the close coupling between wet/dry patterns and fire in this region, but they also provide a strong confirmation of the high fidelity of these two types of dendrochronological reconstructions. The regional reconstructed climate and fire time series are based on independently collected and quite different types of tree-ring data. The climate-sensitive trees used to develop the drought and Niño-3 reconstructions generally come from dry, rocky sites where fire spread is unlikely, and ring widths are the measured variable. The fire-scar records generally come from ponderosa pine and mixed conifer forests in the same region, but typically in sites where grass and pine needle understories could carry widespread fires. Fire scars are the primary observed variable in these time series. Despite these differences, the remarkable coincidence of extreme regional dry (wet) years and high (low) fire occurrence suggests that the two kinds of network-based reconstructions are accurately identifying ecologically effective climatic conditions at annual resolutions.

9.2.2 Western Spruce Budworm Outbreaks and Climatic Entrainment

One of the most widespread forest defoliators in the western United States is the western spruce budworm (*Choristoneura occidentalis*). During its larval stage, this Lepidopteran moth is a voracious feeder on the buds and needles of true firs (*Abies* species) and Douglas-fir (*Pseudotsuga menziesii*) trees. The name 'spruce' budworm is a bit of a misnomer, because although spruce trees (*Picea* spp.) are minor hosts, the true firs and Douglas-fir typically suffer much heavier defoliation and higher mortality rates than spruces. Another closely related budworm species (*Choristoneura fumiferana*) occurs throughout eastern Canada and parts of the northeastern United States, where it is the most widespread and important defoliator of conifers. Numerous tree-ring studies have reconstructed outbreak histories of eastern and western spruce budworm (e.g., Blais 1981; Hadley and Veblen 1993; Morin et al. 1993; Swetnam and Lynch 1993; Krause 1997; Jardon 2001; Ryerson et al. 2003), making this an excellent candidate for development of regional (and perhaps continental) networks of outbreak chronologies.

The tree-ring basis for developing outbreak chronologies is the observation of very sharply reduced ring growth in the host species during the defoliation episode, which typically lasts for a decade or longer (Fig. 9.5). The methods involve multiple steps for confidently identifying reduced growth periods as outbreaks (including distinguishing the outbreak signals from potentially confounding climate effects; e.g., droughts), and compiling the observations into stand-level chronologies of outbreak events and subregional composites (see Swetnam et al. 1985; Swetnam and Lynch 1993; and Ryerson et al. 2003). Specifically, the process we have followed includes: (1) the separate development of 'host' and 'non-host' tree species ring-width chronologies from nearby sites, (2) detailed graphical and statistical comparisons of these chronologies against each other and with independent



Fig. 9.5 Photograph of reduced tree-ring growth in a Douglas-fir tree due to a defoliation event by western spruce budworm. This type of rapid and sustained growth reduction over a distinct period of time, often followed by a growth surge, is characteristic of defoliation events, and generally is not typical of drought responses

instrumental climate data, (3) the use of the non-host chronologies (if comparison tests are satisfactory) to 'correct' the host chronologies (i.e., to remove most or all of the climate effects from the host chronologies), and (4) validation of the corrected chronologies with independent forest entomology records (e.g., maps and observations of defoliation and/or insect population measurements). The correction procedure involves subtraction of a variance rescaled version of the non-host chronology (index values) from the host chronology.

Resulting 'corrected' budworm-affected chronologies from individual trees or stand averages can then be combined into network composites for subregions. Our original work on this focused primarily on stand-level averages of corrected host ring-width chronologies, and the subregional composites were expressed as 'number of chronologies (stands) recording outbreaks' each year (Fig. 9.6). In subsequent work we found that an 'epidemiological approach,' focusing more on the responses of individual trees and their collective responses, rather than solely on stand averages, was more effective in revealing outbreak timing and magnitudes. Composites of these data are expressed as numbers or percentages of trees (by site or in the whole network) recording an outbreak each year (Fig. 9.6).

One of the consistent findings of our western spruce budworm studies in four subregions of the western United States is a general correspondence between budworm outbreaks and wet periods, coupled with reduced budworm outbreaks (endemic periods) during drought episodes (Fig. 9.6). In detailed analysis of precipitation and temperature variables in the southern Rocky Mountains of New Mexico, Swetnam and Lynch (1993) found that a wetter spring through early summer season (March to June) was best correlated with the regional budworm composite record. In addition to concurrence of wet periods and outbreaks, in cross-correlation and cross-spectral analyses we found that increased wet conditions usually preceded the outbreaks by several years up to a decade. These findings suggest that wet conditions predispose trees and forests to budworm outbreaks.



