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Abstract: We remeasured two sets of permanent plots in old-growth, spruce–fir forests on Whiteface Mountain to quantify ongoing vegetation dynamics at sites impacted by spruce decline. One set of plots was a stratified random sample of the vegetation in a subalpine watershed (Baldwin site). The other was selected to represent forest conditions in a high-elevation subset of the spruce–fir forest (Esther site). Between 1987 and 1997, there was a significant increase in aboveground tree biomass at Baldwin with the majority of the increment due to the growth of canopy-sized trees. This growth occurred with little change in either species composition or size structure. The annual mortality rate of 1.28%·year⁻¹ for canopy-sized red spruce (*Picea rubens* Sarg.) in Baldwin almost matched the recruitment rate of 1.4 stems/ha per year. In addition, the relative growth rate of spruce was significantly faster than associated species. In contrast, spruce trees in Esther died at a rate of the 3.6%·year⁻¹ (1985–1995), and survivors grew more slowly than other species. The most obvious community-level trend at Esther (1985–2000) was an increase in overall tree density with most of this increase due to ingrowth of small trees. The demography of the spruce population at Baldwin suggests that the decline is over for at least this population.

Introduction

The decline of red spruce (*Picea rubens* Sarg.) trees in the northeastern United States is a well-documented example of the negative impact chronic atmospheric pollution, specifically acid deposition, can have on the terrestrial biota (Driscoll et al. 2001). Red spruce is a common tree in the upland forests in the northeastern United States. It occurs across a biome transition from eastern deciduous forest to conifer forest (Bormann et al. 1970). Beginning in the mid-1960s, many overstory red spruce trees in the region experienced a progressive loss of vigor. Symptoms included both crown dieback and reduced radial growth rates (Johnson and Siccama 1983). By the 1980s, as many as half the standing spruce trees were dead in some high-elevation stands (Siccama et al. 1982; Scott et al. 1984; Battles et al. 1992). The decline has been directly linked to freezing injury of spruce foliage (Friedland et al. 1984; Johnson et al. 1988; Peart et al. 1992b; Tobi et al. 1995). There is good evidence from both laboratory studies (DeHayes et al. 1999) and field trials (Vann et al. 1992) that exposure to acidic cloud water reduces the freezing tolerance of spruce leaves. In addition, chronic acid input has depleted calcium supplies (Likens et al. 1996) and mobilized additional aluminum (Shortle and...
Smith 1988) in northeastern U.S. forest soils. These changes in soil quality can adversely affect the growth of red spruce trees (Cronan and Grigal 1995). Also spruce–fir forests at high elevation are thought to be among the most sensitive ecosystems to nitrogen pollution. According to one scenario, the continuation of current levels of nitrogen deposition to these high-elevation forests could effect wholesale shifts in community composition with hardwood trees replacing the now-dominant conifer trees (McNulty et al. 1996; Fenn et al. 1998).

The regional trend in pollution shows a modest decline in one of the two primary constituents of acidic deposition. In the U.S. Northeast, the concentration of SO$_4^{2-}$ in precipitation has decreased significantly since 1970, while there has been no discernable change in the wet deposition of N (Likens et al. 2001; Driscoll et al. 2001). For example at the National Atmospheric Deposition Program’s site on Whiteface Mountain, New York (610 m elevation), SO$_4^{2-}$ wet deposition has decreased by almost 4%·year$^{-1}$ between 1985 and 2000 while for the same period there was no detectable trend in the rate of N deposition (data from National Atmospheric Deposition Program Web site). Coincident with these changes in acidic deposition, there is some indication that since the mid-1980s, red spruce at some sites may be recovering from the decline. Silver et al. (1991) reported large increases in dead spruce between 1982 and 1987 in their resurvey of stands on Whiteface Mountain, but noted an improvement in the crown condition of surviving trees. Several researchers have observed marked increases in spruce annual growth rates, as measured from increment cores, since the mid-1980s (Siccama et al. 1994; Reams and Van Deusen 1995).

