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Genetic Variation and Climatic Impacts on Survival and Growth of White Spruce in Alberta, Canada

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Abstract

Because climate has the greatest effect in determining the genetic structure of forest tree species, climatic variables with large effects on growth and survival need to be identified. This would enable proper matching of tree populations to planting sites in the present and future climates. We analysed 24-year survival (S24), height (H24) and diameter (D24) from a series of white spruce provenance trials with 46 populations and 8 test sites in Alberta, Canada. We determined: (1) the amount and pattern of genetic variation, (2) the response of populations to climatic transfer and (3) the potential effects of climate change (2030–2039) on H24 and S24 of the

species in Alberta. We found that: (1) using the intraclass correlation, the between-population genetic variance was 10.6% (H24) and 6.6% (D24) of the betweenpopulation phenotypic variance across sites, (2) three climatic white spruce regions exist in Alberta within which variation in growth potential is strongly clinal, (3) the annual moisture index (AMI) expressed as a ratio of degree days above 5°C (GDD) and mean annual precipitation (MAP) was the major determinant of survival and growth at the test sites, (4) we found that at the level of AMI predicted for the 2030-2039 period, survival and growth would decline substantially in the continental part (northern and central) of Alberta where drought already exists. However, during the same period, survival and growth would increase substantially in the foothills and Rocky Mountains region where growth is currently limited by low GDD due to a short growing season.

Key words: climate change, moisture index, Picea glauca, provenance trial, response function.

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Introduction

Climate is a major determinant of the spatial genetic differentiation of forest tree species. Studies have shown that in the Northern Hemisphere, growth potential declines northward (e.g., LANGLET, 1959; KRUTZSCH, 1992; RUDOLPH and YEATMAN, 1982; XIE and YING, 1995) and from low to high elevation (REHFELDT, 1989; KLEN-SCHMIT and BASTIEN, 1992; BONGARTEN and HANOVER, 1986) partly in response to variation in temperature (MATYAS and YEATMAN, 1992). In addition, provenance trials have shown that climate determine survival and growth when tree populations are planted on a range of field environments (e.g., MATYAS and YEATMAN, 1992; Rehfeldt et al., 1999, 2001, 2002). Therefore, the knowledge of the climatic factors affecting the genetic differentiation of tree species is essential for matching populations to planting sites and adapting to the expected climate change.

In provenance trials, tree populations are planted on a range of sites with different climates and ecology. Consequently, these trials provide the best material for simulating climate change and its effects on survival and growth of individual populations (MATYAS, 1994, 1996; STETTLER and Bradshaw, 1994). Provenance trials have been used in jack pine (Pinus banksiana Lamb.) (MATYAS and YEATMAN, 1992), lodgepole pine (Pinus contorta Dougl.) (REHFELDT et al., 1999, 2001), Scotch pine (Pinus sylvestris L.) (REHFELDT et al., 2002), white spruce (Picea glauca (Moench.) Voss) (ANDALO et al., 2005), Loblolly pine (Pinus taeda L.) and Norway spruce (Picea abies (L.) Kharst) (Schmidtling, 1994) to model the response of species and populations to climatic transfer and predict their survival and growth in a changed climate.

Recent research results have shown that tree populations differ in their climatic optima and that in a changing climate, populations and regions of the species geographic ranges could be affected differently (e. g., Rehfeldt et al., 1999, 2001, 2002). In the present work, we examine the level of population differentiation of white spruce in Alberta, the role of climate in this population structure, the population-specific response to climatic transfer and how the observed level of population climatic sensitivity could affect survival and growth of white spruce in a changing climate.

White spruce is widely distributed across Canada, northeastern USA and Alaska (NIENSTAEDT and ZASADA, 1990). It occurs throughout Alberta, except in the prairies, subalpine and alpine regions. In the foothills and Rocky Mountains of western Alberta, white spruce generally occurs below 1370 metres in the south (OGILVE and VON RUDOLFF, 1968) and 1220 metres in the north (ACHUFF and LA ROI, 1977). This boundary is variable due to the heterogeneity of foothills environments and a distribution related to climate and clinal introgression with Engelmann spruce (Picea engelmannii Parry ex Engelm.), which occurs at higher elevations (LA RoI and DUGLE, 1968; ACHUFF and LA ROI, 1977; RAJORA and DANCIK, 2000). In the Rocky Mountains, the upper boundary of white spruce and lower boundary of Engelmann spruce are intertwined. As a result, both pure white spruce and hybrid populations have been identified at elevations as high as 1450 and 1524 metres (LA ROI and DUGLE, 1968; RAJORA and DANCIK, 2000). In southern Alberta, the limit of white spruce distribution occurs at the woodland – prairie transition where drought is limiting to tree distribution, mean annual precipitation is low (380–510 millimetres) and mean July temperature maxima are greater than 24°C (NIENSTAEDT and ZASADA, 1990).

Although white spruce is the most important forest tree species in Alberta comprising of 45.9% by volume of the coniferous species and 27.9% of coniferous and deciduous species combined (AFLW, 1985), its genetic variation has been studied extensively only in eastern Canada and northeastern USA. Therefore, this study addresses the following questions: (1) How much population differentiation is there for white spruce in Alberta? (2) To what extent is population differentiation of white spruce in Alberta determined by regional climate? (3) What planting site climate variables affect survival and growth of white spruce in Alberta? (4) What are the potential impacts of the expected climate change on future survival and growth of white spruce in Alberta? Throughout the article, the term population refers to provenance or location where seeds were collected.