Fig. 9.6 Composite western spruce budworm histories from four subregions in the western United States, compared with smoothed (13-year symmetrical weighted low pass filter) summer (July–August) Palmer Drought Severity Index (PDSI) reconstructions from independent tree-ring width chronologies (Cook et al. 1999). The smoothed PDSI line in the upper graph is tree-ring reconstructed and the smoothed *dotted line* is based on twentieth century instrumental data. *Vertical lines* connect the PDSI and budworm axes at the minimum (1 standard deviation) years of the smoothed PDSI. The Blue Mountains, Oregon, budworm data are partly described in Swetnam et al. (1995). The other budworm datasets are described in Swetnam and Lynch (1993; southern Rockies, New Mexico); Ryerson et al. (2003; San Juan Mountains, Colorado); and Swetnam, Archambault, and Lynch (unpublished; Sacramento Mountains, New Mexico)

The subject of potential climate entrainment of insect outbreaks—and on animal population dynamics in general—has been debated in the ecological literature for decades (e.g., White 1976; Royama 1984; Martinat 1987; Mattson and Haack 1987). A variety of theories have emerged, including the famous 'Moran effect,' based on Moran's theorizing about the oscillatory behavior and regionally synchronized lynx-hare populations of Canada (Moran 1953). In the case of the western spruce budworm, it seems likely that favorable moisture conditions may act through an increase in the quantity and improvement in the quality of tree foliage, the food base for these defoliators (Swetnam and Lynch 1993). Although the exact mechanisms remain unidentified, we think the consistent, coincident patterns in the tree-ring data of wet periods/outbreaks, and droughts/endemic phases over multiple centuries and large regions are quite compelling evidence that decadal moisture variability is an important driver of budworm populations.

Notable features of forest insect outbreak reconstructions from tree rings are the remarkable cycles or quasi-cycles that are evident in these time series (e.g., see Speer et al. 2001). The famous larch budmoth (Zeiraphera diniana) of central Europe is the clearest example of relatively strong cyclicity in forest insect populations. These outbreaks produce a distinct tree-ring signature of defoliation events, evident as sharply reduced ring widths and densities that have allowed for reconstructions of up to 1200 years in length (Weber 1997; Esper et al. 2007). The recent work of Esper et al. (2007) demonstrates a remarkable regularity and stability of larch budmoth cycles in the European Alps over the past millennium, with an average period of 9.3 years between outbreaks. The most notable finding in this paper is that the larch budmoth reconstruction (832-2004 CE) reveals the recent period (since 1981) is the most unusual in the entire reconstruction, with an unprecedented absence of outbreaks. Esper et al. (2007) show that this period corresponds to an unprecedented increase in temperatures in this region (from an independent tree-ring reconstruction), and conclude that nutrient cycling and other ecosystem processes operating in the Alps may be undergoing a drastic alteration. Although the exact mechanisms of this change are not known, they speculate that warming winter temperatures may have led to earlier emergence of larvae from their eggs in the late winter or spring before larch tree needles have emerged, leading to starvation and failure of the larch budmoth populations to enter an outbreak phase.

Our regional time series of western spruce budworm typically show more variable oscillatory behaviors than is the case with larch budmoth, with considerable variability in the periods between budworm outbreaks. Hence, budworm population fluctuations appear not to be very strongly cyclic. Nevertheless, as much as 50% to 60% of the variance of the regional composite time series is explained by cycles of about 25–35 years (Swetnam and Lynch 1993; Ryerson et al. 2003). Considering that these data are derived from tree rings, where cycles of this strength are virtually never observed in raw ring-width series from non-host trees, this in itself is a remarkable characteristic of defoliated host trees. Moreover, in cross-correlation and cross-spectral analyses there appears to be some coherence of these moderate-strength outbreak periodicities, and much weaker but similar periodicities in climate reconstructions (Swetnam and Lynch 1993).

A notable example of the use of tree rings in evaluations of animal population dynamics and cyclic environmental variables are the studies of Sinclair et al. (1993; Sinclair and Gosline 1997) of hare populations, climate, and sunspot cycles in Canada. Unfortunately, the physical tree-ring basis of the hare population reconstructions (feeding scars on aspen stems) seems obscure and is not well illustrated or described in these papers. Some of the quantitative time series analyses are also difficult to evaluate. More research in this area, particularly with employment of more rigorous dendrochronological sampling and analysis procedures, may be fruitful in evaluating previous findings, and shedding light on the role of climate and animal population dynamics. New and powerful statistical-analytical tools that are increasingly employed by dendrochronologists and paleoclimatologists might be usefully employed in this endeavor (e.g., univariate and bivariate singular spectrum analysis (SSA), or wavelet analysis; e.g., Speer et al. 2001; Esper et al. 2007).

The periodic and aperiodic behaviors evident in tree-ring-based insect outbreak reconstructions, and their potential coherence with climate, deserve much more study. Most insect population dynamics studies have relied on much shorter observational datasets, which in some cases include only one to a few insect outbreaks (e.g., Royama 1984; Myers 1998) or population 'eruptions' (e.g., Ranta et al. 1997). Tree-ring-based insect outbreak and climate reconstructions for population dynamics research are of potentially great value because numerous population oscillations over periods of centuries can be identified at numerous locations over regions, and even continents. We think there are opportunities here for breakthroughs in our understanding of how climatic variations and oscillations act as potential synchronizing (e.g., the Moran effect) or disrupting factors in plant and animal population dynamics. Given recent enormous outbreaks of bark beetles in concert with climatic changes in North America (e.g., Logan et al. 2003; Breshears et al. 2005), and Esper et al.'s (2007) findings about changes in larch budmoth and temperature in the Alps, the importance of this topic is likely to increase in coming years.