In this paper, we provide an update of the status of the subalpine forest community and red spruce populations at Whiteface Mountain, a sentinel site for environmental monitoring in northern New York. Our fundamental premise is that species declines and the role of anthropogenic disturbances must be evaluated in the context of the prevailing vegetation dynamics. Thus, a comprehensive review of the effects of acidic deposition on trees requires recent trends in forest composition and tree demography to complement measures of pollutant emission and deposition rates. We derived this key information from re-measurements of a subset of the permanent sample plots established in the 1980s for the explicit purpose of monitoring the course of spruce decline. Within this framework, we examined two questions: (i) what is the forest-wide implication of a population level phenomenon, such as spruce decline, and (ii) are red spruce populations recovering from the recent episode of decline? We confined this initial investigation to two old-growth stands with a history of detailed ecological and biogeochemical studies. By doing so, we minimized the variability associated with differences in land-use history and capitalized on the wealth of associated data.

**Materials and methods**

**Study site**

Whiteface Mountain (44°22′N, 73°54′W) and its four subpeaks form an isolated massif at the northern end of Adirondack Mountains of New York. The entire massif lies within the boundaries of Adirondack Forest Preserve and the land is managed by the New York State Department of Environmental Conservation. The Adirondacks are in a region of cool, wet summers and cold, snowy winters. Mean growing season temperatures (mid-May to mid-September) on Whiteface Mountain range from 11.3°C at the summit (1483 m elevation) to 17.8°C at 603 m elevation (Battles et al. 1992). The past 15 years have been warmer than the previous 20 years. Mean monthly winter temperatures (November–February) at the observatory on Whiteface Mountain (603 m elevation) averaged 0.5°C warmer in 1984–1999 than in 1964–1983 (D. Wolfe, Atmospheric Science Research Center, State University of New York, Albany, N.Y., personal communication). Mean annual precipitation at 1000 m elevation is 156 cm, of which 30% falls as snow (Friedland and Miller 1999). Moderately deep to thin till covers the bedrock (primarily anorthosite). Soil type varies along the elevation gradient with a broad transition around 1150 m elevation between Spodosols below and Histosols above (Witty 1968).

**Data sources and background**

Between 1964 and 1966, one hundred and eighty-two forest stands on Whiteface Mountain were quantitatively sampled to examine vegetation–environment relations (Holway et al. 1969). These stands were not permanently marked, but their approximate locations are known from detailed field notes and maps. This original work documented that (i) the vegetation of the Whiteface massif is representative of the regional montane flora and (ii) like other mountains in the northeastern United States, the spatial distribution of vegetation is organized along the elevational complex gradient (Coggill and White 1991). Specifically at Whiteface Mountain, red spruce, balsam fir (Abies balsamea (L.) Mill.), and mountain paper birch (Betula papyrifera var. cordifolia (Regel.) Fern.) share dominance in the spruce–fir zone (800–1100 m in elevation). Yellow birch (Betula alleghaniensis Britt.) can be an important component at the lower end of the spruce–fir forest (800–900 m). Below 800 m, conifers are gradually replaced by northern hardwood species. Above 1100 m, balsam fir becomes increasingly important and forms almost pure stands between 1200 and 1350 m (Holway et al. 1969; Battles et al. 1992).

In 1982, Scott et al. (1984) relocated and resurveyed 32 of the subalpine stands at Whiteface Mountain and found a 40–70% decrease in the basal area of spruce with the greatest decrease occurring in the high-elevation (>900 m) stands. Siccama et al. (1982) had earlier reported similar decreases in spruce basal area for a comparable forest in Vermont. These two studies provided the best evidence of a region-wide decline in spruce populations (Pearl et al. 1992a).

In 1987, we completed a network of permanent sample plots in the subalpine forests (700–1350 m) on Whiteface Mountain as part of the U.S. Environmental Protection Agency (EPA) – USDA Forest Response Program (Battles et al. 1992). We placed the centre of a 21-transect grid at a randomly chosen point along the primary north–south ridge of
the Whiteface massif (Fig. 1). To track individual tree growth and population demography, 60 permanent plots (most 400 m² in area) were located along the transects in a stratified random manner. Strata were defined to ensure sampling red spruce populations at different elevations and aspects. All trees ≥5 cm diameter at breast height (DBH, breast height = 1.37 m) were tagged and identified. For tagged trees, DBH, canopy position, and vigor were recorded. Height and height to live crown were measured on a random subset of trees. Understory trees (trees ≥1 m tall and <5 cm DBH) were measured and counted along 2 m wide transects that centered on the diagonals of the plot (109 m²).

