



A Framework for Determining Population-Level Vulnerability to Climate: Evidence for Growth Hysteresis in *Chamaecyparis thyoides* Along Its Contiguous Latitudinal Distribution

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The impact of ecological and climatological factors on individual organisms over time and space is inherently complex and creates substantial uncertainty about how climate change will influence the global biosphere. To understand some of this complexity, we investigated the factors influencing individual growth of *Chamaecyparis thyoides* over 61 years within 18 populations across the ca 1500 km and 11 degrees of latitude. We then applied a vulnerability framework to understand how the variability of tree growth response to climate varies between populations and regions across our network. Surprisingly, we found the growth of trees in the central portion of our network responded more synchronously to warming and drought than trees in the southern end of our network, suggesting greater vulnerability in the central populations with continued warming. Our analyses and framework approach revealed substantial complexity in growth responses to climate within and between populations. We found potential resiliency within all populations, but higher inter-population than intra-population variability in response to climate. We found that latitude was an important proxy for the growth response to temperatures during the non-growing season and spring, but that ecosystem structure can modify the growth response and vulnerability to drought during the summer. The range of growth responses to warming is greater in the southern populations than in more northerly populations. This asymmetrical distribution of growth response across our study network provides evidence for a kind of ecological hysteresis, more southerly populations could be more resilient with warming. Despite the fact that this species primarily lives in wetlands, we found drought stress to be an important constraint on growth. Our study and analyses help to explain the disparities between forecasts of how climatic change might impact tree species and ecosystems over space.

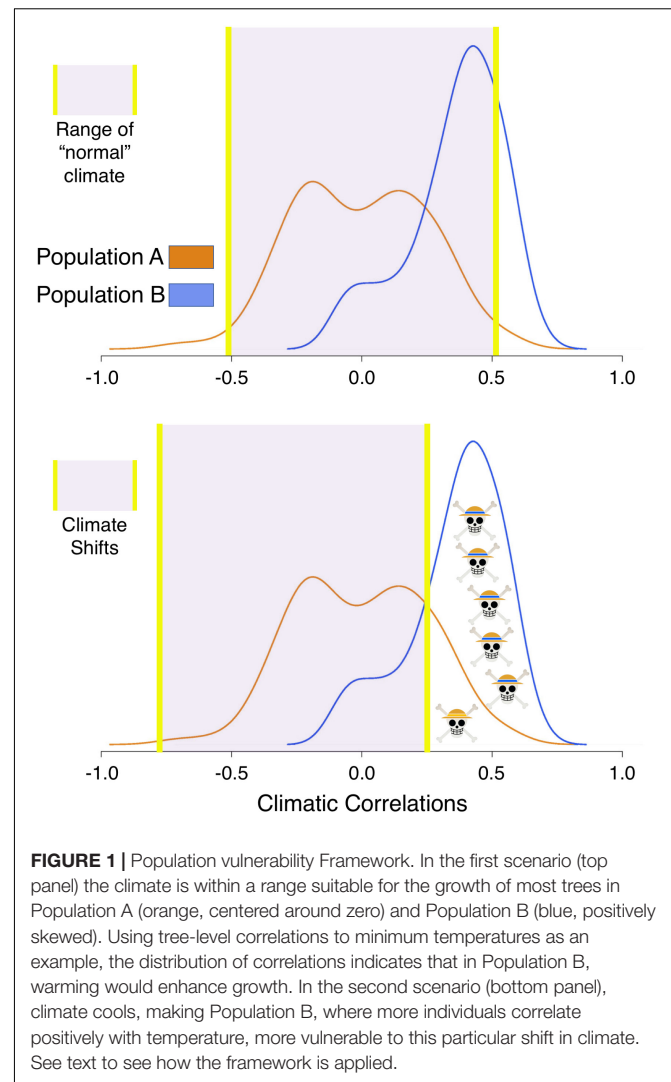
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INTRODUCTION

It is uncertain how global environmental change will alter ecosystems in coming decades (Friedlingstein et al., 2014; Duveneck et al., 2016; Swanston et al., 2017). A component of this uncertainty stems from the unique responses of individual organism to climate. The potential for great variability at the individual scale can add complexity to the dynamics and ecology of ecosystems. The multitude of ecological and climatological factors that influence plant growth are intrinsically complex, as they may compound and change over space and time (Billings, 1952; Turner, 2010; Heffernan et al., 2014; Zscheischler et al., 2018). Given the range of environmental heterogeneity and the uncertainty of future climate variability, forecasts on how the global biosphere will respond to future change are diverse and often contradict each other (e.g., Friedlingstein et al., 2014).

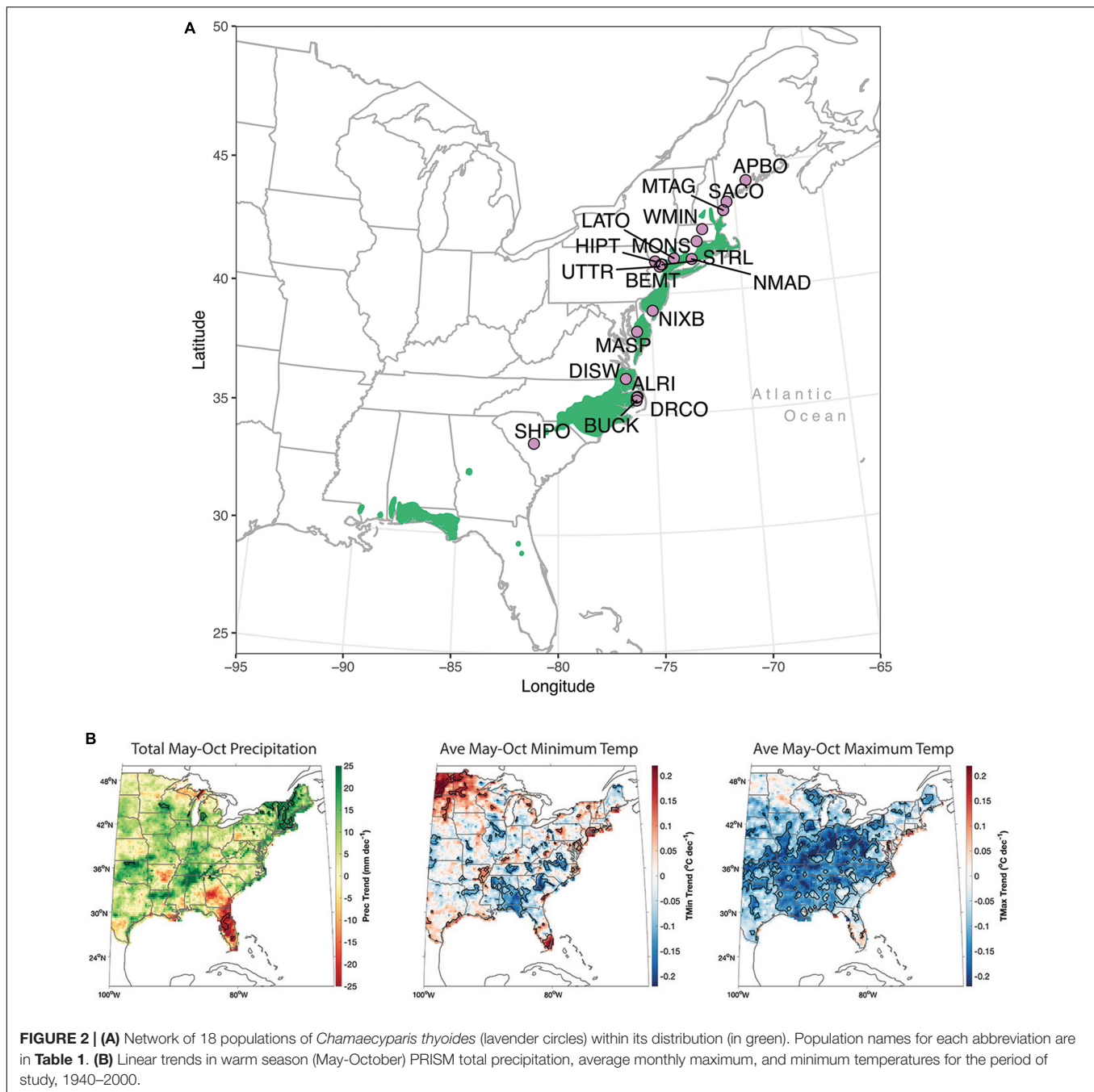
The urgency of understanding climate vulnerability of tree species and forests has stimulated research aimed toward quantifying the climate sensitivity of individual trees (e.g., Carrer, 2011; Clark et al., 2012; Itter et al., 2017; Laskurain et al., 2018). Traditionally, the climatic response of a species or populations has been determined using an averaged time-series of growth at the stand or landscape level (Fritts, 1976). While this approach maximizes the species climate response, it does so by reducing the inherent endogenous, genotypic and phenotypic, and exogenous complexity of individuals in a population. Because stand structure and microtopography have been shown to alter the climatic sensitivity of trees in a variety of forests and environmental conditions (e.g., Cescatti and Piutti, 1998; Bunn et al., 2005, 2018; Martin-Benito et al., 2011; Bottero et al., 2017; Lloyd et al., 2017), investigating the climate response of individual trees can be leveraged to illustrate how these interactions might increase or reduce potential vulnerability of a population to climatic change. Estimating the uncertainty of the response of vegetation to climate change, especially if we are to gain insight on the impact over a broader range of climatic variability, requires research that investigates the growth of individuals from population to broad regional scales over multiple decades.

Here, we introduce a framework to gain inference on population-level vulnerability to climatic variability through the examination of individual tree growth. The framework is based upon the interplay between the range of individual tree response to climate within a population and potential climatic change (Figure 1). To illustrate the framework, let's compare two populations, Populations A and B. Trees in Population A have a symmetrical distribution of climate-growth responses to summer water availability, where responses range from positive to neutral and negative. Trees in Population B have an asymmetrical distribution of climate growth responses with a strong positive skew. Within this framework, more trees in Population B would be expected to decline in growth during a severe, multiannual drought, where some trees in Population A may actually grow more with the same drought. In this case, Population A could persist through the survival of the individuals who respond positively to drought or, for the trees with a neutral response (correlation coefficient around zero), are rather unaffected by



drought. Thus, Population B would be more vulnerable to sustained reductions in water availability than Population A. Scaling up, if multiple populations across a region have a large range of individual growth responses, the species could persist at the regional level through the survival of individuals in multiple populations. The term used hereafter for populations where more trees respond to climate similarly is “climatic growth synchrony,” a term that may also be interpreted as representing populations that are potentially more vulnerable to climate change.