9.2.2.1 Confounding of Dendroclimatic Signals by Insect Outbreaks

One final point regarding insect outbreaks is important in the context of this chapter and book; i.e., the potential confounding influence of insect outbreaks/defoliation on dendroclimatic interpretations from host trees. Dendroclimatologists have long been aware of this as a potential problem (e.g., Morrow and LaMarche 1978), and as a general practice they seek to avoid sampling sites and trees that have a known history of past outbreaks (Fritts 1976). A recent paper by Trotter et al. (2002) evaluated this issue in some detail in pinyon pine (*Pinus edulis*) stands near Flagstaff, Arizona, defoliated by pinyon pine needle scale (*Matsucoccus acalyptus*). They concluded that, indeed, chronic or episodic effects of this insect could alter the dendroclimatic estimates of drought if one were to use tree-ring width series from affected trees.

It is our opinion that the large majority of tree-ring width series that have been sampled and measured for dendroclimatic studies in the western United States do not have a great risk of being confounded with insect outbreak (or chronic insect feeding) signals. This is because most of the major episodic defoliators of conifers (e.g., western spruce budworm and Douglas-fir tussock moth) generally tend to feed in stands that have relatively closed-canopy conditions, in moderate to highly mesic sites, and at mid-elevations. Stands with *Abies* as a dominant or codominant species, for example, are more vulnerable than pure Douglas-fir stands. In general, these kinds of stands are infrequently sampled for dendroclimatic purposes. The classic western United States drought-sensitive site is an open-canopy conifer stand, in a xeric site, at the lower forest border, with steep slopes and shallow soils (Schulman 1956; Fritts 1976). These are typically the least likely stands to be attacked by western spruce budworm. High-elevation, near-tree-line stands that are often sampled for temperature reconstructions also are generally not affected by budworm or other major defoliators. Although some insect defoliators and stem feeders (e.g., bark beetles) can affect large areas, to the best of our knowledge, past outbreaks have tended to occur in a relatively small proportion of the range of host species.

In addition to the reasons listed above, episodic or chronic defoliation, or past insect-induced mortality within stands is usually visually obvious on trees within the stands, or in the sampled ring-width series (e.g., Fig. 9.6; but see Ryerson et al. 2003 for a case study where outbreak signals were not often visually obvious in the 'uncorrected' ring width series). Within stands, past defoliation events are usually apparent in the presence of trees with old dead tops (spikes), and new dominant crown leaders, and many dead branches that are not simply the result of great age. Also, dead trees from past killing events may be present, and in the case of bark beetles, telltale feeding and egg-laying galleries may be visible on the stems of dead trees for many years after the event (e.g., Perkins and Swetnam 1996).

In summary, the potential confounding effects on tree-ring series of past insect attacks—and other disturbances such as fires—are an important consideration that dendroclimatologists should be more keenly aware of than they generally are. Precautions should be taken to investigate what is known from documentary sources and local experts about the potential occurrence of past outbreaks in areas to be sampled. Dendroclimatologists should have some field and laboratory training and experience in identifying signs of past outbreaks within stands and on tree-ring specimens. We encourage greater awareness of this issue by dendroclimatologists. We also recommend further study of the potential 'contamination' of dendroclimatic databases with insect and other disturbance signals. Perhaps a screening of existing dendroclimatic databases is warranted for potentially unknown insect outbreak signals that may exist in some chronologies from host species in areas with known outbreaks. However, it is our expectation that if there are any such problem chronologies, they are likely to be few.

9.2.3 Regional Tree Demography and Climate Effects

Ecologists have long recognized that time series of tree births and deaths are of fundamental value for understanding forest and woodland dynamics (Blackburn and Tueller 1970; Henry and Swan 1974; Harper 1977). Estimating the age of a tree by counting its annual rings is probably the best-known tree-ring application in history, extending at least back to the writings of Aristotle. However, from a dendroecological perspective, the determination of tree ages is more complex than simply counting rings, and understanding forest demographic patterns requires much more than merely sampling what one suspects are the oldest trees in a forest. Because rigorous dendroecological analyses of tree demography are somewhat less well described in the literature than disturbance history methods (e.g., fire-scar and fire-history methods), we briefly describe here some key aspects of this tree-ring application in the context of studies that are relevant to climatic influences on forests and woodlands.

There are several important considerations in determining ages of individual trees and hence the age distribution of a forest stand or landscape. First, obtaining accurate dates for tree-ring sequences is of primary importance for ecological and climatic studies, since identification of synchrony of events and processes requires high temporal resolution (Baumgartner et al. 1989; Fritts and Swetnam 1989). Dendrochronological crossdating of ring sequences (Stokes and Smiley 1968; Swetnam et al. 1985) should always be used (or at least attempted) to derive absolute dates for tree-ring sequences. Even in studies where temporal precision of recruitment dates is on the order of 10–20 years, it is usually impossible to know how much error is involved in ring-counted data without first crossdating at least some of the tree-ring sequences. Crossdating is particularly critical for demographic studies in forests that have been selectively harvested (Brown and Cook 2006). Sampling of only living trees in harvested stands may give a biased perspective on the age distribution of the historical forest, as larger (and, hence, generally older) trees are often removed in selective harvest procedures.