Materials and Methods

Experimental Procedures

Between 1959 and 1976, the Alberta Forest Service collected white spruce seedlots from 46 natural stands distributed across Alberta. These bulk collections were made primarily for operational reforestation and sampled a large number of trees. At the beginning of the Alberta tree improvement programme in 1975, the archived seedlots from these collections were used to initiate a series of provenance trials across Alberta. The populations covered in this article are described in *Table 1* and *Figure 1*. All populations from the sympatric zone of white and Engelmann spruces were treated as white spruce.

Field trials were established with 4-year old seedlings between 1980 and 1983 at 12 sites, each containing 26–30 populations. These included a set of 21 populations that were planted at all 12 sites and 5–9 other populations that were selectively allocated to specific test sites as local controls. Four of the test sites exhibited poor survival and could not be continued beyond age 15 years. Therefore, the present article covers performance on 8 sites that exist today. These sites are described in *Table 1* and their approximate locations in Alberta appear in *Figure 1*.

The field layout was a randomised complete block design with 5 replications and 9-tree row plots at 2.5 x 2.5 metre spacing. A border row was planted around each trial to eliminate the edge effect. Trees were maintained free-to-grow by hand weeding and brushing at one to three-year intervals. The present article covers survival (S24), total height (H24) and diameter at breast height (D24) at the age of 24 years.

Climate Data and Choice of Variables

The 1961–1990 climate data for population and test site locations were derived from the Alberta Climate

Table 1. – Locations and climatic (1961–1990) description for populations and test sites in the Alberta white spruce provenance trials.