We apply our framework to 61 years of tree growth from a network of 18 *Chamaecyparis thyoides* populations that span ca 1500 kilometers of its contiguous latitudinal distribution, starting with its population in central Maine at the northern end of its distribution to a lower latitudinal distribution in central South Carolina (Figure 2A and Table 1). *Chamaecyparis thyoides* (L.) B.S.P. (Atlantic white-cedar) is an obligate wetland species that lives within 200 kilometers of the Atlantic Ocean and Gulf of Mexico. *Chamaecyparis thyoides* does not typically live more than 50 m above mean sea level and is currently surrounded



by a matrix of intensive land-use in a heavily populated portion of eastern North America. The species is a focus for local and regional conservation programs (see the Atlantic White Cedar Initiative¹) due to its foundational role in wetland ecosystems (Laderman, 1989) and its value to local biodiversity. Investigating its vulnerability to climate change is important for local and regional biological conservation goals.

Studies on the long-term growth of trees, including those on *Chamaecyparis thyoides*, have shown contrasting results on the

influence of local and regional influences on growth. Multiple studies suggest regional climate constrains growth such that it will be synchronous over broad regions despite differences in ecosystem structure, topography, and local hydroclimate (e.g., Brubaker, 1980; Cook, 1991; Graumlich, 1993; Meko et al., 1993; Cook et al., 2001). However, microsite topography has been shown to influence climate-growth dynamics of high elevational *Pinus balfouriana* and treeline *Pinus longaeva* (e.g., Bunn et al., 2005, 2018; Lloyd et al., 2017). Some studies of *Chamaecyparis thyoides* reveal large-scale synchrony in the climate-growth dynamics (Hopton and Pederson, 2005;

¹<http://www.atlanticwhitecedar.org>

TABLE 1 | Location, site information, composition, structure, and sample replication of 18 *Chamaecyparis thyoides* populations for this study, arranged from highest to lowest latitudes.

Site	Lat./ Long.	Elev. (m)	Distance from Ocean (km)	No. of total spp.	Basal Area m ² /ha	Percent Basal Area <i>Chamaecyparis thyoides</i>	No. of Trees (series)	Total Growth Time Span	Median Tree Age (min, max)
Appleton Bog, ME	44.333, −69.272	101	21.5	6	75.8	77%	21 (28)	1862–2002	121 (63, 156)
Saco Heath, ME	43.548, −70.470	42	8.6	4	31.0	83%	20 (30)	1874–2002	100 (69, 129)
Mt. Agamenticus, ME	43.231, −70.699	58	8.9	3	46.5	85%	16 (21)	1911–2002	68 (43, 92)
Westminster, MA	42.5267, −71.950	337	77.5	5	56.5	74%	56 (92)	1803–2014	110 (78, 200)
Monson, MA	42.060, −72.301	253	82.2	5	51.0	43%	16 (18)	1865–2002	116 (53, 138)
North Madison, CT	41.360, −72.650	94	10.4	-	-	-	48 (73)	1814–2015	145 (92, 202)
Lake Tonetta Park, NY	41.376, −73.583	164	35.2	-	-	-	22 (44)	1852–2015	139 (83, 164)
Sterling Forest, NY	41.186, −74.281	315	53.5	-	-	-	22 (38)	1818–2001	121 (93, 186)
Bellvale Mountain, NY	41.201, −74.329	353	57.8	-	-	-	40 (69)	1843–2001	131 (43, 159)
High Point, NJ	41.332, −74.658	464	88.6	5	71.2	68%	44 (62)	1807–2002	150 (104, 196)
Uttertown, NJ	41.115, −74.420	343	59.4	7	88.5	81%	41 (63)	1764–2014	134 (72, 238)
Nixon Branch, NJ	39.324, −74.913	9	15.4	-	-	-	10 (13)	1790–2000	110 (68, 211)
Mardela Springs, MD	38.485, −75.791	0	63.0	8	48.8	37%	17 (25)	1848–2002	69 (52, 155)
Dismal Swamp, VA	36.537, −76.469	19	51.9	2	40.0	85%	18 (28)	1921–2003	75 (34, 83)
Alligator River, NC	35.785, −75.885	1	13.2	-	-	-	4 (16)	1940–2002	62 (59, 62)
Buckridge, NC	35.622, −75.945	1	5.2	4	50.0	56%	33 (37)	1905–005	76 (55, 100)
Dare Co. Bombing Range, NC	35.743, −75.932	0	15.1	7	68.2	73%	20 (30)	1929–002	67 (45, 74)
Shealy's Pond, SC	33.863, −81.234	87	173.6	-	-	-	11 (18)	1837–2006	95 (59, 132)

Blank cells represent information where ecosystem-level data was not collected. Populations are arranged from northern to more southern locations within our network. Colors represent regions determined from cluster analyses with northern populations highlighted in pale blue, central populations highlighted in light pea green, and southern populations in pale red.

Pearl et al., 2017), while other studies indicate local hydrology and site conditions are an important factor in the growth, survival, and climate response of *Chamaecyparis thyoides* (e.g., Eagle, 1999; Mylecraine et al., 2005; Seim et al., 2006; DeBerry and Atkinson, 2014; Pearl et al., 2019). Likely, both regional and local climatic constraints operate at different scales or during different

periods in time, as has been shown for *Chamaecyparis thyoides* (Atkinson, 2019).

Important to the derivation of our research questions and hypotheses here, prior research on the long-term growth of trees indicates differences between conifers and broadleaf species in the climatic response over space (e.g., Martin-Benito and

Pederson, 2015; Cavin and Jump, 2017; Seftigen et al., 2018). Notably for conifers, but not for broadleaf species (e.g., Pederson et al., 2004; Tardif et al., 2006) cool temperatures tend to constrain growth in trees near the northern end of their distributions (e.g., Cook and Cole, 1991; Cook et al., 1998; Bhuta et al., 2009). Growth of populations of *Chamaecyparis thyoides* closer to their northern distribution is most strongly constrained by cool temperatures (Pederson et al., 2004; Hopton and Pederson, 2005; Pearl et al., 2017). Preliminary evidence from Hopton et al. (2004) indicates that precipitation and water availability tends to become more important to the growth of *Chamaecyparis thyoides* that are closer to the southern end of the distribution of the species, similar to the response of several species in mesic North America and Europe (Cook and Cole, 1991; Bhuta et al., 2009; Babst et al., 2013; Harvey et al., 2019).

Given these results on the long-term growth of conifers over space, we expect greater variation in the climate response, and perhaps lower correlations, of *Chamaecyparis thyoides* trees in the more southerly populations because reduced regional climate constraints, i.e., higher temperatures and more abundant precipitation, could allow for local variables, such as local hydrology, to play a more important role on growth of trees within each population. Due to the cooling during the summer over much of the southeastern United States during the late 20th century (Mascioli et al., 2017; **Figure 2B**), we expect that heat stress during our period of analysis could be relatively low, potentially reducing the influence of large-scale climate (Hiddink and Kaiser, 2005), and increasing the influence of local conditions even further.

We have four hypotheses that address two broad questions. First, we ask: “How do *Chamaecyparis thyoides* populations respond to climate across its latitudinal range?” We address this question by testing the following hypotheses:

- (i) growth in the majority of *Chamaecyparis thyoides* populations will increase with warmer winter and spring temperatures toward the northern end of our network,
- (ii) growth in the majority of *Chamaecyparis thyoides* populations will increase with more water availability (low temperature, high precipitation, or the combination of both) during the growing season, with stronger correlations to growing season water availability in the more southern end of our network than in the more northern end.

The next set of hypotheses derive from the second question: “Does synchrony in tree-level response to climate in *Chamaecyparis thyoides* change over space?” Building off of previous work cited above (e.g., Cook and Cole, 1991; Cook et al., 1998; Bhuta et al., 2009), we test the following hypotheses:

- (iii) tree-level climate responses in more northerly *Chamaecyparis thyoides* populations are more synchronous than more southerly populations due to the top-down climatic constraint of colder temperatures and lower precipitation limiting growth across the entire northern portion of the network.
- (iv) responses of more southerly *Chamaecyparis thyoides* populations to all climatic variables and seasons will be less

synchronous than northern populations because large-scale climatic constraints will be less limiting for growth.

To test these hypotheses and explore potential vulnerability of populations to climatic change, we calculate climate-growth correlations between annual growth at the individual and population scale, and annual climate variations from 1940 to 2000 across our study network. Because ecological and physical factors can influence the response of trees to climate, we also explore how stand attributes and site characteristics (forest density, age structure, elevation, etc.) and potential mechanisms that would account for the responses of individual, population, and regional vulnerability to climatic change.

MATERIALS AND METHODS

Research Network

We created a network of *Chamaecyparis thyoides* growth records from 18 populations in the eastern United States to assess the relationship between climate and tree growth along its contiguous distribution (**Figure 2A**). We selected *Chamaecyparis thyoides* ecosystems dominated by trees that were likely at least 100 years old. The network spans roughly 11° of latitude (33°–45°N), from South Carolina to population of *Chamaecyparis thyoides* at the highest latitude in Maine (**Figure 2A** and **Table 1**). In addition, sampling in most sites (except Alligator River and Buckridge) targeted canopy dominant trees, so their responses may not fully represent entire populations (Nehrbass-Ahles et al., 2014; Klesse et al., 2018b). We did core across a range of tree diameters and potential tree ages in an attempt to increase the ecological variability in our network (Pederson et al., 2004; Pederson, 2005).