In practice it is difficult to determine the exact germination dates of trees. This is due both to limitations in the ability to obtain ring series at the exact point of seedling germination (the 'root-shoot' boundary) and typically indistinct growth patterns in the very earliest years of seedling growth that make identification of annual ring boundaries difficult. Using the most intensive techniques and under the best circumstances, obtaining annual or nearly annual resolution tree germination dates is possible by destructively sampling and re-sectioning trees at multiple levels across the root/shoot boundary (Savage et al. 1996; League and Veblen 2006). However, most studies are limited to pith dates on ring sequences taken from some height above the original germination point; i.e., above the root-shoot boundary (e.g., Villalba and Veblen 1997b; Brown and Wu 2005; Brown 2006). These studies have generally used the terms 'tree establishment' or 'tree recruitment' to refer to the date or period when a tree became successfully rooted as a seedling, rather than the exact date of germination. Often establishment periods of 5–20 years' resolution are used to reflect the relatively low precision in estimates of tree ages.

It is also often difficult to obtain exact death dates of remnant trees. Reasons include difficulties in obtaining usable tree-ring samples from often highly decayed wood, and irregular ring characteristics near the outside, especially in very old trees that were stressed at the time of death. Death dates on remnant trees are also often difficult to determine because sapwood typically decays rapidly after tree death (especially in logs on the ground), which removes an unknown number of rings from the outside of the stem. Again, however, using the most intensive techniques

and with careful consideration of available evidence, annual or near-annual resolution of death dates can be obtained for examination of stand- to landscape-scale mortality events (e.g., Margolis et al. 2007).

Another requirement of tree demographic studies is an adequate sample size. However, no studies to our knowledge have addressed how many trees may be needed to adequately characterize population age structure in varying forests, and more work is needed in this regard. In old, uneven-aged forests, sometimes hundreds of trees must be sampled to obtain adequate characterization of age structure distributions. This requirement is mainly due to the decline in survivorship of trees as they age (often following an inverse J-shaped curve). This decline in quantity or quality of evidence with increasing time before the present is what has been referred to as the 'fading-record' problem, and it is common to paleoecological studies. The oldest trees that represent germination pulses, or 'cohorts,' in the earliest periods may not be detected unless numerous trees are sampled, but again, few studies have addressed this question in any type of systematic manner (but see Johnson et al. 1994). Furthermore, many studies have found relatively poor relationships between tree size and age. Therefore, some sort of systematic sampling procedure should be used to select trees for aging rather than merely selecting the largest trees in a stand.

Similar to tree recruitment dates, preservation of dead trees is a declining function with time before present, and obtaining estimates of past mortality events depends both on persistence of woody material and the ability to adequately sample the material to obtain death dates. Old forests usually contain complex recruitment and mortality patterns, and to temporally resolve the different pulses and hiatus periods of establishment and mortality events often requires very large sample sizes and careful sample techniques, particularly with sampling difficulties of obtaining the bark rings on dead trees.

However, despite these and other difficulties in obtaining high-resolution tree demographic data, concentrated efforts have resulted in very useful time series for evaluating population dynamics related to climatic influences. One of the most successful of these studies to date is the work of Villalba and Veblen (1997a) on *Austrocedrus chilensis* woodlands in Argentina. They demonstrated a clear set of linkages between favorable moisture conditions promoting regional tree cohort establishment, and unfavorable (drought) conditions promoting reduced tree establishment and mortality (Fig. 9.7). Another successful example is a recent study by Brown and Wu (2005) that illustrated the contingent effects of climate and fire occurrence on tree establishment in a ponderosa pine (*Pinus ponderosa*) forest in southwestern Colorado (Fig. 9.8).

Examples of regional- to landscape-scale tree natality, mortality, and climate associations found by Villalba and Veblen (1997a), Brown and Wu (2005), and Brown (2006) inspired us to undertake a 'meta-analysis' of regional tree establishment data in ponderosa pine forests from the southwestern United States. For this analysis, we compiled data from 12 studies that sampled hundreds of ponderosa pine trees distributed around the Southwest (Fig. 9.9). Some of these studies estimated stand densities (number of stems or trees per hectare) from a sampling of trees, while others reported total counts of trees establishing by date or period over



Fig. 9.7 Tree establishment and mortality dates from *Austrocedrus chilensis* woodlands in northern Argentina driven by fluctuations in effective moisture (Villalba and Veblen 1997a). (Reprinted from Villalba and Veblen, with permission from Journal of Ecology, Wiley-Blackwell)

a certain size class within fixed plots or belt transects. There were also differences in the establishment date (period) resolution in the datasets, including 5-, 10-, and 20-year periods.