In 1997, we measured all the permanent sample plots (11) that were located in the watershed southwest of Baldwin Hill (Fig. 1). At the Baldwin site, there is no historical or ecological evidence that the spruce–fir forest has ever been logged or burned (Battles and Fahey 2000). The plots were normally distributed across the elevation gradient with a mean elevation of 943 m, near the midpoint of the elevation gradient that encompasses the spruce–fir forest (800–1100 m). The slope of the plots was 14 ± 7° (mean ± SD), and the Baldwin watershed was on the northwestern face of the mountain. In 1987, forty-three percent of the standing red spruce trees in the 11 remeasured plots were dead, and 15% of the forested area was affected by fine-scale, canopy gaps (Battles and Fahey 1996). In 1985, an Integrated Forest Study site was established near Esther Mountain on the northwestern slope of Whiteface Mountain (Fig. 1). The primary objective was to analyze the effects of atmospheric deposition on nutrient cycling in a variety of forest ecosystems (Johnson and Lindberg 1992). The Esther site was located in a high-elevation, old-growth, spruce–fir forest. Four 0.1-ha circular plots were set up to monitor vegetation influence on nutrient flux. In these plots we identified, measured, and tagged every woody stem ≥2 cm DBH. Plots were selected to represent the range of structural conditions observed in the spruce–fir forest (Friedland et al. 1991). These plots have been remeasured at least every 5 years since 1985. However, since 1993 only canopy-sized trees (DBH ≥9.5 cm) were tracked individually, while the smaller trees in the plots were subsampled in five radial transects (total sample area = 178 m²). The Esther plots are at a mean elevation of 1031 m with a narrow range from 1000 to 1063 m. The slope was 22 ± 4°. In 1985, forty-three percent of the standing red spruce trees in the Esther plots were dead.
Neither site was seriously impacted by the destructive, regional ice storm of 1998 (Irland 1998). The ice storm struck the northeastern United States and southeastern Canada in January 1998 and caused damage to 1.86 × 10^6 ha in New York. On Whiteface Mountain, icing was confined to elevations below 830 m and to forests on the east side of the masif (New York Department of Environmental Conservation Web site; D. Wolfe personal communication).

Data analysis

We used these three data sources (Scott stands, Baldwin plots, and Esther plots) to assess change in the old-growth spruce–fir forests at two sites on Whiteface Mountain. Results from the Scott stands serve as a baseline description of the predecline forest. We took care to summarize the composition in 1964–1966 for stands in the same general area as the Baldwin and Esther sites. Following the ecological sampling standards of the time, Holway and Scott (1969) measured only stands without recent major disturbances. Thus, the 1964–1966 results represent only the mature phase of forest development.

From the repeated measures in the Esther and Baldwin plots, we calculated annual mortality, recruitment, and growth rates. We followed the methods in Sheil and May (1996) for evaluating mortality and recruitment and took care to avoid census-period dependence by making comparisons only over census intervals of the same length. Specifically annual mortality (λ, %·year⁻¹) and recruitment (k, individuals/ha per year) were computed as follows:

\[ \lambda = \frac{\ln(N_0) - \ln(N_t)}{t} \]

\[ k = -\frac{N^*}{At} \left[ 1 - \frac{N^*_t(t)}{N^*} \right] \]

where \( N_t \) is the number of live individuals at the end of the census interval, \( N_0 \) is the number of live individuals at the beginning of the census, \( t \) is the length of the census interval, \( A \) is the area sampled, and \( N^*_t(t) \) is the total number of new stems (recruits) present at the end of the census. To account for those individuals that are newly recruited but die before they can be recorded at the end of the census interval, we assumed that the 1964–1966 estimates of tree density were the asymptotic steady-state values to which the periodic recruitment rates tend (\( N^* \) in eq. 2). While any assumption of near steady-state dynamics is suspect for a forest recovering from a decline–disturbance (Battles and Fahey 1996), in practice our calculated recruitment rates were not sensitive to large changes in \( N^* \). For example, a 50% decrease in \( N^* \) for subcanopy spruce trees in Baldwin only raised the estimated recruitment rate by 12%; doubling \( N^* \) only decreased the recruitment estimate by 4%.