The climate along the latitudinal gradient our network varies substantially. We used monthly precipitation totals and monthly mean maximum and minimum temperatures derived from the 800 m resolution PRISM Climate Group gridded dataset to characterize regional climatology across the network (Daly et al., 2008). Climate data was averaged for temperature and summed for precipitation seasonally for winter (December–February), spring (March–May), summer (June–August), fall (September–November), and the warm season (May–October). During our period of study (1940–2000), the coolest portion of our network was at higher latitudes (average daily winter minimum temperature of -9.7°C , average daily summer maximum temperature of 13.1°C) while the southern portion was the warmest portion of our network (-0.1°C in the winter, 19.2°C in the summer). Focusing on seasons that might be most pertinent to tree growth, areas at greater latitudes received the least amount of precipitation (550.7 mm of precipitation during the warm season, 267.1 mm during summer, and 1146.8 mm annually), while areas at lower latitudes received the most during the warm season and summer (669.7 mm and 378.3 mm, respectively) and the core of our study area received the most annually (1237.0 mm) (**Supplementary Table S1**).

Trends in precipitation and temperature from 1940 to 2000 vary substantially over space (**Figure 2B** and

Supplementary Figure S1). While warm-season average daily maximum temperatures have cooled, most strongly in the southern portion of our network, average minimum temperatures have warmed significantly annually, during the warm season, and during the summer along the coast (annual trends are similar, but not presented). The most consistent warming trends are rising minimum temperatures in southern New England and New York State (**Supplementary Figure S1**). It is important to note that our study period follows the Dust Bowl of the 1930s, one of the warmer periods across a large portion of the contiguous United States. Our study period also includes the 1950s to 1970s, a period of enhanced summer and winter cooling (Mascioli et al., 2017). In stark contrast to anthropogenic global warming, regional cooling in this part of the United States from 1940 to 2000 was likely due to natural climate variability (Mascioli et al., 2017). While there is drying during the summer in the lowest latitudes of our network, notably in southern Virginia and northeastern North Carolina (**Supplementary Figure S1**), our analyses of instrumental climate data across the eastern United States indicates there is a general pattern of wetting over our network annually and during the warm season (Usgcrp, 2018), consistent with Pederson et al. (2015) and Bishop and Pederson (2015).

Tree-Growth Data

Age structure of our data indicates that most sampled forests were likely recovering from either a period of timber extraction or forest management; management has been more recent in the more southern forests of our network (Scott Smith [resource manager of Dare Co. Bombing Range in 2003], John Taggart and Mike Schafale [Dismal Swamp], personal communication, June 2004). Despite this, the median age of sampled trees in each population ranged from 62 to 150 years (**Table 1**). Forest composition, tree density, and basal area were estimated using basal area prism measurements in 11 of our 18 populations. Diameter at breast height (DBH) was recorded for each cored tree as a part of a forest structure census. Population-level forest composition, tree density, elevation, and distance from the Atlantic Ocean are in **Table 1**.

Increment cores were dried, stabilized, sanded, and visually crossdated within each population using standard dendrochronological methods (Stokes and Smiley, 1968). Radial increments were measured to the nearest ± 0.001 mm. Hypothetical calendar dates for each ring, as derived from visual crossdating, were statistically tested and confirmed using the program COFECHA (Holmes, 1983).

To homogenize variance in all *Chamaecyparis thyoides* growth series associated with tree allometry, size, tree age, and potential growth releases or declines, we applied a two-step standardization process. First, we applied a modified exponential curve, or a straight line constrained to having a slope \leq zero, and then we used a cubic smoothing spline with a 50% frequency response cut-off set at two-thirds the length of each series (Cook and Peters, 1981; Cook, 1985). Prior to detrending, variance was stabilized by applying a power-transformation to all series (Osborn et al., 1997), so that residuals could be calculated by

subtraction from the curves, instead of ratios, to produce tree-ring indices (Cook and Peters, 1997). Final time-series of growth was produced using the robust mean of all indices within each population. We used the same standardization procedure on all cores within each tree to produce tree-level time-series of growth. Residual chronologies produced by the program ARSTAN (Cook and Krusic, 2008) were used in all analyses. Residual chronologies are processed to remove as much serial autocorrelation as possible, optimizing comparisons of interannual variability of radial increment without the influence of previous growth legacies or trends (Cook, 1985). Biweight robust means of all tree-ring indices were computed at the population level to develop population records of annual radial increment.

Our detrending approach, to reduce the influences of tree size, a dominant factor of growth (Foster et al., 2015), and competitive and disturbance processes (Cook, 1985; Cook and Kairiukstis, 1990), emphasizes the annual variation in growth. Part of the range in responses at tree level, therefore, may have to do with detrending leaving more non-climate signal in each tree, which evens out at population level.

Cluster Analysis

To determine the extent of common interannual variability in growth among trees across the *Chamaecyparis thyoides* network, we performed hierarchical cluster analysis (hclust package in R; Murtagh and Legendre, 2014; R core team, 2017) and correlation analysis on all time-series of growth at the population level. By using residual chronologies for these analyses, identified clusters represent populations with similar responses to interannual climatic variation. Based on a visual assessment of the dendrogram, we partitioned the network into regions based on a k-means approach to determine their spatial characteristics. We conducted a second analysis to identify potential regional clustering in the interannual variability of radial increment using a correlation matrix across the network. Clusters in this analysis were determined using the “corrplot” function in R (Wei et al., 2017).

Determining Climate-Growth Relations

Tree growth does not adhere to the human calendar. It might be initiated by factors and environmental cues that coincide with the human construction of seasons, but investigations into stem growth and climate-radial growth dynamics indicate that using climate seasons might not best capture when trees grow and what aspects of climate constrain or propel growth (Kim and Siccama, 1986). For example, most radial growth increment often occurs within 2–3 months and is mostly complete by late July in the Northern Hemisphere (Deslauriers et al., 2007; McMahon and Parker, 2015; Ziaco and Liang, 2019). High resolution analysis indicates the peak of climate-radial growth dynamics can occur in narrow bands that straddle month boundaries, like in Italy where peak dynamics occurs at the end of May and beginning of July (Carrer et al., 2017). Complicating this within our network, there are likely important differences in the timing of tree growth between populations due to differing seasonality over 11 degrees of latitude. That is, moisture stress might be more vital in May to trees in the southern end of our network while June is a

critical period of moisture availability for trees in the northern portions of our network. Restricting our analysis to the same common seasons for all populations regardless of their latitude could mask differences in how changing seasonality over larger scales could affect growth. With these concerns as a guide, we first sought to test whether an approach that reflected potential local regional growing seasons might aid us in discerning differences in climate sensitivity between populations over our study area. For simplicity in this test case, we used daily meteorological data from PRISM to define the growing season for each based on the time after the average last freezing day ($<0^{\circ}\text{C}$) of the year and the first freezing day in the fall of that same year between 1981 and 2017, the full period available at the time of this analysis (Prism Climate Group, 2020). Once this period was determined for each population, we summed or averaged all potential climatic variables during the growing and non-growing seasons for correlation analysis with population-level growth. This method prevented us from being able to assess sub-seasonal climate-growth relationships and masked sub-seasonal climate-growth relationships and the comparison of climate-growth relationships at the population levels. As a result, we abandoned this approach and re-started our analyses using more traditional approaches.

Our next approach to investigate potential differences in the response of growth to climate was to use traditional seasons: winter (prior-year December through current-year February), spring (March–May), and summer (June–August). Because stored non-structural carbohydrates are used for growth in subsequent years (Trumbore et al., 2002; Kagawa et al., 2006; Carbone et al., 2013), we included climate during the prior summer as a potential factor in the tree growth. We summarized this analysis at the regional level using the regions determined by cluster analysis. To compare and complement that approach, we then evaluated the climate response of *Chamaecyparis thyoides* growth at the population level using monthly climate variables. The nearest grid point to each population was selected to calculate correlations between monthly climate data and the residual chronologies for the 1940–2000 common period. We computed monthly Pearson's correlation coefficients against all climate variables. Note: climate analyses for individual tree growth was determined using this same methodology.

Using what we learned from the seasonal and monthly analyses, we then developed a data-driven approach where we identified the seasons that most strongly and most commonly drove population-level growth within each region that was determined by cluster analysis. In this way, seasons could be defined broadly and, occasionally, longer than the traditional 3-month season. We then made reciprocal comparisons for the dominant seasons in a particular region against the trees and populations in other regions within our network. For example, the strongest, most consistent responses to climate for populations in the central region were with total May–August precipitation and April–August average maximum monthly temperature. The growth of trees and populations in the northern and southern portions of our network were then correlated to these two seasons to compare all trees and populations using the same metric for determining potential differences

across our network. The strongest response of tree growth to climate in northern populations was average minimum monthly temperature from prior-year November through May of the current year. The growth response of trees and populations in the southern and central regions were compared to this season.

To compare the distribution of seasonal climatic responses within populations and between regions, we estimated whether the continuous probability distribution functions of climate correlations at individual-tree level differed between regions using the non-parametric two-sample Kolmogorov-Smirnov test. We also tested the stability of the continuous probability distribution functions of climate correlations using bootstrapping techniques. For the distribution of tree-growth correlations to climate at the regional scale, we ran 100 bootstrapped PDFs, removing 25% of the trees in each region. We also conducted bootstrap analysis on these distributions on populations at the regional level by removing 1 population for each region to determine similarities and differences between regions.

Potential Geophysical and Ecological Factors Influence Climate-Growth Relations

We used linear mixed-effect models (LMEM) to determine the effect of site factors on the climate responses of individual trees (Zuur et al., 2009), and to account for the lack of independence between observations caused by the hierarchical structure of data (i.e., trees grouped within populations). In addition, linear mixed effect models allow us to analyze the variance not explained by fixed effects by allocating the variance to different hierarchical levels or random effects, in our case trees and populations.