Regional patterns of synchrony emerge in an overall compilation of the 12 time series in a ponderosa pine 'establishment index' (Fig. 9.10). This time series was derived from the logarithms of the time series values (tree numbers and tree densities; 1.0 was added to all values so that logarithms could be computed for years/decades with zero values) and the ratio of each log-transformed value to the mean of the entire log-transformed series. The regional time series is the average of all the establishment indices computed for each series (site). Logarithmic transformations are useful here because there is a strong tendency for recent periods (especially after 1900) to have one to several orders of magnitude more trees establishing (or surviving) than before this period. Hence, to evaluate relative patterns, the log transform and ratio index help to standardize all datasets for visual comparisons and combination in the regional index time series.

Overall, the regional composite (Fig. 9.10) shows that ponderosa pine forests are typically uneven-aged and the age structures are often dominated by episodes of tree establishment (cohorts) that alternated with multi-decade periods when relatively fewer trees established in the stands. Moreover, the composite suggests that there was substantial regional synchrony of these episodes of establishment (and lack of trees establishing), indicating the likely influence of broadscale climate forcing. In general, the largest cohorts correspond with wetter conditions, as is shown in the comparison with dendroclimatic reconstructions of drought indices (Cook et al. 2004) from the region (Fig. 9.10). In particular, the 1810s–1860s and 1890s–1930s cohorts coincide with exceptionally wet periods. Decreases in establishment



Fig. 9.8 Fire history, age structure, and climate variations in a ponderosa pine forest in southwestern Colorado, United States (Brown and Wu 2005). (a) ENSO time series (thin solid line is Niño-3 sea surface temperature (SST) index [D'Arrigo et al. 2005]; thin dashed line is Southern Oscillation Index [SOI, Stahle et al. 1998]). SOI is reversed to be consistent with other moisture indices. Years of significant triennial wet/dry oscillations identified by superposed epoch analyses (SEA) are shown by up arrowheads centered on the drought years. Biennial oscillations also were tested and found to be largely absent during the fire-quiescent periods of 1684–1724 and 1818– 1851. Thick solid and dashed lines show reconstructed hydroclimate time series, smoothed with 20-year cubic splines. The *solid line* shows the annual precipitation in northeastern New Mexico (Grissino-Mayer 1996), and the dashed line shows the Palmer Drought Severity Index (PDSI) for the Four Corners area (Cook et al. 2004). (b) Fire-year chronology for Archuleta Mesa. Horizontal lines mark time spans of individual trees, with fire scars designated by inverted triangles. Fire years at bottom are those recorded on more than two trees. (c, d) Tree recruitment dates by 5-year periods for (c) ponderosa pine and (d) other tree species. The shaded vertical bars connecting graphs in (a)-(d) mark relatively quiescent ENSO variability, wet periods, reduced fire occurrence and pulses of recruitment in ponderosa pine (modified from Brown and Wu 2005, reprinted with permission of Ecology, Ecological Society of America)



Fig. 9.9 Map of the Southwest and southern Rocky Mountains showing locations of the 12 sites where ponderosa pine age structure data have been collected and composted for analysis in this chapter. The *gray shaded* area is the approximate range of ponderosa pine in this region

of trees also appear to approximately coincide with drier periods in the 1750s– 1760s, 1850s–1860s, and 1950s–1960s. Note also that a most recent cohort in the 1980s–1990s also coincides with a wet period in the Southwest that occurred from approximately 1977 to 1992 (Swetnam and Betancourt 1998), although many of the datasets used here lack counts of tree seedlings and saplings in these recent decades. Earlier wet periods (1610s–1640s and 1690s–1710s) may coincide with very slight establishment episodes, but these cohorts are not well resolved, probably because fewer trees overall are included in this earlier part of the record.

Although this is a relatively coarse-scale spatial and temporal analysis of ponderosa pine age structure patterns in the Southwest, the results (Fig. 9.10) suggest that there are several key features of local to regional recruitment dynamics that were responsive to climatic variability. It is well known that ponderosa pine produces large cone crops only erratically, and that successful germination of the seeds is enhanced by warm, wet summers (Pearson 1950; Savage et al. 1996). Successful establishment and survival of seedlings into saplings is dependent on (1) favorable moisture conditions (i.e., lack of drought), (2) the absence of surface fires for a sufficient length of time to allow the saplings to develop thicker, more heat-resistant bark (e.g., Fig 9.8), and (3) sufficient time for meristems to be elevated above grasses and



Fig. 9.10 Composite age structure data from a network of 12 ponderosa pine stands and reconstructed summer Palmer Drought Severity Index (PDSI) from corresponding grid points in the southwestern United States (Cook et al. 1999, 2004)

other surface fuels. Hence, it seems quite likely that historically, the favorable germination and establishment periods of ponderosa pine in the Southwest corresponded with wetter conditions for the reasons described above; i.e., more successful germination and survival during these periods, and relatively longer intervals between fires during these wet periods (see also Fig. 9.8). It is likely that climate affected recruitment in many of these forests both through direct effects on demographic processes (seed production, germination, and seedling growth) as well as by modifying fire frequency and timing (and perhaps other disturbances, such as insect outbreaks; e.g., Fig. 9.6). In a later section of this chapter, we will discuss timing of longer fire-free intervals, and their relationships to both climate and patterns of regional tree recruitment shown in Fig. 9.10.