	1 1											
ID	Location	LAT.	LON.	ELE.	MAT		NDD	TWM	GDD	CI		AMI
_		(°N)	(°W)	(m)	(°C)	(°C)		(°C)		(°C)	(mm)	
	<u>dations</u>	#00444				21.0	2660				400	• •
2+	Footner Lake	58°44'	117°15'	335	-1.6	-21.9	2668	16.0	1190	37.9	408	2.9
3	Footner Lake	57°55'	117°55'	305	-0.3	-20.8	2371	16.4	1303	37.2	401	3.2
4+	Footner Lake	57°55'	115°30'	360	-0.4	-20.9	2353	16.3	1275	37.2	428	3.0
5	Footner Lake	58°33'	114°14'	235	-1.0	-21.9	2502	16.3	1239	38.2	370	3.3
6+	Athabasca	58°44'	111°15'	235	-1.7	-23.9	2759	16.9	1242	40.8	386	3.2
7	Athabasca	58°12'	111°23'	229	-0.9	-22.5	2554	17.3	1330	39.8	417	3.2
8+	Athabasca	57°08'	111°38′	274	-0.2	-21.1	2351	16.9	1340	38.0	439	3.1
9	Athabasca	56°38'	111°10′	370	0.2	-19.7	2203	16.6	1320	36.3	473	2.8
10+	Athabasca	59°53'	111°43'	183	-2.6	-25.2	2984	16.7	1174	41.9	331	3.5
11	Lac La Biche	54°22'	110°45'	550	0.8	-18.4	2035	16.4	1351	34.8	418	3.2
12+	Lac La Biche	54°38'	110°13'	610	0.6	-18.0	2039	16.2	1300	34.2	462	2.8
13	Lac La Biche	54°58'	112°10'	551	1.3	-16.8	1835	16.1	1310	32.9	503	2.6
14	Lac La Biche	55°13'	113°12'	610	1.0	-17.4	1889	15.9	1261	33.3	482	2.6
15+	Slave Lake	56°38'	114°35'	731	-0.1	-17.5	2054	14.8	1083	32.3	439	2.0
16+	Slave Lake	54°48'	116°59'	731	2.5	-12.7	1432	15.6	1294	28.3	553	2.3
17+	Slave Lake	55°14'	114°46'	610	1.5	-15.0	1701	15.7	1259	30.7	520	2.4
18	Slave Lake	54°32'	114°05'	640	1.6	-15.5	1725	15.9	1291	31.4	509	2.5
19	Slave Lake	55°29'	116°05'	610	1.4	-16.4	1794	15.8	1287	32.2	481	2.7
20	Slave Lake	55°46'	113°18'	579	0.8	-17.7	1946	16.0	1263	33.7	509	2.5
21+	Peace River	57°36'	117°31'	460	-0.5	-20.6	2340	15.8	1223	36.4	443	2.8
22	Peace River	56°59'	117°50'	610	-0.3	-19.3	2230	15.4	1174	34.7	445	2.6
23+	Peace River	56°34'	119°40'	762	-0.2	-18.0	2101	14.7	1086	32.7	454	2.4
24	Peace River	56°28'	118°05'	823	-0.1	-17.4	2073	14.8	1114	32.2	480	2.3
26	Grande Prairie	55°35'	119°35'	838	0.8	-15.0	1771	14.5	1104	29.5	522	2.1
27+	Grande Prairie	55°35'	118°18'	640	1.6	-15.7	1747	15.8	1320	31.5	483	2.7
28	Grande Prairie	55°08'	117°17'	667	2.4	-13.9	1528	16.1	1365	30.0	490	2.8
29+	Grande Prairie	54°27'	117°38'	940	1.8	-13.9	1474	14.5	1112	27.4	611	1.8
30	Grande Prairie	54°38'	117°57'	990	1.6	-12.9	1507	14.3	1088	27.2	614	1.8
31+	Edson	53°46'	118°48'	1402	0.7	-11.7	1462	12.1	773	23.8	627	1.2
32+	Edson	53°14'	117°28'	1342	1.5	-11.4	1372	13.3	932	24.7	601	1.6
34	Edson	53°19'	117°51'	1067	2.4	-10.7	1232	14.2	1079	24.9	566	1.9
35	Whitecourt	53°39'	115°42'	838	1.7	-13.7	1547	14.8	1166	28.5	568	2.1
36	Whitecourt	54°16'	115°18'	762	1.8	-13.9	1574	15.3	1219	29.2	535	2.3
37+	Whitecourt	54°11'	116°37'	945	2.3	-11.1	1341	14.8	1159	25.9	591	2.0
38	Whitecourt	54°22'	114°40' 115°30'	610	1.9	-11.1	1647	16.0	1327	31.0	480	2.8
39 40+	Roc/Clearwater	52°33'	115°30' 115°28'	1067	2.4	-15.0	1304	14.8	1153	26.2	610	1.9
	Roc/Clearwater	52°10′	115 28° 115°15'	1341	1.4	-11.5	1380	13.2 13.3	911 947	24.7 24.4	620 624	1.5
41 42	Roc/Clearwater	52°00'	115 15 115°25'	1280	1.8	-11.1	1294	15.1				1.5
42	Roc/Clearwater	52°43'	115 25' 115°47'	1036 1067	2.5	-11.6	1328 1367	14.8	1205 1161	26.7 26.6	614	2.0 1.9
	Roc/Clearwater	52°55'	113 47 114°36'		2.2	-11.8					618	
45	Bow/Crow	50°48' 51°24'	114 36 115°13'	1463	2.0	-9.8	1201	13.2	921	23.0	599 502	1.5
46+	Bow/Crow			1600	0.8	-11.5	1466	12.5	807	24.0	593	1.4
47	Bow/Crow	50°05' 49°39'	114°30'	1830	0.8	-10.3	1407	12.5	780	22.8	689	1.1
48+	Bow/Crow		114°37'	1585	2.2	-9.2	1168	13.6	943	22.8	756	1.2
446+	V 1	49°38'	110°14′	1310	2.4	-11.7	1375 2874	15.7	1243	27.4	484	2.6
/84+	Footner Lake	59°32'	117°13'	305	-2.3	-23.6	28/4	15.9	1171	39.5	382	3.1
T4	G#											
Test	<u>Sites</u> Hay River	500001	117024	270	2.2	22.1	2062	15.7	1127	200	410	20
В		59°08'	117°34'	370 670	-2.3	-23.1	2862	15.7	1137	38.8	410	2.8
C	Zeidler Mills	55°33'	114°50'	670	1.2	-15.5	1765	15.4	1210	30.9	551	2.2
D	Sexsmith	55°31'	118°30'	805	1.2	-14.9	1733	15.1	1199	30.0	509	2.4
Е	Swartz Creek	53°23'	116°30'	990	2.0	-11.6	1380	14.5	1123	26.1	580	1.9
F	Prairie Creek	52°15'	115°21'	1220	1.6	-11.7	1376	13.5	965	25.2	625	1.5
G	Chinchaga	57°50'	118°12'	470 625	-0.8	-20.8	2396	15.7	1188	36.5	448	2.7
H	Calling Lake	55°17'	113°09'	625	0.9	-17.5	1901	15.9	1254	33.4	488	2.6
J	Hanginstone	56°23'	111°26′	540	0.2	-18.8	2112	16.1	1251	34.8	534	2.3

MAT – mean annual temperature; TCM – mean temperature for coldest month; TWM –mean temperature for the warmest month; NDD degree days below $5\,^{\circ}$ C; GDD – degree days above $5\,^{\circ}$ C; CI – continentality index (TWM *minus* TCM); MAP – mean annual precipitation, AMI – annual moisture index (GDD \div MAP); LAT-latitude; LON – longitude; ELE – elevation + -populations planted on all sites.

Model (ACM I) as described by Alberta Environment (2005). Future climate data were derived from ACM II (Alberta Environment, 2004). The future climate scenario adopted in the present study corresponds closely to the HadCM3 climate change scenario for Alberta described by Barrow and Yu (2005). This scenario represents a moderate change in temperature and precipi-

tation. The future climate period adopted in this article is the year 2030–2039.