We expressed individual tree growth as stem volume increment. To compare the growth rate among species and populations with different size structures, we needed a metric that accounted for the allometric scaling between plant growth and plant size (Enquist et al. 1999; Niklas and Enquist 2001). Based on inspection of the data, we chose to measure growth as the slope of the regression line describing stem volume increment as a linear function of initial stem volume. All regression were forced through zero to preserve biological reality (i.e., trees with no volume do not grow). Specifically, for each species at each site we fit the following equation using a least-squares procedure:

\[ V_t - V_0 = RGI(V_0) \]

where \( RGI \) is the relative growth increment (m^3·m^-3), \( V_t \) is the stem volume at the end of the census interval, and \( V_0 \) is the stem volume at the beginning of the census interval. In all cases, the census interval was 10 years. For all regressions, there was a significant linear relationship between stem volume increment and initial stem volume \((p < 0.01)\). The fits ranged from a high of \( R^2 = 0.65 \) for balsam fir at Esther to a low of \( R^2 = 0.29 \) for red spruce at Esther. Stem volume was computed using a parabolic approximation: one-half the stem cross-sectional area at breast height times the tree height (Whittaker et al. 1974). Tree heights were estimated using site-specific allometric equations that predict height as a function of species, DBH, and elevation (Battles et al. 1995).

Changes in aboveground tree biomass were assessed using the standard methodology described in Whittaker et al. (1974). For canopy-sized trees (trees ≥9.5 cm DBH), we estimated parabolic volume as described above and then used volume as the independent variable to predict tree biomass from the allometric equations in Whittaker et al. (1974) as updated by Siccama et al. (1994). This approach is the same as that applied by Friedland et al. (1991) and Friedland and Miller (1999) for biogeochemical analyses at the Esther site. However, we used separate species-specific equations for trees smaller than 9.5 cm DBH to account for ontogenetic change in tree allometry. A suite of equations for the dominant species at Whiteface Mountain were available from work done in the White Mountains of New Hampshire (Fahey et al. 1998; Reiners 1992; T. Siccama, unpublished data).

Statistical analysis

To assess changes in forest composition and tree biomass, we used the appropriate repeated-measures design where the subjects (in this case, the plots) serve as their own controls (Neter et al. 1990). Specifically for Baldwin, we used paired \( t \) tests to detect significant changes between 1987 and 1997. For Esther, we used general linear models with plots as a random factor to detect significant trends between 1985, 1990, 1995, and 2000. Changes in diameter distributions were tested with Kolmogorov–Smirnov tests; changes in median height of canopy-sized trees with Mann–Whitney U tests. Significant differences in RGI were defined by the nonoverlap of 95% confidence intervals based on the \( t \) distribution. While we applied similar statistical analyses to the Baldwin and Esther data, it is important to note that the Baldwin plots represent a stratified random sample of the spruce–fir forest in the Baldwin watershed. Thus, the results from the Baldwin plots are representative of the entire watershed. In contrast, the Esther plots were selected to capture the range of conditions observed, and thus, inferences cannot be extrapolated beyond what is happening to the trees in these plots.

In some cases, mortality rates were based on very small sample sizes. We assessed the reliability of these estimates...
with randomization tests based on binomial probabilities (sensu Condit et al. 1995). For 99 random samples, we developed the distribution of expected mortalities using the observed sample size and the observed ratio of surviving trees. We then took the 2.5 and 97.5% quantiles as the bounds for the 95% confidence interval for the observed mortality rate. Since the trees from the Esther plots are not intended to be population-level estimates, we did not calculate empirical confidence intervals for Esther mortality rates.

### Results

#### Community dynamics

In the Baldwin watershed, there was no change in canopy tree (DBH ≥9.5 cm) density between 1987 and 1997. However, canopy tree basal area increased by 13% during the same period (Table 1; paired \(t\) test, \(p < 0.01\)). Paper birch experienced the largest relative increase (24%) but the basal area of all three dominant species increased by at least 15% (Table 1). Yellow birch was the only canopy tree that declined in relative basal area during the decade. Despite this basal area increase, there were only minor changes in canopy composition. Balsam fir was the dominant species in 1987; its basal area was more than double that of spruce and paper birch. The same allocation of species dominance was true in 1997 (Table 1). The composition of the subcanopy trees was remarkably stable between 1987 and 1997 (Table 2). Also, neither the total density nor the composition of the smallest trees (trees <5 cm DBH and ≤9.5 cm DBH) more than doubled in the Esther plots between 1985 and 2000. Mean understory density increased from 2131 ± 714 (mean ± SE) to 4418 ± 908 stems/ha; basal area went from 3.5 ± 1.0 to 8.8 ± 2.6 m²·ha⁻¹. Although more than 95% of the understory trees were fir, spruce and paper birch either maintained (spruce) or increased (birch) their relative abundance in the understory size class between 1985 and 2000. This general rise in understory trees contributed to net increases in basal area, density, and aboveground tree biomass in the Esther plots (Table 4). However, only the 81% increase in density was statistically detectable.