As in most dendrochronological studies involving networks of chronologies, regional-scale emergent patterns of ecological synchrony may be reasonably interpreted to reflect, at least in part, the important role of climatic variability. This is not to discount the important role of local factors in causing unique disturbances or demographic peculiarities at the tree or stand level. Interactions between

disturbances and demography (e.g., fire, insect outbreaks, and tree mortality and subsequent natality) can add multiple layers of complexity and non-climatic signals in composite, ecological time series. Nevertheless, depending on the extent and degree of spatial coherence and synchrony of disturbances or demographic patterns, the relative role of climate can be assessed by the nature of the aggregated time series, and their relations to independent climate reconstructions. The moderate degree of regional synchrony in southwestern US ponderosa pine establishment over the past several centuries, and approximate correspondence with reconstructed drought indices (Fig. 9.10), suggests that climate is at least partly controlling. We recognize, however, that multiple other environment factors probably also determine the nonsynchronous variations, especially at finer spatial scales of individual stands.

9.3 The Late Eighteenth-Century, Early Nineteenth-Century Fire Gap

We close our discussion of dendroecological and climate reconstructions with a specific example of an interesting climatic episode that has been identified in multiple fire chronologies. Fire historians in North and South America have long recognized a very interesting 'gap,' or hiatus, in fire events during the period from about 1780 to 1840. The gap appears as an unusually long interval (relative to average intervals in the rest of the chronologies) between fire events. This gap is evident in many forest stands from the Southwest (e.g., Swetnam and Dieterich 1985; Fig. 9.11; see also Fig. 9.8), Baja California (Stephens et al. 2003; Fig. 9.12), and northeastern Oregon (Heyerdahl et al. 2002). The gap seems to be detectable primarily in the intermediate fire frequency fire regimes (i.e., mean fire return intervals between about 7 and 15 years). The timing and length of the gap varies somewhat from area to area, starting or ending some years earlier or later, but generally within the period from the late 1700s through the middle1800s.

Kitzberger et al. (2001) noted that a similar gap of reduced fire frequency was present during approximately the same time period (i.e., 1780s–1840s) in fire histories from *Austrocedrus chilensis* woodlands in northern Patagonia, Argentina (Fig. 9.13). Perhaps most remarkably, the regional fire-scar time series from the Southwest and Patagonia were statistically coherent (i.e., synchronous), especially in the frequency range of about 2–7 years (determined in a bivariate spectral analysis). Kitzberger et al. (2001) hypothesized that the common response of fire regimes in the Southwest and Patagonia to the El Niño/Southern Oscillation (Swetnam and Betancourt 1990; Kitzberger et al. 1997; Veblen et al. 1999, 2000; Veblen and Kitzberger 2002) was likely the reason for coherence of the two fire occurrence time series from different hemispheres. Moreover, they proposed that the hiatus during the late eighteenth and early nineteenth centuries could be due to a combination of (1) reduced amplitude and/or frequency of ENSO events (especially La Niña events) during this period (Fig. 9.14; see also Fig. 9.8), and (2) a coincidence of this period with a major global cooling phase in the early 1800s (i.e., 1810s–1820s),



Fig. 9.11 Fire-scar chronologies from ponderosa pine and mixed-conifer forests in the Gila Wilderness, New Mexico (Swetnam and Dieterich 1985; Abolt 1997). *Horizontal lines* are treering records from individual fire-scarred trees along an elevational gradient, and the *vertical tick marks* are the fire-scar dates recorded on the trees. Note the 'gap' in fire occurrence in the period 1811–1837. The lack of fire scars after 1904 was due to livestock grazing and fire exclusion by government firefighters, and is a common feature of many surface fire chronologies in the western United States

which might have been associated with volcanic eruptions, and is clearly indicated in Northern Hemisphere temperature reconstructions (e.g., Mann et al. 1998, 1999).

Additional detailed analysis of the Southwest regional fire-scar series (Fig. 9.2), and a subset of this series, offers further insight on the nature and possible causes of the gap. Swetnam and Baisan (1996) found that an index of regional fire occurrence in the Southwest based on 63 fire-scar sites was highly correlated with tree-ring-width-based Palmer Drought Severity Index reconstructions from the region during the early to mid-1700s and after the mid-1800s (Pearson's r > 0.8). However, this correlation declined precipitously during the late 1700s and early 1800s (r < 0.3) before rising again after about 1850. This finding seems to support the interpretation that interannual climate variations were reduced, perhaps associated with a quiescent ENSO, and this may have caused an uncoupling from the fire occurrence pattern in the Southwest (see also Fig. 9.8, and Brown and Wu [2005] findings of reduced 'biennial' oscillations during this period).