The climate variables used for biological analyses were chosen to cover a range of climatic conditions in terms of precipitation, winter and growing season temperatures, temperatures-moisture relationships and their relationship with survival and annual growth cycle

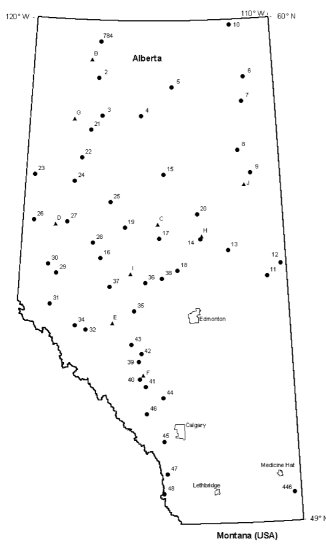


Figure 1. – White spruce populations (\bullet) and test site (Δ) locations in Alberta. Numbers and capital letters represent population and test site ID, respectively.

of conifers. Because most of the climate variables were derived from each other and were therefore highly intercorrelated, an attempt was made to reduce these variables into few biologically interpretable ones. To do this, the Pearson's correlation coefficients were calculated from the climate of population and test site locations. These correlations were then used to select one or two climate variables that best represent a particular season of the year. The selected variables were mean annual temperature (MAT), degree days below 0°C (NDD), mean temperature for the coldest month (TCM), mean temperature for the warmest month (TWM), degree days above 5°C (GDD), the continentality index (CI) expressed as temperature difference between the warmest and coldest months (CI = TWM minus TCM), mean annual precipitation (MAP) and the annual moisture index (AMI) expressed as GDD ÷ MAP.

AMI was chosen to represent a balance between heat and moisture during the growing season. High values of AMI represent dry conditions due high heat (GDD) relative to moisture (MAP) and, therefore, a potential for drought. High GDD could result from either hot sum-

mers or warm and long summers. Likewise, low values of AMI represent cool conditions relative to the available moisture, which are characteristic of regions with short growing seasons. These selected variables describe populations and test site locations in terms of their average annual temperature and moisture, average and extreme winter temperatures, average and extreme summer temperatures, and heat-moisture relationship during the growing season. In addition, studies by Rehfeldt et al. (1999, 2001, 2002), Matyas and Yeatman (1992) and our unpublished studies on the major Alberta conifers have shown that these are the main climate variables controlling growth and survival in northern conifers.

Analysis of Genetic Variation

Before performing the analysis of variance (ANOVA), data for H24 and D24 were examined and found to conform to the homogeneity of variance and normal distribution assumptions. Single-site ANOVA was performed as described in the following model:

$$Y_{ijn} = \mu + R_i + P_j + RP_{ij} + E_{ijn}$$
 [1]

where Y = response (H24 or D24), μ = site mean; R_i , P_j , RP_{ij} and E_{ijn} are effect due to ith replication, jth population, experimental error and residual for nth tree in jth population in ith replication, respectively. Except μ , all effects on the right side of equation 1 were considered random with zero expectation and variances $\sigma^2_{r_{ss}}$ (replication), $\sigma^2_{p_{ss}}$ (population), $\sigma^2_{rp_{ss}}$ (experimental error) and $\sigma^2_{e_{ss}}$ (residual). Likewise, the combined-site ANOVA were performed as described in the following model:

$$Y_{kijn} = \mu + S_k + R_{j(k)} + P_j + SP_{kj} + RP_{j(k)} + E_{kijn}$$
 [2]

where, Y = response (H24 or D24), μ = general mean; S_k , $R_{i(k)}$, SP_{kj} , $RP_{ij(k)}$ and E_{kijn} are effect due to kth site, ith replication-within kth site, interaction due to kth site and jth population, interaction due the ith replicationwithin kth site and jth population, and residual for nth tree in *jth* population in *ith* replication at *kth* test site. Except u, all effects on the right side of equation 2 were considered random with zero expectation and variances σ^2_s , (site), $\sigma^2_{r(s)}$ (replication-within site), $\sigma^2_{p_{ac}}$ (population), σ^2_{sp} (site x population interaction), $\sigma^2_{rp(s)}$ (replication-within site x population interaction) and $\sigma_{e_{nc}}^2$ (residual). SAS PROC MIXED (SAS INST., 2004) was used to obtain restricted maximum likelihood (REML) estimates of variance components and to test if these variance components were significantly different from zero. This test used the Wald Z test available in PROC MIXED. All single-site analyses used all populations (26-30) that were planted on specific sites, whereas combined-site analyses used only the 21 populations that were common to all sites.

The amount of genetic variation was estimated using an intraclass correlation, which is the ratio of the between-population genetic variance component to the between-population phenotypic variance component expressed as follows:

$$\rho_{ss} = \frac{\sigma_{\rho_{ss}}^{2}}{\sigma_{\ell_{ss}}^{2} + \sigma_{pp}^{2} + \sigma_{\rho_{ss}}^{2}}$$
[3]

$$\rho_{ac} = \frac{\sigma_{\rho_{ac}}^{2}}{\sigma_{e..}^{2} + \sigma_{zp(s)}^{2} + \sigma_{sp}^{2} + \sigma_{e..}^{2}},$$
[4]

where all variance components are as already defined. The subscripts ss and ac are used to identify single site and combined-site statistics where the two cannot be distinguished from the primary notations.