From 1987 to 1997, tree basal area and aboveground tree biomass in the Baldwin plots increased significantly with only a minor concomitant change in tree density (Table 4, Fig. 2). Also, the size distribution of trees in the Baldwin plots (based on DBH) were nearly identical between 1987 and 1997 (Fig. 2; Kolmogorov–Smirnov, \(p = 1.0\)). The same was true for the diameter distributions at Esther between 1985 and 2000 (Fig. 2; Kolmogorov–Smirnov, \(p = 0.96\)), although the numerical gain in total tree density at Esther was apparent.

#### Population dynamics

For canopy red spruce trees in Baldwin during the 10-year census interval (1987–1997), the annual mortality rate was 1.2%·year⁻¹, and annual recruitment rate was 1.4 stems/ha per year (Table 5). The rates nearly offset each other with the net result being a small (4%) reduction in spruce canopy tree density (Table 1). Based on the RGI, spruce grew significantly faster than associated species (Table 5). This rapid growth accounted for most of the 15% increase in spruce basal area (Table 1). For both balsam fir and paper birch, canopy recruitment exceeded mortality. Thus, a combination of recruitment and growth led to the observed increases in fir and birch basal area (Tables 1 and 5). With the exception

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Table 1. Changes in canopy tree composition at the Baldwin site, an old-growth, spruce–fir forest on Whiteface Mountain, New York.

<table>
<thead>
<tr>
<th>Tree species</th>
<th>Density (stems/ha)</th>
<th>Basal area (m²·ha⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Red spruce</td>
<td>253</td>
<td>154</td>
</tr>
<tr>
<td>Balsam fir</td>
<td>521</td>
<td>612</td>
</tr>
<tr>
<td>Paper birch</td>
<td>158</td>
<td>193</td>
</tr>
<tr>
<td>Yellow birch</td>
<td>10</td>
<td>64</td>
</tr>
<tr>
<td>Total</td>
<td>976</td>
<td>1068 (126)</td>
</tr>
</tbody>
</table>

Note: Canopy trees defined as stems ≥9.5 cm DBH. The 1964–1966 data were from seven Scott stands in or near the Baldwin site. Values in parentheses are SEs (\(N = 11\) plots between 800 and 1100 m elevation). Other tree species present and included in the total but not listed separately were *Acer saccharum* Marsh., *A. pensylvanicum* L., *A. spicatum* Lam., *Sorbus americana* Marsh., and *Pennisetum pennsylvanica* L.

Table 2. Changes in subcanopy tree composition at the Baldwin site, an old-growth, spruce–fir forest on Whiteface Mountain, New York.

<table>
<thead>
<tr>
<th>Tree species</th>
<th>Density (stems/ha)</th>
<th>Basal area (m²·ha⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Red spruce</td>
<td>100</td>
<td>109</td>
</tr>
<tr>
<td>Balsam fir</td>
<td>736</td>
<td>742</td>
</tr>
<tr>
<td>Paper birch</td>
<td>171</td>
<td>126</td>
</tr>
<tr>
<td>Yellow birch</td>
<td>44</td>
<td>29</td>
</tr>
<tr>
<td>Total</td>
<td>1092 (246)</td>
<td>1028 (172)</td>
</tr>
</tbody>
</table>

Note: Subcanopy trees are 5 cm DBH < 9.5 cm. Values in parentheses are SEs (\(N = 11\) plots between 800 and 1100 m elevation). Other tree species present and included in the total but not listed separately were *A. saccharum*, *A. pensylvanicum*, *A. spicatum*, and *S. americana*.
of yellow birch, mortality and recruitment rates of the sub-
canopy trees in Baldwin were higher than canopy trees (Ta-
ble 6). However, there was little change in the composition
of this stratum of the tree community between 1987 and 1997 (Table 2).