Other dendroecological evidence also points to possible shifts in fire regimes during the late eighteenth- and early nineteenth-century period. An analysis of a subset of fire-scar chronologies from the El Malpais area of west-central New Mexico



Fig. 9.12 Fire-scar chronology from Sierra San Pedro Mártir, Baja California (Stephens et al. 2003). Note the gap in fire occurrence from about 1790 to 1830. Stephens et al. suggest the gap may be related to climate, or potentially land-use history (i.e., early livestock grazing by Spanish missionaries). Note that in contrast with the Gila Wilderness chronologies (Fig. 9.11), fires generally continue to burn in this area during the twentieth century, probably indicating that livestock grazing and/or fire suppression were minimal in this area of northern Mexico until after about 1950 (modified from Fig. 1 in Stephens et al. [2003], reprinted with permission from Canadian Journal of Forest Research, NRC Research Press)

indicated that the seasonal timing of fires also shifted during the late eighteenth century (Grissino-Mayer and Swetnam 2000). This analysis of the distribution of fire-scar seasonality found that late season scars were more prevalent in the period before the late 1700s than after. In this remote and rugged area (with impassable lava flows in places), human-set fires were unlikely. It is probable that climate shifts were responsible for a change in seasonal timing of fires. Grissino-Mayer and Swetnam (2000) hypothesized that the arid fore-summer/monsoon pattern may have strengthened after the late 1700s, resulting in more fires during the early to middle growing season (i.e., the arid fore-summer, May and June), but fewer fires during and after the late summer monsoon (July to August). Prior to this time, a weaker monsoon may have persisted for some decades (early to mid-1700s), allowing more fires to occur later in the growing season (i.e., July to August).

Other synoptic climate patterns also may have played a role in the fire gap. In recent multivariate and contingency analyses of 238 fire-scar chronologies from



Fig. 9.13 Fire frequency changes in Patagonia, Argentina, and the southwestern United States. Both regional composites from fire-scar networks show reduced fire frequency during the 1780s–1840s period (*vertical shaded bar*) (Kitzberger et al. 2001). The *dashed* and *solid lines* show 'moving' fire frequencies, computed as the sum of fire events that were synchronously recorded by the indicated numbers or percentage of sites within the networks during 49-year windows, plotted on the central (25th) year of the window (reprinted from *Global Ecology and Biogeography*, permission granted from Wiley-Blackwell)

across western North America, extending from southern British Columbia to northern Mexico, Kitzberger et al. (2007) showed that the most coherent regional signal in the fire-scar dataset was well correlated with ENSO (Pearson r = -0.47, p < 0.001, between the first principal component of the 238 series and the D'Arrigo et al. [2005] reconstruction of the Niño-3 sea surface temperature index). An independent



Fig. 9.14 Multiple reconstructions of the El Niño/Southern Oscillation (ENSO) show a reduced occurrence or amplitude during the 1780s–1840s period (*vertical shaded bar*) (Kitzberger et al. 2001). (a) La Niña and El Niño events reconstructed from tree-ring chronologies in Patagonia and central Chile (Villalba 1994). (b) Recurrence of moderate to very strong El Niño events reconstructed from archival documents (Quinn and Neal 1992). (c) El Niño recurrence based on years when d¹⁸O was > -16% (i.e., warm events) in the Quelccaya summit ice core record (Michaelsen and Thompson 1992). Plots are mean number of events per year based on moving 49-year sums of all indices. (d) Record of ENSO-related central Pacific upwelling based on d¹⁸O (%) coral from Urvina Bay, Galapagos Islands (Dunbar et al. 1994; 49-year running mean). In all cases the horizontal dotted line represents long-term mean values (reprinted from *Global Ecology and Biogeography*, permission granted from Wiley-Blackwell)

tree-ring reconstruction of the Pacific Decadal Oscillation (PDO; D'Arrigo et al. 2001) was weakly correlated with the second principal component (r = 0.17, p < 0.01). Moreover, an independent tree-ring width reconstruction of the Atlantic Multidecadal Oscillation (AMO; Gray et al. 2004) appeared to be associated with

periods of maximum coherence (cross correlations) among the fire-scar chronologies during warm phases of the AMO, and minimum coherence during cool phases. The longest and coldest AMO phase occurred during the period 1750–1849, encompassing the fire-scar gap observed in many of the sites in the network. These findings corroborate and expand a growing list of tree-ring studies demonstrating fire-climate teleconnections at mountain range to subregional scales in the western United States involving ENSO (e.g., Swetnam and Betancourt 1990; Veblen et al. 1999, 2000; Donnegan et al. 2001; Heyerdahl et al. 2002; Norman and Taylor 2003; Schoennagel et al. 2005), PDO (e.g., Norman and Taylor 2003; Westerling and Swetnam 2003; Hessl et al. 2004; Taylor and Beaty 2005; Schoennagel et al. 2005), and AMO (Brown 2006; Sibold and Veblen 2006).