These models were fit for the 11 populations for which stand density was available (Table 1); stand density was inferred from measured basal area per hectare at the time of sampling. For our purposes here, stand density was considered as a population-invariant variable measured at sampling time. Other variables included for the initial tests were: tree species diversity; classes of understory shrub density (0, little to no shrubs to 4, high density); wetland type, genetic factors; median, minimum, and maximum diameter at breast height; median, minimum, and maximum tree age; latitude; longitude; elevation (m); distance to the Atlantic Coast (m); site slope; site aspect; distance to open water; and species-level plant traits derived from the TRY database (Kattge et al., 2011). Distance to open water is used as a proxy for rooting height above the water table. Distance to the coast represents the nearest point of the Atlantic Ocean in any direction from each site. Genetic factors were derived from Mylecraine et al. (2004). While the genetics of our populations in their analysis fell into one region, we used clusters within that region to put populations in classes from 1 to 10, where classes 1 and 2 were most related to each other and classes 1 and 10 were most distantly related to one another. Some populations in our study were sampled by Mylecraine et al. (2004), but several were not. For the populations not sampled for genetics, we assigned classes based on their proximity to the nearest population sampled for genetics. Wetland type was derived from Laderman (1989) and U.S. Fish and Wildlife Service (2019). Shrub density was classed

in categories of 0–25%, 25–50%, 50–75%, and 75–100% based on photographs from each site. Forest density and tree diversity was collected in 11 ecosystems using 2- and 3-meter factor cruise prisms (**Table 1**).

We started by fitting a model of growth indices with selected climate variables as fixed effects from those that are highly correlated with growth at tree and population level (see Results, **Supplementary Figures S3–S5**). Prior to fitting the models, all seasonal climate variables were centered and scaled while maintaining trends in each series. A level of $p < 0.05$ was used for significance testing of variables in the model. Radial-growth data can be highly autocorrelated, which could result in biased estimations of confidence intervals of parameter estimates in the models (Grégoire et al., 1995). Thus, we included an autoregressive parameter of order 1 to account for the potential temporal autocorrelation in growth indices within the same tree. Along these lines, because we used the residual growth indices, the autoregressive structure is greatly reduced (Cook and Kairiukstis, 1990). At this stage of model comparison, models were fit with maximum likelihood estimations. At each step, models were compared to the previous, more parsimonious model based on Akaike's Information Criterion (AIC) decrease to check for significant model improvements. This best-fit model was checked for several assumptions such as temporal correlation, normal distributions, and homoscedasticity of residuals (homogenous variance), as well as non-collinearity of independent variables. Multicollinearity of variables was checked by assuring that the variance inflation factor remained below a recommended threshold value of three (Zuur et al., 2010).

After a best-fit model with only fixed effects was found, we explored random effects by considering both intercepts and slopes at two hierarchical levels (tree, population). Because our aim was to analyze the variability of the effect of climate at individual tree level within populations, we included random intercepts and slopes for all explanatory variables, excluding interactions. We used the log-likelihood ratio chi-squared test to compare the importance of different random effect structure, i.e., the same variables, but different hierarchical levels sites or both sites and trees. All linear mixed effects models were fitted using the package *nlme* (Pinheiro et al., 2012). We estimated marginal and conditional R^2 for the model (including fixed effects only or both fixed and random effects) following Nakagawa and Schielzeth (2013) extended for random slopes by Johnson (2014).

All R packages used here are from in R statistical software (version 3.4.2; R core team, 2017).

RESULTS

Regional Radial Growth Patterns

Hierarchical cluster analysis and correlation analysis conducted on residual growth chronologies indicate four clusters of populations within our network (**Figure 3**). Moving from the northern end of our network to the southern end, the first group is composed of four or five sites, depending on the analysis, ranging from central New England to coastal Maine. One outlier population for this group, Saco Heath, Maine, correlates

positively with other northern populations, but hierarchical cluster analysis groups it with the southernmost populations. Correlation reveals similarly weak, but positive, correlations of the Saco Heath populations to the populations in North Carolina on the Albemarle Peninsula, Buckridge, and the Dare County Bombing Range (**Figure 3C**). Given the weaker correlation statistics and proximity to the more northern populations, we group Saco Heath with those populations, a grouping called hereafter the northern region. The next grouping falls in the north-central portion of our network and hereafter is called the central region. Both hierarchical cluster and correlation analyses cleave this portion of our network to the same size populations. We note that the annual growth of populations in the central region correlate from 0.2 to 0.4 to the growth of populations in the northern region (**Figure 3C**).

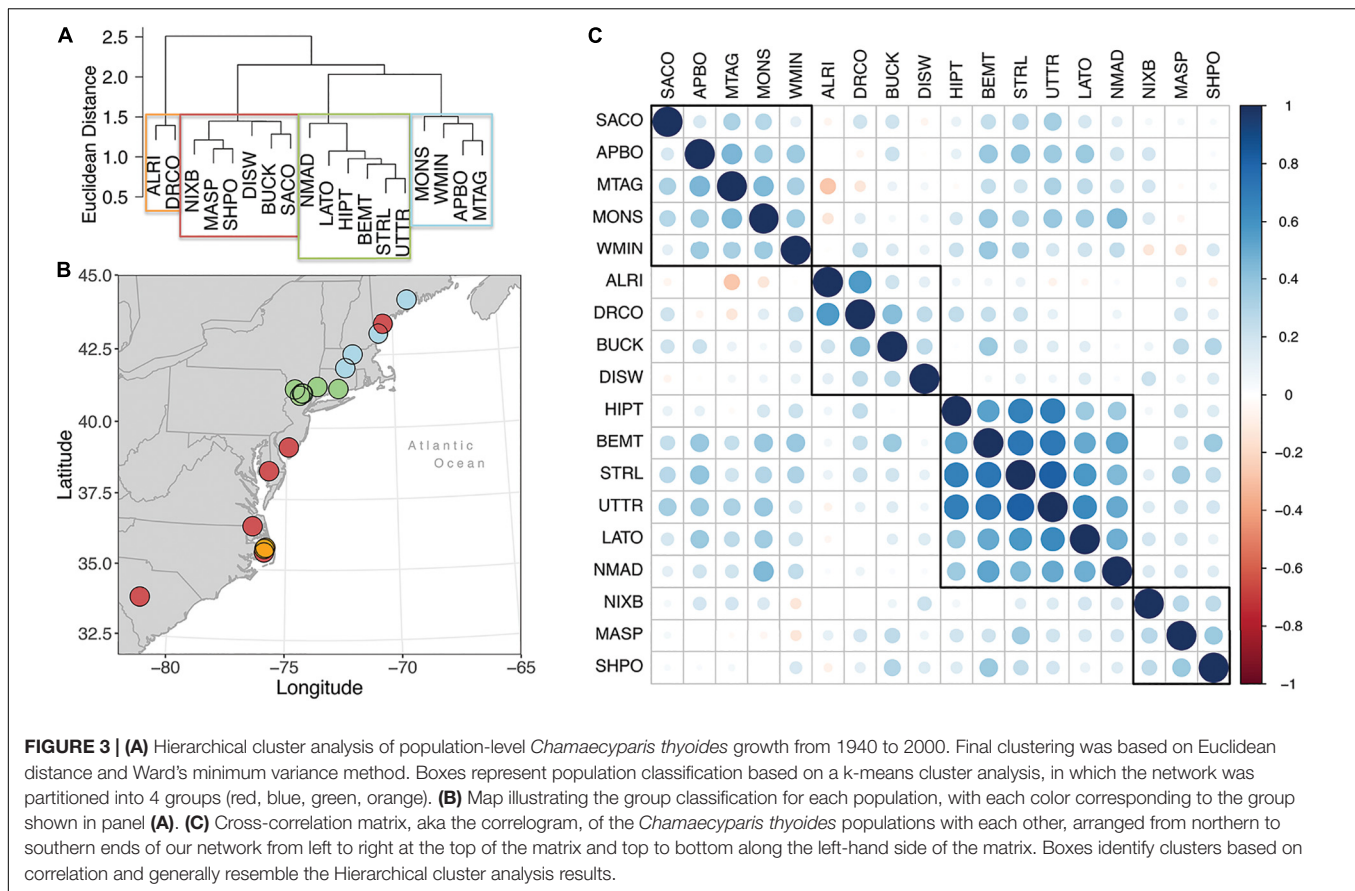
Cluster analysis separated the remaining sites less consistently. Hierarchical analysis indicates a cluster of five populations distributed across the southern half of our network, the aforementioned Saco Heath, and a closely related second cluster of two populations in northeastern North Carolina. Correlation analysis also identifies two groups in the southern end of our network and also indicates that the hierarchical cluster of these three population correlates nearly equally with both the hierarchical cluster of five population and population in the central region (**Figure 3C**). To simplify further analyses, all population from southern New Jersey to South Carolina are considered the same group, hereafter the southern region. The interannual to decadal-scale radial growth variation of all trees highlighting the mean, minimum, maximum, upper 95%, and lower 5% distribution for each year in each region is plotted in **Supplementary Figure S2**.

Regional and Population Growth Response to Seasonal Climate

Here we present population-level correlations to climate at the regional level (**Figure 4**). Additional figures showing the climate responses for each population within each region are shown in supplementary material (**Supplementary Figures S3–S5**). The **Supplementary Figures** generally support the findings presented in **Figure 4**. Results displayed in **Supplementary Figures S3–S5** helped us decide seasons for testing.

Precipitation

At the regional level, there are differences in the response of *Chamaecyparis thyoides* growth to total seasonal precipitation (**Figure 4**). Central region populations have the strongest relation to summer precipitation ($p < 0.05$; $r > 0.4$ in some cases), suggesting that precipitation increases growth in this region. Though weaker, the opposite pattern is apparent for responses to the prior summer. Precipitation weakly drives *Chamaecyparis thyoides* growth in the northern and southern populations. In contrast to most other populations, growth of southern populations generally increases with precipitation during winter (prior December–February), spring (March–May), and summer. Most notably, populations in this region show a greater spread in relation to precipitation during these seasons.



Maximum Temperature

The response of population-level *Chamaecyparis thyoides* radial growth to maximum temperatures at the regional level generally reflects patterns observed for total precipitation (**Figure 4**). The strongest and most consistent response of population-level growth to maximum temperatures is in the central region where higher temperatures during summer reduce growth, there is a negative correlation. As might be expected from the relation of growth among central populations to precipitation, there generally are positive, weak, but more significant relations to average seasonal maximum temperatures during the prior summer than to total precipitation during the same region. *Chamaecyparis thyoides* populations in the southern end of our network generally have negative relationships with maximum temperatures during fall of the prior year (prior September–prior November), current spring, and current summer. Positive relations between growth and winter maximum temperatures are seen in northern and central populations.