In contrast to Baldwin, annual mortality of red spruce
canopy trees in the Esther plots consistently outpaced can-
oopy recruitment (Table 7). For a comparable period (1985–
1995), spruce mortality in Esther was more than double and
recruitment less than 20% of the rates observed in Baldwin
(Tables 5 and 7). On the other hand, while balsam fir experi-
cenced relatively high annual mortality (2.0%·year –1 between
1985 and 1995), it had even higher rates of recruitment
(24 stems/ha per year between 1985 and 1995). Balsam fir
also did not find any evidence of collateral damage to bal-
sam fir or paper birch in the wake of widespread spruce
mortality (Perkins et al. 1992; McLaughlin and Percy 1999).
However, there were important changes in the rate and na-
ture of biomass accumulation since the mid-1980s. Between
1987 and 1997, the spruce–fir forest in the Baldwin water-
shed was an aggrading ecosystem in terms of aboveground
tree biomass with the majority of the increment due to the
growth of canopy-sized trees. This growth occurred with
only minor shifts in composition or structure (Tables 1 and
4, Fig. 2). Trends in the Esther plots between 1985 and 2000 in-
cluded an increase in overall tree density (Fig. 2, Table 4)
with most of this increase occurring in the understory size
class (9.5 cm > DBH ≥ 2 cm). There was also an increase in
the importance of fir relative to spruce in the canopy (Ta-
ble 3). Friedland and Miller (1999) noted the importance of
t fir component in terms of nitrogen biogeochemistry. Nitro-
gen concentration in balsam fir foliage was 40% higher than
red spruce. Although there was only a small increase in tree
biomass between 1985 and 1995 (3%), there was a 27% in-
crease in N stored in vegetation (Friedland and Miller 1999).
Since 1995, the trees in the Esther plots have continued to
accumulate biomass by the same means, namely an ingrowth
of small fir trees.

The 0.5-m reduction in the stature of the Esther canopy
could influence population, community, and ecosystem pro-
cesses at the site. For example, shorter canopy trees experi-
ence a reduced risk of mortality from both chronic and
catastrophic winds (Peltola et al. 1999; Canham et al. 2001).
Miller and Friedland (1999) documented the downward shift

### Table 3. Trends in canopy tree composition at the Esther site, a high-elevation, spruce–fir forest on Whiteface Mountain, New York.

<table>
<thead>
<tr>
<th>Tree species</th>
<th>Density (stems/ha)</th>
<th>Basal area (m²·ha⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Red spruce</td>
<td>356</td>
<td>81</td>
</tr>
<tr>
<td>Balsam fir</td>
<td>886</td>
<td>593</td>
</tr>
<tr>
<td>Paper birch</td>
<td>200</td>
<td>57</td>
</tr>
<tr>
<td>Total</td>
<td>1475</td>
<td>746</td>
</tr>
</tbody>
</table>

Note: Canopy trees defined as stems ≥ 9.5 cm DBH. The 1964–1966 data are from seven Scott stands in or near the Esther site. Values in parentheses are SEs (N = 4 plots between 1000 and 1063 m elevation). Sorbus americana was a tree species present and included in the total but not listed separately in the table.

### Table 4. Changes in forest community structure in old-growth spruce–fir sites on Whiteface Mountain, New York: (A) Baldwin and (B) Esther Mountain sites.

#### (A) Baldwin site (trees ≥ 5 cm DBH).

<table>
<thead>
<tr>
<th>Parameter</th>
<th>1987</th>
<th>2000</th>
<th>Change (%)</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Basal area (m²·ha⁻¹)</td>
<td>31.0</td>
<td>34.2</td>
<td>10</td>
<td>0.02</td>
</tr>
<tr>
<td>Density (stems/ha)</td>
<td>2160</td>
<td>2123</td>
<td>−2</td>
<td>0.85</td>
</tr>
<tr>
<td>Aboveground tree biomass (Mg·ha⁻¹)</td>
<td>124</td>
<td>138</td>
<td>11</td>
<td>0.02</td>
</tr>
</tbody>
</table>

#### (B) Esther site (trees ≥ 2 cm DBH).

<table>
<thead>
<tr>
<th>Parameter</th>
<th>1985</th>
<th>2000</th>
<th>Change (%)</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Basal area (m²·ha⁻¹)</td>
<td>30.3</td>
<td>35.6</td>
<td>17</td>
<td>0.08</td>
</tr>
<tr>
<td>Density (stems/ha)</td>
<td>2877</td>
<td>5328</td>
<td>81</td>
<td>0.03</td>
</tr>
<tr>
<td>Aboveground tree biomass (Mg·ha⁻¹)</td>
<td>107</td>
<td>115</td>
<td>8</td>
<td>0.28</td>
</tr>
</tbody>
</table>

Note: Statistical significance was evaluated using paired t tests for each structural parameter.