Preliminary stepwise regression analyses indicated that H24 and D24 were significantly ($P \le 0.05$) related to latitude, longitude and elevation at a place of seed origin ($R^2 = 0.37-0.86$). However, it is generally accepted that, because geographic transfer of seeds involve climatic transfer, growth and survival tend to be related to seed source geography by their relationship with seed source climate (MATYAS, 1994, 1996). Therefore, a principal component analysis (PCA) was performed using MAT, TCM, NDD, TWM, GDD, CI and MAP to examine the climatic grouping of white spruce populations in Alberta and relate this to survival and growth potential of these populations. This PCA analysis used PROC PRINCOMP (SAS INST., 2004).

Analysis of Population Response to Climatic Transfer

Although the trial at site C occupies one of the best sites climatically, it exhibited the lowest H24 and D24. Because this trials was established on an abandoned sawmill landing site, it has a great potential for soil compaction, which alters soil-water-air relationships causing tree mortality and poor growth (KOZLOWSKI, 1999). Although compacted soils may provide a special type of environment for studying genetic variation, growth and survival on these soils cannot be used for modelling population response to climatic transfer, where climate not soil characteristics is considered the main cause of the observed performance. Therefore, site C was excluded from the fitted climatic response functions. In addition, because H24 and D24 showed similar response to climatic transfer, only H24 is used in this paper to illustrate the effect of climate change on tree growth.

Preliminary analysis indicated that S24 and H24 were the lowest at site B and F, which are on the opposite side of the moisture and temperature continuum for most of the selected climatic variables. Therefore, a second degree polynomial regression was considered appropriate for examining the response of tree populations to climate change. This response function would enable us to determine the potential climatic optima for each population. Thus, H24 and S24 for individual populations were separately regressed on individual climate variables at the test site as follows:

$$y_i = b_0 + b_1 x_i + b_2 x_i^2 + e$$
 [5]

where y_i is the population mean at the ith site, x_i is MAT, TCM, NDD, TWM, GDD, CI, MAP or AMI at the ith site, b_0 , b_1 and b_2 are regression coefficients and e is the residual. The potential optimum climate for each population on individual climate variables was calculated as the climate corresponding to maximum S24 or H24 on the response function.

Although polynomial regressions have limited accuracy in predicting responses beyond the range of climate covered by field trials, they allow us to cautiously examine the potential effects of climate change on forest productivity (see Rehfeldt et al., 2001, 2002; Andalo et al., 2005; Schmidtling, 1994). In the present study, the fitted response functions were used to predict S24 and H24 for trees planted and growing in the climate expected for the 2030–2039 period. Climate predicted for the 2030–2039 period were either within the range of the tested climate for some sites or slightly beyond the range for others.

The impact of climate change was calculated as a percentage difference between performances (S24 and H24) in the 1961–1990 and in the future climate as described below:

$$\Delta_x = \frac{y_x - y_{00}}{y_{00}} \times 100 \tag{6}$$

where x is future climate, Δ_x is percentage change in H24 or S24 for trees planted and growing to age 24 years in the future climate, y_x is H24 or S24 for trees planted and growing to age 24 years in the future climate, and y_{00} is H24 or S24 for trees planted and growing in the 1961–1990 baseline climate.

We will show later that, a balance between MAP and GDD at the test site largely influenced growth and survival. During this century, the Alberta climate is expected to warm substantially. For example, by the 2030-2039 period, MAT and TCM could, respectively, increase by an average of 2°C and 3.2°C over the 1961–1990 climate. During the same period, GDD, MAP and AMI could increase by an average of 342 (28%), 20 mm (4.1%) and 0.6 (22.2%), respectively (ALBERTA Environment, 2005, 2004). This shows that in a changing climate, winters could become warmer posing only a minimum risk to tree survival. On the contrary, a disproportional increase in GDD and MAP means that growing seasons could become increasingly warmer and drier, causing tree mortality in young stands and lower annual growth for both young and old stands. Therefore, AMI was considered the most appropriate variable for predicting S24 and H24 in the future Alberta climate. Thus, future predictions of survival and growth were limited to AMI response functions.

Results and Discussion

Genetic Variability

Table 2 summarises mean growth and survival, and the levels of genetic variation for H24 and D24. Survival was moderate at sites B (55.2%) and C (54.4%) and high (75.2–96.4%) at all other sites. Significant ($P \leq 0.05$) variation among populations for S24 existed only at site D and J where the between-population genetic variance was 16.1% and 23.5% of the between-population phenotypic variance. Generally, S24 was not significantly (P > 0.05) related to geography and climate of seed origin.

Variation among populations for H24 and D24 was significant ($P \le 0.05$) on individual and across-sites

Table 2. – General means with standard errors and range of population means for growth and survival, and levels of population variation for height and diameter growth.