Minimum Temperature

The response of population-level *Chamaecyparis thyoides* radial growth to average seasonal minimum temperatures generally breaks from the patterns for precipitation and maximum temperatures. The strongest and most consistent response is in northern populations, where higher temperatures during the winter and spring increase growth (i.e., a positive correlation)

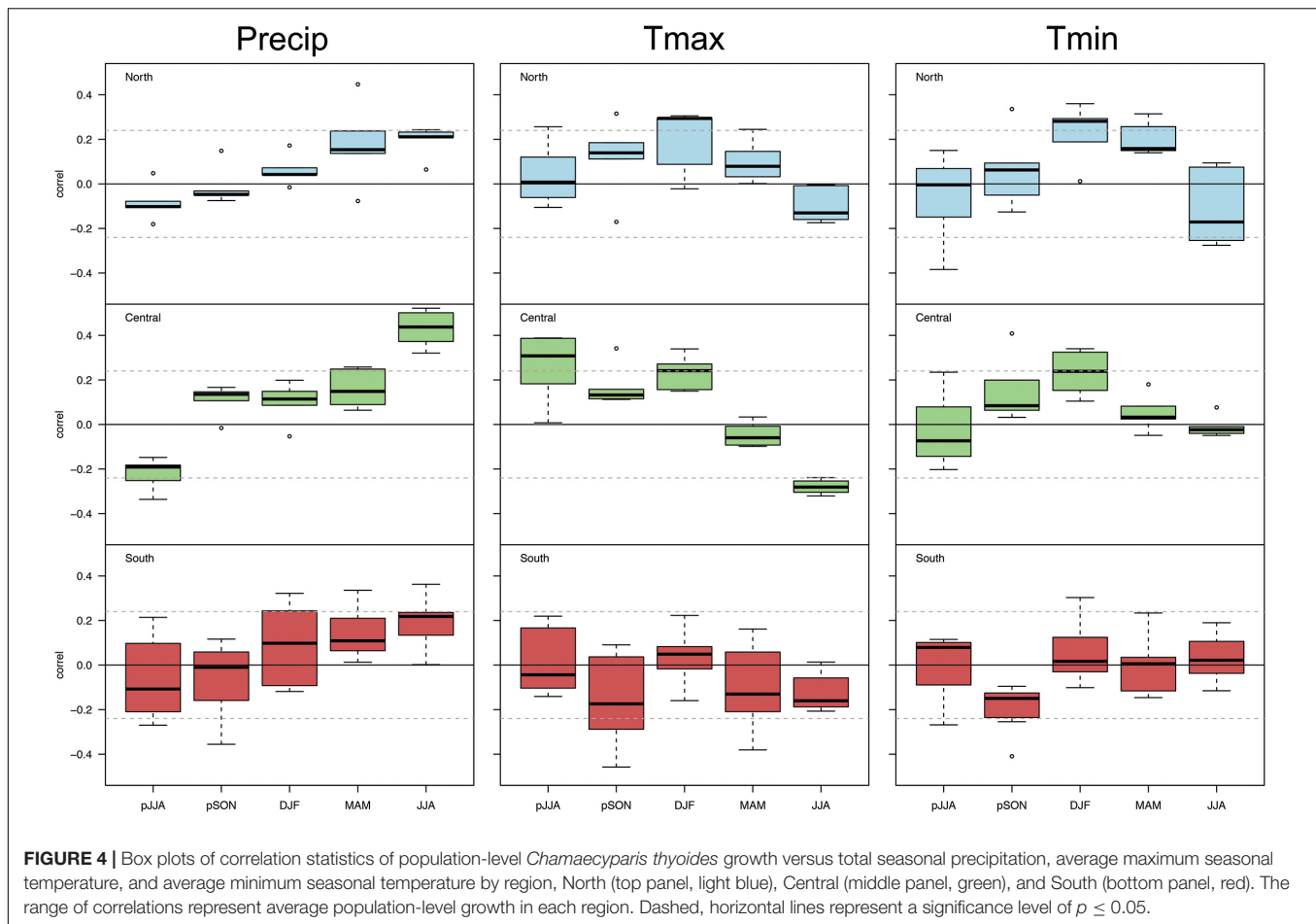
(**Figure 4**); the relationship during the spring is weaker than during the winter in this region. Within the central *Chamaecyparis thyoides* populations, the strongest and most consistent growth relations to minimum winter temperatures are found during the winter. In southern populations, growth relations were weak and inconsistent.

Tree and Population Growth Response to Seasonal Climate at the Regional Level

Here we present bootstrapped probability distribution functions of individual-tree and population-level correlations to climate at the regional level, **Figures 5, 6**, respectively. Additional figures of the probability distribution functions for each population within each region are shown in supplementary material (**Supplementary Figures S6–S8**). The **Supplementary Figures** generally support the findings presented in **Figures 5, 6**. We focus the results on the tree-level distributions, but include the population level results in the main text to show that responses to climate are rather consistent across scales.

Precipitation

The distribution of correlations with total precipitation align strongly with the previous seasonal and monthly population level analyses. The distributions at the tree and population level increased toward more positive values moving from winter to summer in all populations. This trend strongest in the central



populations. In fact, almost all trees in the central region are positively correlated to total precipitation in April–August, May–August and summer, indicating the most synchrony during those seasons (Figures 5, 6). Confidence intervals and cumulative distribution functions show that the central region precipitation response is also more consistently and statistically different from other regions during these warm seasons (Figures 5, 6 and Supplementary Figure S6). The distribution of tree correlations in northern populations are most different from those in other regions from prior November through current May and March through May where negative or neutral correlations are more prevalent.

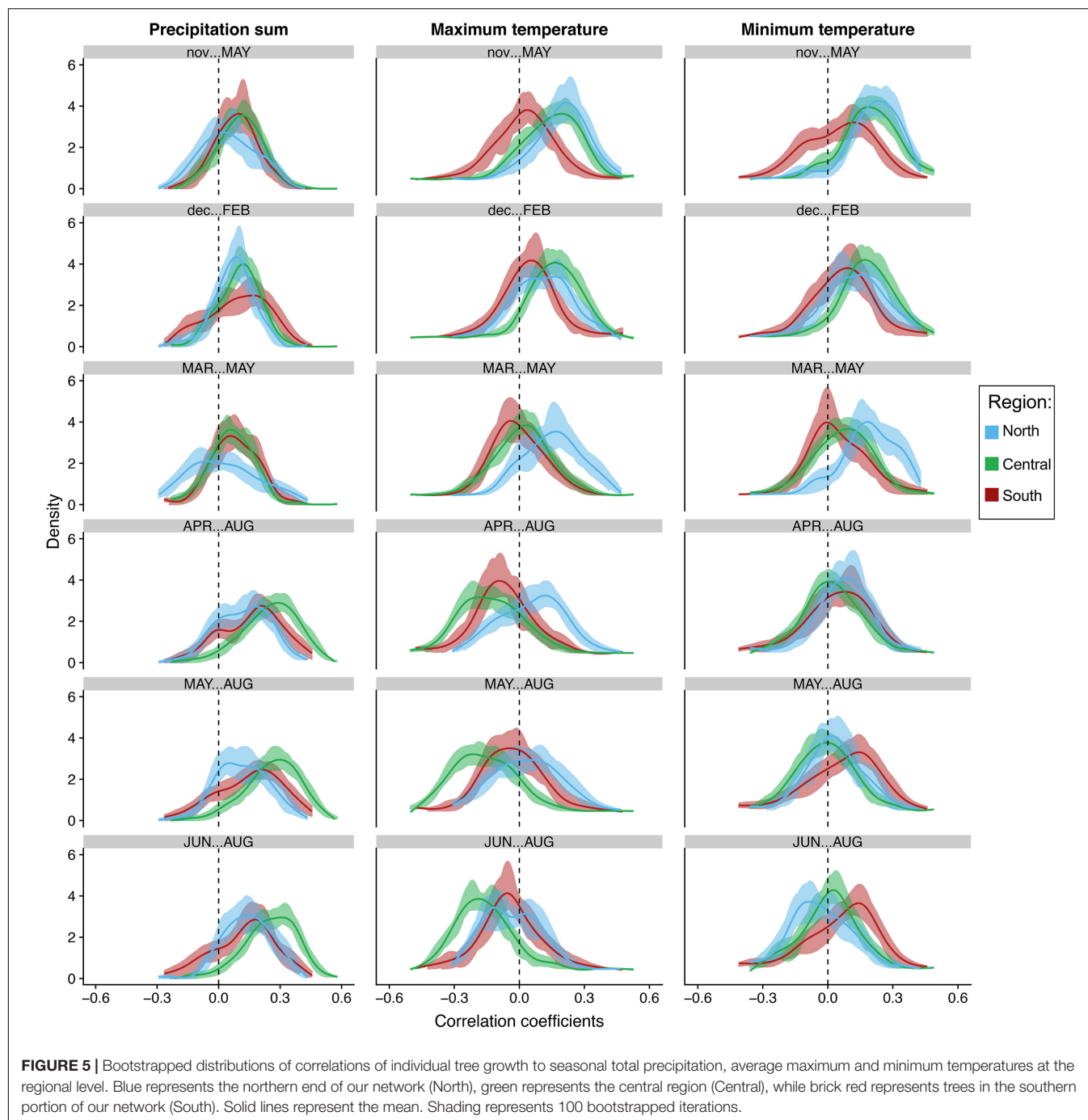
Maximum Temperatures

Positive correlations with maximum temperatures during the cool season shift to either more negative distributions or center around zero during the warmest seasons, except for in the more northern populations where they are more positive and significantly different than other populations until summer (Figures 5, 6). Correlation distributions in northern populations were more positive, and significantly different from other populations during March–May, April–August, and May–August. In contrast, distributions in the central generally are more negative during the warm seasons. In more

southern portion of our network, distributions against maximum temperatures trends more toward neutral, i.e., peak distribution closer to zero, from prior November–current May and in summer. Confidence intervals and cumulative distributions indicate substantial differences between all populations in five of the six seasons analyzed, prior November–current May, March–May, April–August, May–August, and June–August (Figures 5, 6 and Supplementary Figure S7).