### Discussion

#### Forestwide implications of spruce decline

The recent community-level changes in these two spruce–
fir forests suggest that spruce decline did not initiate whole-
sale transformations of tree composition and structure. We
also did not find any evidence of collateral damage to bal-
sam fir or paper birch in the wake of widespread spruce
mortality (Perkins et al. 1992; McLaughlin and Percy 1999).
However, there were important changes in the rate and na-
ture of biomass accumulation since the mid-1980s. Between
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biomass between 1985 and 1995 (3%), there was a 27% in-
crease in N stored in vegetation (Friedland and Miller 1999).
Since 1995, the trees in the Esther plots have continued to
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of small fir trees.

The 0.5-m reduction in the stature of the Esther canopy
could influence population, community, and ecosystem pro-
cesses at the site. For example, shorter canopy trees experi-
ence a reduced risk of mortality from both chronic and
catastrophic winds (Peltola et al. 1999; Canham et al. 2001).
Miller and Friedland (1999) documented the downward shift
in the vertical distribution of leaf area associated with the changes in canopy structure in the Esther plots. In terms of the forest community, the amount of foliage in the layers available to vertebrate browsers (0–2 m in height) doubled from 0.6 to 1.2 m²·m⁻² between 1985 and 1995. At the ecosystem level, any alterations in the composition and distribution of leaf area has the potential to modify the atmospheric deposition regime (Miller and Friedland 1999).

Assessing recovery from decline

Standards of proof sufficient to document the existence of a forest decline were much debated with reference to red spruce trees (e.g., Loehle 1988). Eventually a consensus was reached that relied on results from repeated censuses of relocated vegetation plots (Peart et al. 1992a). There also needs to be criteria to determine the end of a decline event and to assess the potential for recovery. Thus, we suggest that an
end to the decline would require both a cessation of high mortality and a return to vigorous growth. A recovering population would need to exhibit canopy recruitment that exceeds or at least matches canopy mortality, and a regeneration rate sufficient to maintain its future presence in the community. Previously, mortality had been approximated by changes in basal area (Siccama et al. 1982; Scott et al. 1984) or percent standing dead trees (Silver et al. 1991; Battles et al. 1992). However, to get the threefold increase in standing dead spruce observed by Silver et al. (1991) from 1982 to 1987 at their Whiteface site, spruce mortality would have had to be at least 5%·year\(^{-1}\). The current mortality rate at Baldwin is comparable with the mortality rates reported for a stable red spruce population (i.e., nondeclining) in an old-growth forest in the southern Appalachians (Busing and Pauley 1994). Growth rates in the Baldwin population have also rallied to the point where red spruce was the fastest growing species in the canopy during the decade 1987–1997 (Table 5).

The case for recovery of the Baldwin spruce population is less certain. Canopy recruitment matched spruce mortality (Table 5). Overall spruce basal area increased because of the robust growth of surviving trees (Table 1). For subcanopy spruce (5 cm ≤ DBH < 9.5 cm), a relatively high mortality rate (3.6%·year\(^{-1}\)) was offset by equally high recruitment (9.6 stems/ha per year; Table 6) with the result being a small net change in subcanopy density (Table 2). The smallest size class of spruce trees (trees ≥ 1 m tall and ≤ 5 cm DBH) remained at approximately 700 stems/ha during the sampling interval. The demographics suggest that recruitment and regeneration rates are sufficient to replace the current generation of canopy trees, but there is yet no indication that red spruce is regaining its former status in the community. Cause for concern includes the relatively high mortality of spruce in the subcanopy size class. Given the differences in the autecology of the two conifers, spruce is expected to have lower mortality than fir in the smaller size classes (White and Cogbill 1992), yet the opposite was true for the subcanopy trees at Baldwin (Table 6).

Compared with the Baldwin population, spruce at Esther are dying faster, growing less, and recruiting more slowly (Tables 1, 3, 5, and 7). In their analysis of element cycling at the Esther site, Friedland and Miller (1999) reported a significant decrease in red spruce basal area between 1985 and 1995. Indeed, annual spruce mortality during the interval was 3.6%·year\(^{-1}\), and spruce basal area decreased by 38% (Tables 3 and 7). The most recent 5-year estimate of mortality (0.9%·year\(^{-1}\), 1995–2000) implies a return to a more sustainable level of mortality, but this apparent reduction may be an artefact of the census-period dependence in vital rates noted by Sheil and May (1996). The evidence from the Esther plots indicates that spruce decline is not yet over.