Site]	H24 (m)			D24 (cm)		S24	(%)
	Mean	Range	ρ	Mean	Range	ρ	Mean	Range
В	4.69 ± 0.11	3.67 - 5.21	0.130**	6.57 ± 0.19	5.02 - 7.42	0.117***	55.2 ± 1.6	31.1 – 84.4
C	3.85 ± 0.25	3.27 - 4.36	0.089**	5.81 ± 0.50	4.94 - 7.10	0.073*	54.4 ± 3.2	40.0 - 75.6
D	6.58 ± 0.16	5.52 - 7.71	0.174***	10.69 ± 0.24	9.80 - 11.41	0.043	82.9 ± 3.5	55.6 - 97.8
E	6.59 ± 0.13	5.38 - 7.20	0.179***	10.34 ± 0.24	9.05 - 11.13	0.091***	96.4 ± 0.8	88.9 - 100
F	5.04 ± 0.20	4.15 - 5.70	0.118***	8.13 ± 0.36	6.62 - 9.49	0.098***	75.2 ± 4.0	64.4 - 93.3
G	6.31 ± 0.24	5.40 - 6.78	0.100***	8.78 ± 0.38	7.21 - 9.30	0.067***	85.0 ± 2.5	71.1 - 97.8
Η	5.62 ± 0.15	4.28 - 6.48	0.209***	8.87 ± 0.23	6.80 - 10.08	0.153***	95.7 ± 1.0	82.2 - 100
J	6.50 ± 0.19	4.46 - 7.15	0.211***	9.72 ± 0.39	6.11 - 10.58	0.134***	89.0 ± 2.3	46.7 - 100
AC	5.57 ± 0.38	4.84 - 5.99	0.106**	8.52 ± 0.65	7.19 - 9.18	0.066**	71.0 ± 10.3	3 58.7 – 77.4

AC – Across sites; ρ (ρ_{ss} or ρ_{ac}) – population variance as a fraction of the phenotypic variance (intraclass correlation); * – P \leq 0.05; ** – P \leq 0.01; *** – P \leq 0.001; H24, D24, S24 – height, dbh and survival at 24 years; other codes see *Table 1*.

Table 3. – Correlations among climate variables at the population origin and PCA analysis examining the climatic grouping of white spruce populations in Alberta.

Variable			Co	orrelation Coe	fficients			Eigenvect	ors
	MAT	TCM	NDD	TWM	GDD	CI	MAP	Pl	P2
MAT	1.00							-0.351	0.500
TCM	0.92	1.00						-0.411	0.160
NDD	-0.95	-0.99	1.00					0.403	-0.244
TWM	-0.46	-0.76	0.70	1.00				0.362	0.451
GDD	-0.17	-0.54	0.46	0.94	1.00			0.282	0.678
CI	-0.85	-0.99	0.97	0.85	0.65	1.00		0.418	-0.026
MAP	0.79	0.92	-0.90	-0.79	-0.63	-0.93	1.00	-0.399	0.010
Eigenvalue	e							5.69	1.18
% of varia	nce							81.3	17.8

P1 and P2 – first and second principal components; other codes see Table 1.

(Table 2). On individual sites, the between-population genetic variance was 8.9-21.1% (H24) and 4.3-15.3% (D24) of the between-population phenotypic variance. When the population x site interaction was removed by the combined-site analysis, the between-population genetic variance was 10.6% (H24) and 6.6% (D24) of the between-population phenotypic variance. Comparable estimates of genetic variation among white spruce populations are from nursery experiments and young field provenance trials in the eastern part of the species' natural range (see LI et al., 1993; LI et al., 1997; LESSER and Parker, 2004). Because of large age differences, the amount of genetic variability observed in the present Alberta study may not be directly compared with that observed in previous studies elsewhere. However, given the older age of the Alberta study, white spruce may be more genetically variable in the western than eastern part of its natural range.

In the present study, we assumed that populations from the hybrid zone between white and Engelmann spruce were white spruce. If any of these populations were hybrids with different growth potential from that of white spruce populations in the region, the level of genetic variation observed in the present study may be slightly inflated.

Because the PCA had similar results for most sites, only results from a combined-site analysis with 21 populations are discussed. *Table 3* shows that the first (P1)

and second (P2) principal component accounted for 81.3% and 17.8% of the variance in the data, respectively. P1 had similar loading for all variables, whereas P2 had high loading for GDD. A plot of P1 and P2 shows that based on climate, Alberta white spruce could be divided into southern, central and northern regions (Figure 2). This grouping is clearly demonstrated along the P1 axis. In terms of climate, these three regions represent the cold, dry and highly continental region

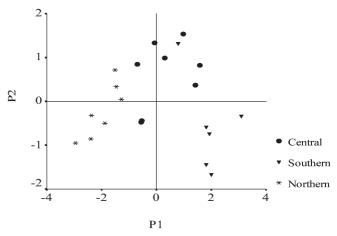


Figure 2. — Plot of the first (P1) and second (P2) principal components illustrating the division of Alberta into three white spruce climatic regions.

Table 4. – Pearson's correlations for population means between biological variables and climate of population origins.

Region		MAT	TCM	NDD	TWM	GDD	CI	MAP	AMI
Northern	H24	0.90**	0.96***	-0.94**	-0.34	0.56	-0.94**	0.90**	-0.75
	D24	0.83*	0.92**	-0.87*	-0.43	0.46	-0.92**	0.84*	-0.78*
	S24	-0.42	-0.17	0.35	-0.58	-0.60	0.02	-0.23	-0.05
Central	H24	0.44	0.23	-0.32	0.41	0.51	-0.12	0.29	0.03
	D24	0.74*	0.56	-0.65	0.31	0.53	-0.45	0.66	0.01
	S24	-0.69	-0.74*	0.77*	0.32	0.01	0.75*	-0.82*	0.46
Southern	H24	0.23	-0.68	0.35	0.53	0.58	0.85*	-0.78	0.71
	D24	0.12	-0.78	0.49	0.47	0.52	0.87*	-0.87*	0.72
	S24	-0.04	-0.87*	0.64	0.35	0.40	0.83*	-0.88*	0.65

 $^{* -} P \le 0.05$; $** - P \le 0.01$; $*** - P \le 0.001$; other codes see *Table 1* and 2.