Minimum Temperatures

The distribution of correlations to minimum temperatures are skewed toward positive values during the cool season in all regions and tend to become less positive and more neutral during the warm seasons (Figures 5, 6). The trend from positive to more values centered around zero is most strongly expressed in northern and central populations. Confidence intervals and cumulative distributions show the strongest differences in northern and central populations compared to more southern populations from prior November through current May (Figures 5, 6 and Supplementary Figure S8). Like the distributions for maximum temperatures, the northern populations skew significantly more positive during spring. It is interesting to note, however, that southern populations skew more positively

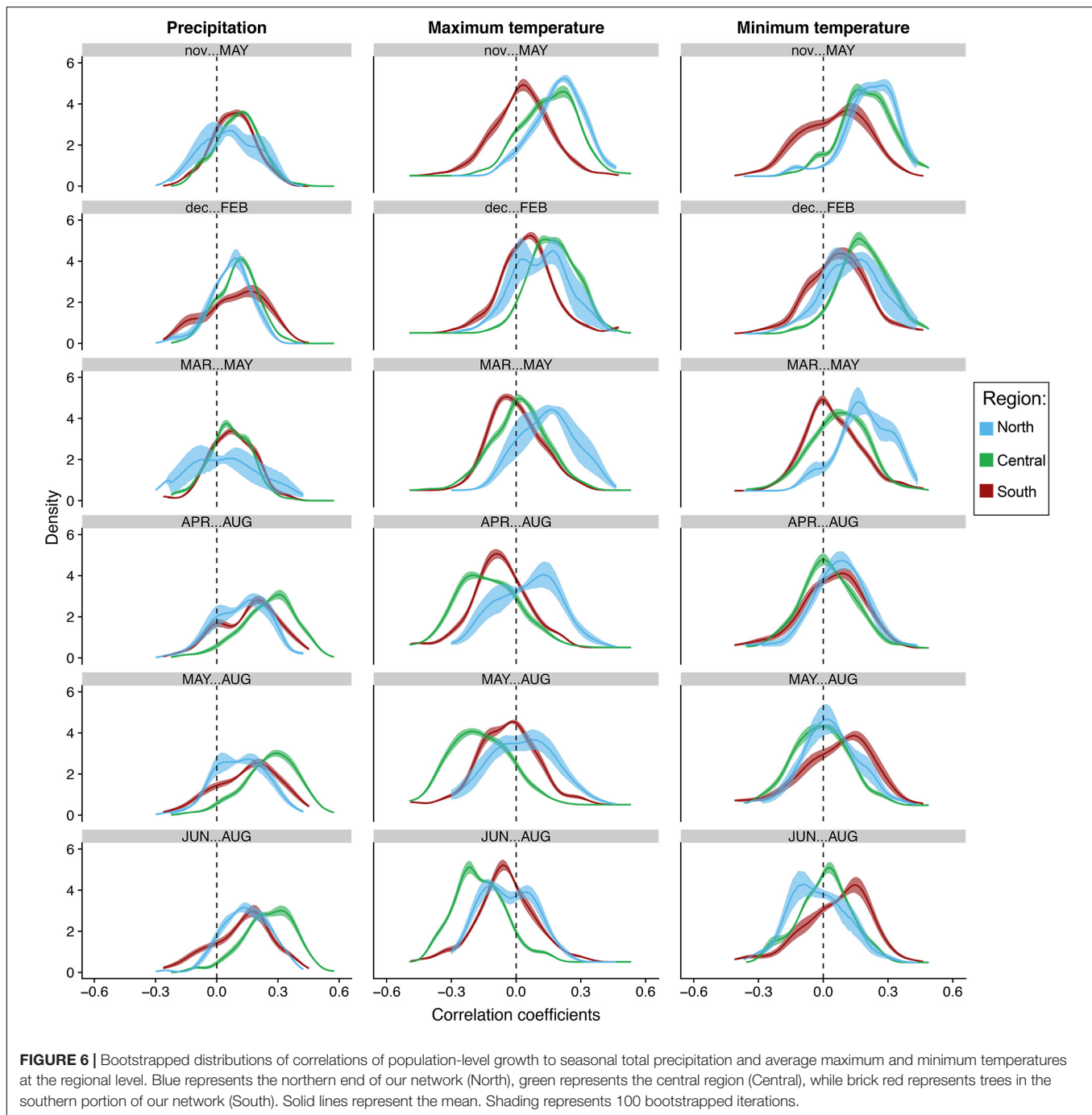


than the central and northern populations from May-Aug and during summer.

Geophysical and Ecological Factors Influence Climate-Growth Relations

Based on previous results from correlation of climate with mean growth at the population- and individual-tree level, our final linear mixed effect model included the three seasonal climate variables with some of the strongest and most divergent responses

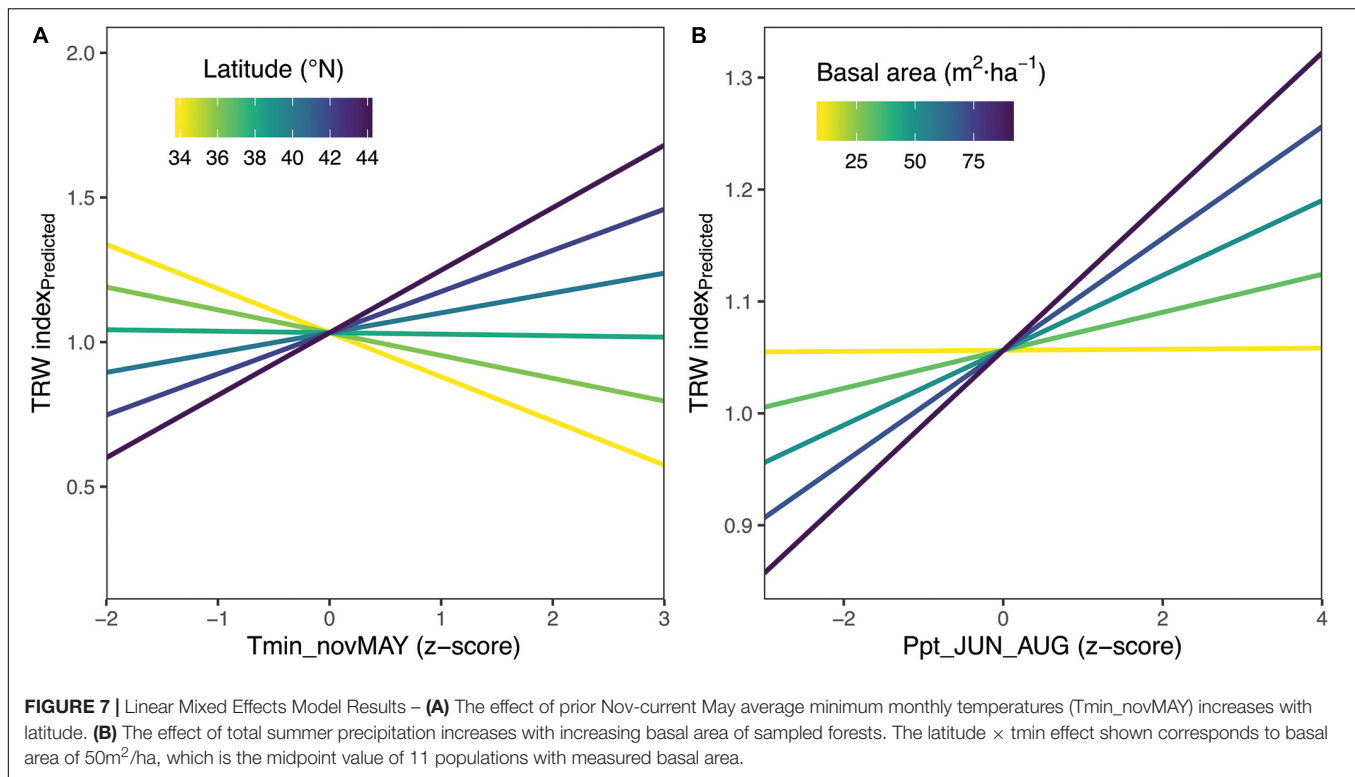
between regions and populations: total summer precipitation, average summer maximum monthly temperature, and average monthly minimum temperature from prior November through current May. Two of these climatic variables showed significant interactions with site-invariant variables such as basal area and latitude. Prior November through current May minimum monthly temperatures had a positive interaction with latitude ($p < 0.01$) so that its effect increased with latitude shifting from a small negative influence in the southern populations (latitude $< 36^\circ$) to a positive effect in the northern populations (**Figure 7**),



which is broadly consistent with other analyses. The growth of some southern trees and a few southern populations have some of the stronger negative relations to temperatures during the prior non-growing season (prior November through current May), especially prior November and current March and April (**Figures 4–6** and **Supplementary Figures S4, S5, S7, S8**).

The effect of summer precipitation was positively modified by stand density ($p < 0.01$), making tree growth more sensitive to precipitation with increasing stand density (**Figure 7**). In this case, total summer precipitation had a positive effect on

tree growth for the range of conditions considered except at the lowest stand densities where the effect was neutral. Two of the fixed effects, total summer precipitation and average summer maximum monthly temperature, were insignificant when only fixed effects were considered. However, allowing for varying slopes for different sites for the explanatory variables considered (i.e., our final model with fixed and random effects) showed that all of these variables were significantly related ($p < 0.05$) (**Table 2**). Fixed effects represent the mean response for *Chamaecyparis thyoides* along our entire gradient. Thus, the lack



of significance in the fixed effects of total summer precipitation and average summer maximum monthly temperature was due to the higher variability of population-level growth response along the gradient for those two effects as shown by the random effects (see next paragraph). Overall the model showed a marginal R^2 of 0.12 (considering fixed effects only) and a conditional R^2 of 0.47 (considering fixed and random effects). Residuals for this model showed no departure from the assumption of normally distributed error with homogenous variance.