It is important to note that the trees in the Esther plots represent a high-elevation subset of the population. In the montane forest of the U.S. Northeast, the more demanding environment encountered at higher elevations is associated with lower growth and higher mortality of the dominant tree species (Siccama 1974; Foster and Reiners 1983; Battles et al. 1992). In addition, there was a well-documented elevation gradient in the severity of spruce decline (Craig and Friedland 1991). Specifically at Whiteface Mountain, there was a significantly higher fraction of standing dead spruce above 1000 m in elevation (Battles et al. 1992). Also, Miller and Friedland (1999) could not detect any significant decrease in either S or N deposition rates between 1986 and 1996 at the Esther site, a result in contrast to the regionwide trend and the trend from the NADP site at a lower elevation on Whiteface Mountain. The comparison suggests that given the documented importance of the elevational gradient in structuring the vegetation of these mountain ecosystems (Bormann et al. 1970; White and Cogbill 1992), there most likely will be differences in the nature and timing of the forest recovery from spruce decline.

### Future implications

Compared with the composition of the predecline forest (1964–1966), the contemporary forest at Baldwin and Esther had fewer spruce and a greater relative dominance of balsam fir (Tables 1 and 3). We recognize that values based on the Scott stands are biased toward larger basal area, lower tree density, and greater spruce dominance given the explicit exclusion of disturbed patches from their sample (Holway et al. 1969). Yet, these historical estimates do set an upper bound for the trajectory of any recovering stands. As of yet, none of the 15 contemporary plots were approaching the “mean” composition of the spruce–fir forest in 1964–1966.

For canopy red spruce trees at Whiteface Mountain, visual assessment of crown condition (Johnson and Siccama 1983; Silver et al. 1991; Peart et al. 1992) has proven to be a reliable indicator of tree vigor. For the Baldwin plots, less than 5% of the spruce trees in 1987 and 1997 were in the severe crown damage class (>50% of crown foliage missing or discolored); for the Esther plots, more than 25% of the trees had crowns with severe damage in 1985 and 1995. The difference between sites in the proportion of trees with unhealthy crowns was reflected in the corresponding difference in spruce growth rates (Tables 5 and 7).

Looking toward the future, the return of these red spruce populations to their predecline dominance is uncertain. Weather plays a fundamental role in spruce decline. It modulates the input of the predisposing stressor (i.e., acidic de-
position) and determines the severity of a key inciting factor (i.e., midwinter temperatures cold enough to induce freezing injury to foliage). Because of the unpredictability of the weather component of this decline, current trends that suggest an incipient recovery could be reversed by a series of cold winters.

Red spruce decline is a regional phenomenon with a complex etiology (Johnson et al. 1992). To measure and to understand any recovery will require a regionwide evaluation. Here, we present the first estimates of mortality and recruitment for the subalpine forest in the U.S. Northeast. However, many more communities and populations need to be sampled to test the spatial and temporal consistency of the trends observed for two sites on Whiteface Mountain.

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References


Table 7. Demography of canopy trees (≥29.5 cm DBH) at the Esther site, a high-elevation, spruce–fir on Whiteface Mountain, New York.

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<td>( N_0 )</td>
<td>( \lambda )</td>
<td>( k )</td>
<td>( N_0 )</td>
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<tr>
<td>Red spruce</td>
<td>30</td>
<td>4.5</td>
<td>0</td>
<td>24</td>
</tr>
<tr>
<td>Balsam fir</td>
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<td>2.5</td>
<td>17.6</td>
<td>266</td>
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<td>Paper birch</td>
<td>21</td>
<td>0.5</td>
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Note: Results are from measurements of four permanent sample plots between 1000 and 1063 m elevation. Canopy recruitment includes correction for recruits that could have entered canopy class after 1987 but died before remeasurement in 1997 (after Sheil and May 1996). Values in parentheses are one-half the 95% confidence interval of \( RGI \), annual mortality (%·year –1); \( k \), recruitment (stems/ha per year); \( N_0 \), number of living trees at the beginning of the census interval; \( RGI \), per-volume growth increment during the interval (m 3·m–3).


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