(northern Alberta), the relatively moist and warm region at medium latitudes and elevations (central Alberta) and the moist, warm and less continental region at high elevations in the foothills and Rocky Mountains region (southwestern Alberta). Although the southern region has the highest precipitation and the least extreme temperatures, it has the lowest GDD and therefore the shortest growing season.

It should be noted that the existence of three climatic groups of white spruce populations is merely imposed by the Alberta topography and does not indicate the existence of white spruce ecotypes in Alberta. Within the three regions, variation for growth potential is strongly clinal in relation to climate of seed origin. Clines are the steepest in the northern and southern regions (*Table 4*). Growth potential was linearly related to winter temperatures (MAT, TCM, NDD), continentality (CI) and precipitation (MAP) at the place of seed origin. Similar clines were demonstrated for geographic variables in the eastern part of the species natural range (see KHALIL,

1986; Furnier et al., 1991; Nienstaedt and Riemenschneider, 1985).

Table 4 also shows that for all three regions, growth potential was not related to summer temperatures (TWM and GDD). This suggests that cool season temperatures and moisture limitations are the major climatic factors controlling genetic differentiation of white spruce in Alberta.

Population Response to Climatic Transfer

Table 5 summarises H24 and S24 response functions for the eight climate variables, which shows that both growth and survival responded most to MAP and AMI. Table 5 also shows that cool season temperatures may have more control on survival than height growth. In terms of \mathbb{R}^2 , the relationships of GDD with S24 and H24 appear to be relatively low. However, the joint effect of MAP, GDD and consequently AMI on survival and growth was apparent, especially when survival and growth at the two extreme sites (B and F) are consid-

Table 5. – Mean and range of R^2 for population response functions and the predicted potential optimum climate for annual moisture index (AMI) based on the second order polynomial regression.

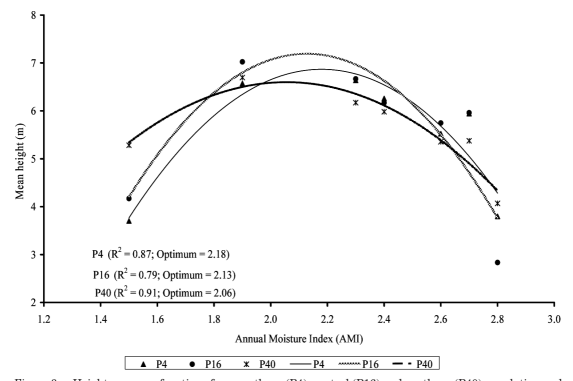
Variable	\overline{R}^2	R ² Range	SN	Predicted Average Optimum AMI	
				Region $X_{\rm I}$ $X_{\rm op}$ Δ	(%)
Survival (S24)				Survival (S24)	
MAT	0.37	0.06 - 0.78	1	Northern 3.1 2.2 -	29.0***
TCM	0.45	0.07 - 0.84	2	Central 2.3 2.2	-4.3
NDD	0.47	0.08 - 0.87	3	Southern 1.6 2.1 +	31.3*
TWM	0.33	0.06 - 0.91	2		
GDD	0.41	0.03 - 0.95	2		
CI	0.52	0.07 - 0.91	3		
MAP	0.65	0.19 - 0.92	8		
AMI	0.61	0.23 - 0.87	4		
Height (H24)				Height (H24)	
MAT	0.31	0.00 - 0.64	0		29.0***
TCM	0.33	0.01 - 0.60	0	Central 2.3 2.1	-8.7
NDD	0.37	0.03 - 0.63	0	Southern 1.6 2.1 +	31.3*
TWM	0.33	0.04 - 0.88	1		
CI	0.42	0.04 - 0.70	0		
GDD	0.37	0.04 - 0.80	0		
MAP	0.61	0.29 - 0.90	7		
AMI	0.71	0.40 - 0.93	9		

SN – number of significant regressions P \leq 0.05); R^2 – coefficient of determination; X_I – inhabited climate; X_{op} – predicted optimum climate; $\Delta(\%)$ – difference between X_I and X_{op} expressed as percentage of X_I ; * – P \leq 0.05; *** – P \leq 0.001; \bar{R}^2 – average coefficient of determination; other codes see Table 1.

ered. At site B where GDD is high, low MAP limited growth, whereas at a high elevation site F where MAP is high, growth was limited by low GDD.