The use of random effects in our model as intercept and slopes allows us to estimate the importance of the variability within a population and among trees. The standard deviation of the random effects can be compared with the parameter estimates of the fixed effects in order to estimate the variability of response between different levels of hierarchy, trees or populations. Standard deviation of the random effect for all three climate variables for this analysis at tree level (<0.0005) was much lower compared to standard deviation for the random effects for populations (>0.018) (Table 2). Comparing models with random effects for sites only or for sites and trees using a log-likelihood ratio chi-squared test showed no support for the inclusion of tree random effects. These results indicate a higher inter- than intra-population variability in climate response. In fact, standard deviations for random effect for total summer precipitation and average summer maximum monthly temperature were larger than the corresponding fixed effects in line with the large spread of correlation coefficients for all three regions combined (Figures 5, 6 and Supplementary Figures S6–S8).

The standard deviation for the random effect prior November through current May minimum monthly temperatures compared

to its fixed effect was lower than for the other two climate variables. Adding the interaction between prior November through current May minimum monthly temperatures and latitude reduces the variability associated with population response because part of that variability is related to latitude in the model. In contrast, the interaction between stand density and total summer precipitation seemed to only explain a small part of the variability in the response to total summer precipitation, suggesting that other environmental variables that we were unable to test (soil depth, stand topography, etc.) might be responsible for the different responses. The full final model can be found in **Supplementary Material**.

DISCUSSION

Synthesizing six decades of climate-growth dynamics in 18 *Chamaecyparis thyoides* populations across its contiguous distribution using our vulnerability framework allowed us to address two broad questions: (Q1) “How do *Chamaecyparis thyoides* populations respond to climate across its latitudinal range?” and, (Q2) “Does the synchrony in tree-level response to climate in *Chamaecyparis thyoides* change over space?” Regarding (Q1), we found that the growth of trees in northern populations was most limited by cooler temperatures during the non-growing season and early in the growing season, but that trees in the central populations of our study network had stronger positive correlations to precipitation and negative correlations to maximum temperatures during the growing season (Figures 4–6 and Supplementary Figures S3, S4).

TABLE 2 | Summary of the selected linear-mixed effects model of tree-ring width indices.

Fixed effects	Value	SE	DF	t-value	p-value	Random and fixed effects p-value
Intercept	1.0323	0.0403	21625	25.5890	<0.001	<0.0001
Ave Summer Maximum Temps	−0.0080	0.0073	21625	−1.0985	0.2720	<0.0001
Tot. Summer Precipitation	−0.0078	0.0165	21625	−0.4708	0.6378	<0.0001
Prior Nov-current May Minimum Temps	−1.4063	0.1710	21625	−8.2256	<0.001	0.0154
Tot. Summer Precipitation x Stand Density	0.0008	0.0003	21625	3.1098	0.0019	0.0317
Prior Nov-current May Mini Temps x Latitude	0.0369	0.0042	21625	8.7568	<0.001	<0.0001
Random effects						
	SD site	SD tree	Marginal R ²		R ² Conditional	
Intercept	0.13105	0.00021	0.12		0.47	
Ave Summer Maximum Temps	0.02006	0.00025				
Tot. Summer Precipitation	0.01801	0.00030				
Prior Nov-current May Minimum Temps	0.09293	0.00015				

Regarding (Q2), we found less climate-growth synchrony among populations in the southern end of our network. However, we also found that climate-growth synchrony varied with the climatic variable being tested and other environmental and ecological factors. Latitude appears to drive the climate growth synchrony of northern populations to non-growing and spring season temperatures. This was in contrast to the central populations had the greatest amount of climate growth synchrony in response to precipitation and maximum temperatures during the growing season (Figures 5, 6 and Supplementary Figures S6, S7), which together induce drought stress in plants. While we found greater vulnerability to future climatic change in some populations, we found levels of potential resilience within all populations.

The distribution of individual tree-growth responses to climate across our network provides evidence for a kind of ecological hysteresis. One definition of ecological hysteresis considers how external conditions can induce different changes in state and that these changes are conditional on the state of the system and direction of external change, that there is a linear response over time (Litzow and Hunsicker, 2016; Li et al., 2019). Another definition considers to how resistant or slow a system is to change than might be expected (Jesse, 1999; Loehle, 2000; van Nes and Scheffer, 2005). Yet, another definition of ecological hysteresis considers asymmetrical and opposing responses to the same climate factor (Da Silveira Lobo Sternberg, 2001; Zhang et al., 2003). Here we invoke the asymmetrical response of growth to warming across our study area as ecological hysteresis: most trees in the northern end of our network increase in growth in response to warming while a substantial number of trees in several populations in the southern end of our network would be non-responsive or even increase in growth with warming while some will decline in growth (Figures 5 and Supplementary Figures S7, S8). The asymmetrical response to temperature in the two ends of our network indicates the potential for greater growth stability within populations on the southern edge of *Chamaecyparis thyoides*' distribution than might be presumed under future warming (Loehle, 2000). Notably, our findings provide further evidence

that climate envelope modeling approaches are not nuanced enough to model populations at broad scales, at least for the 61-year period tested here, from 1940 to 2000. Generalization of our results suggests greater potential resilience or, at least, reduced negative impacts due to lagged or slower than expected response from projected warming, a hysteresis, than what might be expected for southern *Chamaecyparis thyoides* populations if climate envelope modeling held true.

How do *Chamaecyparis Thyoides* Populations Respond to Climate Across Its Latitudinal Range?

Our findings generally support hypothesis, *H(i)*, that growth in the majority of *Chamaecyparis thyoides* populations will increase with warmer winter and spring temperatures toward the northern end of our network. Growth of *Chamaecyparis thyoides* trees in northern populations, from ~41–44 degrees north latitude, increased with warmer winter and spring temperatures more than trees in the southern end of our network (Figures 4–7 and Supplementary Figures S4, S5, S7, S8). These findings not only support work with *Chamaecyparis thyoides* in our study region, they also support findings that growth of conifers nearer to the northern end of their distribution is more limited by non-growing season minimum temperatures than more southerly populations (Cook and Cole, 1991; Cook et al., 1998; Hopton and Pederson, 2005; Bhuta et al., 2009).

Cleaving the greatest difference between the response of trees to temperature within the contiguous distribution of *Chamaecyparis thyoides*, growth in the northern end of our network is significantly more limited by minimum temperatures than trees in the central and more southern regions during spring and from April–August. While this generally supports Hypothesis (i), the response of trees in the central portion of our network to winter temperatures is virtually indistinguishable from more northern trees (Figures 5, 6 and Supplementary Figures S7, S8), undercutting the strict geographic expectations of Hypothesis (i).

For Hypothesis (ii), we hypothesized that growth in the majority of southern *Chamaecyparis thyoides* populations will

increase more with more water availability during the growing season than populations in the northern end of our network. While this is a weak trend moving north to south, the strongest responses to drought are seen in the central portion of our network, which does not support this hypothesis. The implications of these findings are fully discussed below in see section “Implications and Mechanisms.”

Does the Synchrony in Tree-Level Response to Climate in *Chamaecyparis Thyoides* Change Over Space?

Our findings partially support *Hypothesis (iii)*, that the tree-level climate responses in more northern *Chamaecyparis thyoides* populations are more synchronous than more southern populations. Linear mixed effect models indicated a top-down constraint of minimum temperatures in the northern end of *Chamaecyparis thyoides*’ distribution (Figure 7), supporting findings of more synchronous responses to minimum temperatures at tree and population levels (Figures 5, 6 and Supplementary Figure S8). However, the modulation of the growth response to precipitation by geomorphology (ombrotrophic wetlands) and ecological structure (stand density) does not support *Hypothesis (iii)*. Stand density increased *Chamaecyparis thyoides* drought stress (Figure 7), providing evidence that water availability appears to have a bottom-up constraint on growth. The density and age structure of our study forests are indicative of the stem-exclusion stage of forest development. Stem-exclusion stage begins not too long after disturbance (within 2–3 decades) and is often characterized as having high stand density and low regeneration (Oliver and Larson, 1990). Our study forests dated back to either the time of intensive logging of *Chamaecyparis thyoides* ecosystems during the late 19th century (Laderman, 1989; Table 1) or in the more southern populations, recent forest management. Research on a larger network that includes other *Chamaecyparis thyoides* populations in the northern end of our network indicates that average population-level growth on ombrotrophic wetlands is mainly constrained by water availability (Pearl et al., 2019), a response we replicate with the Saco Heath population (Figures 5 and Supplementary Figures S3, S6). Trees growing on ombrotrophic wetlands gain water from incoming precipitation, not from groundwater, indicating how growth is constrained by water availability.

Our findings primarily support *Hypothesis (iv)*, that the responses of southern *Chamaecyparis thyoides* populations to all climatic variables and seasons will be less synchronous than more northern populations. The climatic responses of and within the southern populations were generally more muted and more variable than other populations (Figures 4–6 and Supplementary Figures S3–S8). These findings support prior work that indicted local factors as important factors of *Chamaecyparis thyoides* growth within this region (e.g., Eagle, 1999; Hopton et al., 2004; Hopton and Pederson, 2005; Mylecraine et al., 2005; Seim et al., 2006; DeBerry and Atkinson, 2014). However, we must also note that the more southern populations within our network tended to have more consistent

negative correlations to maximum temperatures, giving some support to the general expectation that southern populations in northern temperate regions will be vulnerable to rising temperatures in the future and indicating some potential synchronicity in the southern populations. We also must note that our analysis occurred to a period of broad cooling in the southern portion of our region (Figure 2; Mascioli et al., 2017). Hypothetically, hotter conditions earlier in the 20th century, ca 5000–10,000 years BP (Marcott et al., 2013), or the not too distant future might result in greater climatic growth synchrony than observed here.

Other Potential Factors of the Climate Response of *Chamaecyparis thyoides*

Our study considered some of the many potential factors influencing growth of *Chamaecyparis thyoides*. Only stand density was selected as an important factor on the climatic response of trees in this study. It is not clear why other factors were not selected. One reason could be that the accuracy of some of the environmental data is too coarse or similar variations between sites (e.g., distance to open water, slope, or aspect) to have a detectable impact on tree growth. It might also be that the range of environmental data we have for our 11 sites might be too small to capture differences between populations (e.g., similar distance to open water, slope, or aspect), which might limit the strength of the data. At the least, it seems we can say that overfitting is not a serious issue with the model results.