9.4 Ecologically Effective Climate Change

It appears likely that changes in both fire regimes and increases in tree recruitment during the early nineteenth century in western North America and South America are related to changes in timing and strength of ENSO events (Brown and Wu 2005), with further possible modulation of the effects of ENSO (at least in North America) by the AMO (Kitzberger et al. 2007). The early 1800s regional cohort synchrony (Fig. 9.10) may be the best example of the contingency of favorable regional climate and fire conditions occurring together to promote successful establishment; i.e., wetter conditions and longer fire intervals (e.g., Figs. 9.8, 9.11, and 9.12). We suggest that overall, these findings and the widespread occurrence of the 'gap' in fire-scar chronologies in North and South America are compelling examples of an 'ecologically effective climate change' deserving of more focused analysis by paleoclimatologists and others. In particular, we posit that these ecological responses that occurred at the turn of the eighteenth to nineteenth centuries reflect one of the most important and ecologically effective climatic changes in the past three centuries in western North America and southern South America, causing changes in forest structure that persist to the present.

Most dendroecological studies involving climatic analysis, and most of the preceding examples in this chapter, focus primarily on the ecological implications of the responses of disturbances or demographic processes to climate variations and changes. This is natural and appropriate as a focus of dendroecological research. However, now that numerous case studies have been conducted demonstrating and evaluating ecological responses to climate, and broadscale network approaches using dendroecological data are increasingly feasible (and with regional datasets starting to become available), we propose that such dendroecological datasets and analyses be used more broadly for identifying important climatological events and processes. Glacier- and lake-level fluctuations are commonly used by dendroclimatologists and paleoclimatologists as corroborating lines of evidence for identifying climate events and changes. Fire and other dendroecological evidence are used less frequently for such validation (but see Cook et al. 2004 for an example where both fire-scar and lake-level data were compared with tree-ring climate reconstructions). In some cases, regional ecological time series may be less noisy and/or higher-resolution indicators of climate variability than glacier- and lake-level fluctuations.

It is clear that dendroecological data, especially networks of such data, can be very useful for clarifying and identifying climatic variations and changes *that* are important to ecosystems. We refer to this as 'ecologically effective climate change,' and we think that this category of climatic variability deserves special attention by ecologists and other researchers. As climate change due to greenhouse gas-induced warming continues and perhaps accelerates, it will be increasingly important that climatologists, ecologists, managers, policy makers, and the public focus their attention on understanding and anticipating the impacts of climatic change on ecosystems. This will require more effective integration and comparative analyses of dendroclimatic and dendroecological studies. It may well be that the ecologically effective climatic changes that dendroecology can address will provide some of the most sensitive and insightful information about climate change impacts. The great potential for ecological indicators is exemplified by time series of plant phenology, such as the timing of plant flowering (Cayan et al. 2001). These records (especially in networks) are now widely recognized and embraced as a key and valuable ecological indicator of climate change.

The final example we offer of the potential of using dendroecological responses to identify ecologically effective climate changes for guiding future research, and for applications, is the finding of the importance of wet/dry oscillations in fire occurrence in the western United States (e.g., Fig. 9.4). It was tree-ring analysis that first identified a strong statistical association between the wet/dry sequence of 1-3 wet years, followed by a dry year, and widespread fire occurrence during the dry year (Baisan and Swetnam 1990; Swetnam and Betancourt 1992; Swetnam and Baisan 1996). This finding has been replicated by numerous dendroecology studies, particularly in relatively dry ponderosa pine ecosystems where grass and other fine fuel production was important for historical fire ignition and spread (e.g., Veblen et al. 2000; Brown et al. 2001; Donnegan et al. 2001; Stephens et al. 2003; Brown and Wu 2005). Subsequently, analysis of modern fire and climate records (i.e., twentieth- and twenty-first-century data) confirmed that this pattern was important over large portions of the western United States during recent times (Knapp 1995; Westerling et al. 2002, 2003; Crimmins and Comrie 2004). Now, this wet/dry pattern, in combination with ENSO-based climate forecasts (and ENSO-fire associations), is used to develop seasonal 'outlook' (forecasting) products for fire managers (see Predictive Services at National Interagency Coordination Center, Boise, Idaho: http://www.nifc.gov/nicc/predictive/outlooks/outlooks.htm).

In conclusion, we are optimistic that dendroecological contributions to dendoclimatological research, and vice versa, will greatly expand in coming years. The interest in ecological responses to climatic variability and change is already very large and will certainly increase as global warming continues and its impacts increase. The signs of this scientific interest are already evident in increasing numbers of dendroclimatic and dendroecological papers with a climate focus in leading forest science, ecology, and climatology journals. A challenge to dendroecologists is to work collaboratively and to promptly contribute datasets to publicly accessible archives, in the same generous, ethical, and forward-looking spirit as the hundreds of dendrochronologists around the world who have contributed ring width and ring density chronologies to the International Tree-Ring Data Bank. A challenge to dendroclimatologists is to focus their studies on parameters and questions with relevance to ecologists, and in turn to utilize the climatically relevant findings and datasets that dendroecologists are developing to address new questions in historical climatology.

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