Some research suggests an age effect on the climate-growth relations in trees (Szeicz and MacDonald, 1994; Carrer and Urbinati, 2004; Martínez del Castillo et al., 2018). While there are differences in age structure between our regions, particularly with older trees in the central region (Table 1), linear effects modeling analyses suggests these differences do not strongly influence interannual to decadal scale growth, not more than the other environmental or ecological factors that we were able to test. It is possible that natural thinning during the early phases of ecosystem development removed individuals that represented a greater range of individual trees in the older central populations to reduce the overall range of growth response in this region. It is currently impossible to estimate how the trees that have died and are long faded from the landscape, so-called ghost trees (Foster et al., 2014), might have responded to climatic variations. It appears that age structure has a limited impact on climate-growth dynamics of *Chamaecyparis thyoides* in our network.

Implications and Mechanisms

Individuals in the southern portion of a species’ distribution are generally expected to be more vulnerable to a warming climate than individuals at higher latitudes in the distribution (e.g., Woodward and Woodward, 1987). In contrast, the stronger and more synchronous drought constraint on central *Chamaecyparis thyoides* trees during the warm season suggests these populations would be the most vulnerable to “hot droughts,” or periods characterized by both low precipitation and elevated temperatures. These so-called “hot droughts” typically increase drought stress in trees (Adams et al., 2017).

Increased warming is expected to lead to drier conditions via increased surface evapotranspiration (Cook et al., 2014; Mankin et al., 2019). Multiannual droughts can lead to growth declines (Berdanier and Clark, 2016) and increase tree mortality rates in our study region (Clark et al., 2016; Druckenbrod et al., 2019). Dead trees have been shown to decline in growth for years and sometimes decades prior to mortality (Pedersen, 1998; Cailleret et al., 2017). The physical and ecological implications suggest that the central *Chamaecyparis thyoides* trees would be more vulnerable with continued warming and if the current cooling trend during the growing season is reversed.

While our findings appear counter to theoretical understandings, they generally follow findings that core populations of *Fagus sylvatica* might be more vulnerable to climatic variation than edge populations (Cavin and Jump, 2017). In that study, it was thought that *Fagus sylvatica* living at the species' distributional margins had a greater resistance to drought due to local conditions. Our work provides some evidence to support that thesis: *Chamaecyparis thyoides* trees in forests with high stand density (basal area) are more likely to experience greater drought stress (Figure 7). Interestingly, trees in the central and southern regions of our network received a similar amount of precipitation during the period of analysis, but central region trees grew under summer conditions that were on average 4°C cooler than more southern trees (26.5°C vs. 30.5°C; Supplementary Table S1). That warming conditions during our study period did not synchronize the climate response of trees provides additional evidence that stand density can modulate the potential relief of a cooler climate. Our findings fall in line with the growing body of literature indicating that higher forest density can increase drought stress or vulnerability of trees to drought (Cescatti and Piutti, 1998; Martin-Benito et al., 2011; D'Amato et al., 2013; Bottero et al., 2017).

It might appear counterintuitive that *Chamaecyparis thyoides* trees growing in wetlands could be drought stressed, research has long shown that the annual growth of trees in mesic conditions or wetlands in mesic regions is often constrained by water availability (e.g., Hawley et al., 1941; Fritts, 1962; Stahle et al., 1988; Cook, 1991; Graumlich, 1993; D'Orangeville et al., 2018). As it appears now, the combination of future climatic projections and the ecology of drought stress in dense forests would seem to make dense *Chamaecyparis thyoides* populations more vulnerable to continued warming.

While the central populations appear vulnerable to summer warming, winter warming might also increase growth such that it could offset some of this potential negative impact on growth. Warm winter temperatures have been found to be associated with increased growth in the Northern Hemisphere for decades (e.g., Brubaker, 1980; Graumlich, 1993; Pederson et al., 2004; Babst et al., 2013; Martin-Benito et al., 2018; Weigel et al., 2018; Alexander et al., 2019; LeBlanc et al., accepted). Mechanisms that explain how dormant season temperatures might influence growth in conifers are, however, not an area of consensus. Warmer temperatures during the non-growing season can stimulate photosynthesis and carbon

assimilation in conifers (Dolman et al., 2002; Ensminger et al., 2004) while low temperatures can inhibit photosynthesis and delay spring recovery of winter-stressed trees (Ottander and Öquist, 1991; Ensminger et al., 2004). The distributions of winter temperature correlations in central populations are statistically indistinguishable from more northern populations, indicating the potential for increased growth with warming. Overall, the vulnerability of growth in central populations will depend on the balance of winter warming and increased water stress during periods of growing-season warming and reduced precipitation.

There is greater climatic synchrony to cool temperatures during the non-growing season and spring in the more northern portion of our network (Figures 5–7 and Supplementary Figures S7, S8), which would make them more vulnerable to an abrupt reversal of the current warming to an extreme cold climate. Though less likely to occur in the near future, paleoecological records indicate a reduction of southern temperate species and an increase of boreal species at the beginning of the Younger Dryas (Peteet et al., 1990; Peteet, 2000) occurred when the warming trend at the end of the Pleistocene abruptly reversed to extremely and persistent cold conditions (Broecker et al., 1988). Our findings indicate the growth of *Chamaecyparis thyoides* in the more northern end of our network would be more vulnerable to cooler winter and spring temperatures.

Advantages of the Vulnerability Framework and Examinations Across Scale

Investigating six decades of individual tree growth response of *Chamaecyparis thyoides* to climate in 18 populations over 11 degrees of latitude and ca 1500 km revealed the complexity of how individual trees respond to climate. Investigating the ecology of the individual is statistically more fraught, messy one might say, than distilling the response of individual trees to a population mean. Using a combination of population mean and individual tree approaches over the spatial scale of a network like ours has important advantages. Determining the average population response to climate allowed us to determine what seasons influence tree growth over a study region with differing seasonality. Comparing disparate populations using only one season would likely mask important ecological differences to trees and populations across our network. For example, while central populations appear more heat-stressed during the summer, population level analyses indicated a strong tendency of heat stress in the southern populations across most of the months tested.

By moving beyond the average population response and characterizing the range of growth responses within a population, we were able to discern which populations had trees that responded to growth more synchronously than other populations. This insight allowed us to identify populations that might be more vulnerable to extreme climatic shifts. Theoretically, climate, population vulnerability, and adaptation

influence the range of variability at the individual level more than the population level (Carrer, 2011; Clark et al., 2012). In fact, we found that understanding the levels of vulnerability and potential impact of continued climatic change depends on how bottom-up factors like stand density can modulate top-down factors like temperature within populations. Yet, surprisingly, we show how the climate-growth dynamics at the tree level reflects much of the broader patterns at the population and regional levels. In fact, linear mixed modeling indicated that climate-growth dynamics between populations at the average population level is greater than within populations. Despite this finding, we cannot help but think of the ecological nuance that would be extracted from large-scale studies of climate-growth dynamics had individual trees been analyzed instead of populations means (Martin-Benito and Pederson, 2015; Charney et al., 2016; Klesse et al., 2018a).

In thinking about vegetation modeling and our results, data assimilation between models, even Earth system models, and these kinds of studies could result in more nuanced understanding of climate constraints on tree growth. Our results show that, population-level responses of *Chamaecyparis thyoides* are more important than individual tree response. If this holds true for a larger number of species, considering the population level response in models may improve them enough so that individual tree responses might not be sufficiently justified given the trade-off between effort and gain. However, we also note that results for a species of restricted distribution, such as *Chamaecyparis thyoides*, limited to the specific sites might be hard to extrapolate to other systems. A good pursuit would be to continue efforts like ours and continued data-model assimilations (Dietze et al., 2013; Rollinson et al., 2017; Fisher et al., 2018).

Summary

By examining climate-growth responses at the individual tree level over the contiguous distribution of *Chamaecyparis thyoides*, we have shown that:

- (1) The central portion of the species' range has the most synchronous heat stress and drought response, which is counter to the common expectation that species are more responsive and vulnerable to climatic change at the southern end of their range in the Northern Hemisphere;
- (2) There is potential resilience to warming and drought within all populations;
- (3) Asymmetrical growth responses to projected warming across our study network provides evidence of ecological hysteresis;
- (4) Local environmental conditions can override climatic controls toward the southern end of its distribution;
- (5) Drought stress is an important constraint on growth, even for a species that primarily lives in wetland ecosystems;
- (6) Variability of the influence of climate on tree growth among populations dominated over the common species influence or individual tree effects, indicating that endogenous (e.g., genetics) or exogenous (site conditions) characteristics for each individual population need to be

considered for a thorough understanding of the species response to climate.

Researching growth-climate relationships at the individual tree level helps to understand how forests might respond to climatic change, and their potential productivity moving into the future (e.g., Carrer, 2011; Clark et al., 2012). However, some of the next steps in understanding the potential level of persistence and resilience of trees includes understanding how climate-growth relations might change over time (Cook and Zedaker, 1992; Visser et al., 2010; Carrer, 2011), the nonlinear response of tree growth to climate (Rollinson et al., 2017), and then linking these responses to individual genetics within a population (Housset et al., 2018). The ultimate growth vulnerability and potential mortality of tree populations likely depends on relative changes in climate, the genetic, phenotypic, and morphological adaptability of trees to change, and ecosystem development processes that influence ecological factors of growth. Overall, we find a complex structure of climate-tree growth responses within populations and over latitude that provides insight on the inherent difficulties in forecasting how climatic change will impact populations and forests over space.

DATA AVAILABILITY STATEMENT

The datasets for this study will be found in the Harvard Forest Archive upon publication <https://harvardforest.fas.harvard.edu/harvard-forest-data-archive>.

AUTHOR CONTRIBUTIONS

NP, DM-B, CL, TM, and DB designed the study. NP, DM-B, CL, DB, KA, and TM conducted data analyses and created data presentations. NP, MH-A, DB, and JP conducted field work and processed data. NP, DM-B, CL, JP, TM, and DB wrote the manuscript.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: <https://www.frontiersin.org/articles/10.3389/ffgc.2020.00039/full#supplementary-material>

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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