Figures 3 and 4 illustrate the response of H24 and S24 to AMI for three populations selected from three distinct regions. Site B and F are at the upper and lower end of the response functions, respectively. Using all 21 popu-

lations, it was determined that the average potential optimum AMI for H24 and S24 was 2.2 compared with the average inhabited AMI of 2.4. Thus, on average, the tested white spruce requires 8.3% moister environment than its current environment, although large differences exist among regions (*Table 5*). These results show that populations in northern Alberta (highly continental and



 $\label{eq:Figure 3.-Height response functions for a northern (P4), central (P16) and southern (P40) population and the predicted optimum AMI.$

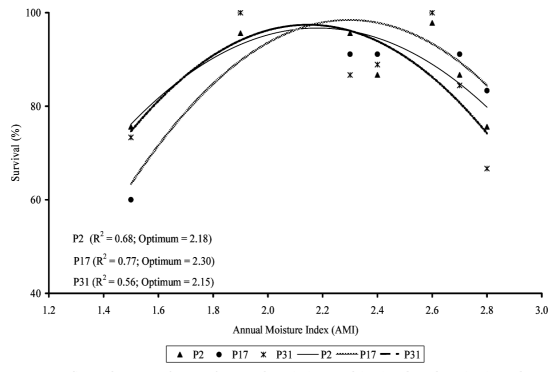


Figure 4. – Survival response function for a northern (P2), central (P17) and southern (P31) population and the predicted optimum AMI.

dry) and those in the foothills and Rocky Mountains (less continental and low GDD) inhabits less than optimal climate compared to those in central Alberta (warm and moist).

The pattern of population response to climatic transfer observed in the present study is similar to that observed in lodgepole pine in British Columbia (REHFELDT et al.; 1999, 2001) and Scotch pine from Russia (Rehfeldt et al., 2002). In both species, populations from high latitudes and other climatically marginal environments were far from their optimum climate with respect to MAT (REHFELDT et al., 1999, 2001), GDD and degree of continentality (REHFELDT et al., 2002) compared to those from low latitudes, low elevation and other climatically mild environments. In white ash (Fraxinus americana L.), Roberds et al. (1990) showed that populations from northern latitudes attained optimum height when moved 5° south of their origin, southern latitude populations reached optimum height 1.5° north of their origin and populations from the centre of the species' distribution reached optimum height in their native environments. Earlier, MATYAS and YEAT-MAN (1992) showed the same for jack pine.

The present study also showed that for white spruce in Alberta, the AMI climate for optimum survival and height growth were the same. This is somehow contrary to lodgepole pine in British Columbia where Rehfeldt et al. (1999) found that optimum growth required a more continental climate than optimum survival.

Potential Effects of Climate Change

In the present study, we used the response functions developed for the 1961-1990 AMI climate to predict H24 and S24 in a future climate scenario. The results are summarised in Table 6. They show that, as early as 2030–2039, warming could cause serious decline in survival and growth of white spruce in Alberta. Warming could intensify drought in northern and central Alberta making it difficult for white spruce to attain substantial survival and growth. At higher elevations in the foothills and Rocky Mountains region, warming could either have less negative effects or promote survival and growth. For example, at site E (990 metres above sea level), the decline in S24 and H24 would only be 2.6% and 4.7%, respectively. At much higher elevations such as site F (1220 metres above sea level), S24 and H24 could increase by an average of 27.6% and 57.7%, respectively.

Because populations from different climatic regions of Alberta exhibited different AMI optima, we found regional differences in the effect of climate change on survival and growth. *Table 6* shows that except for site E, the decline in S24 and H24 could be the greatest among populations originating from central Alberta, which currently have near optimal AMI. These populations may not be drought-hardy to survive and grow adequately in the predicted future climate. Similar inferences were made by Rehfeldt et al. (2001, 2002) who predicted that lodgepole pine and Scotch pine populations inhabiting near optimal climates would be the most affected by climate change.

In the present climate, growth and survival at higher elevations is constrained by low degree days not drought. Consistently, REHFELDT et al. (1999, 2001, 2002) argued that warming at high elevations would drive moisture indices toward their optimal values and therefore increase forest productivity. The predicted drastic increase in S24 and H24 at site F attests this. It has also been argued that as warming progresses, cool regions at high latitudes and elevations could become favourable to populations from low latitudes and elevations, respectively (Persson, 1998). Rehfeldt et al. (1999, 2001, 2002) showed that in the future warmer climate, populations from low latitudes could be the best in growth at high latitudes. Likewise, populations from low elevations could be the best at higher elevations. Table 6 shows that at site F, warming could favour populations from relatively drier regions of central and northern Alberta than local ones. Therefore, the present study confirms that warming at higher elevation regions could increase productivity due to improved growing conditions for local populations or through transfer of populations from low elevation areas.

Caution should be taken in assimilating the predictions of changes in growth and survival of white spruce in Alberta due to climate change. Although there are agreements in predictions of future temperatures, many general circulation models (GCM) differ greatly in prediction of future precipitation (LOEHLE and LEBRANC, 1996) and long-term predictions exhibit the highest uncertainties (see Barrow and Yu, 2005). In addition, predictions of H24 and S24 for future climate were derived by extrapolating the existing response functions beyond the existing white spruce test site AMI climate.

Although the predicted declines in growth and survival of white spruce due to climate change are large,

Table 6. – Mean S24 and H24 for all populations and for populations from three regions as predicted by the response functions for current (1961-1990) and future (2030-2039) AMI climates.

Site	Mean S24 (%)		Percentage Change in S24			_	Mean H24 (m)			Percentage Change in H24				
	P_{00}	P ₃₀	Mean	NT	CT	ST		P_{00}	P ₃₀	Mean	NT	CT	ST	
В	75.2	9.8	-88.5	-82.3	-94.7	-86.4		4.14	0.00	-100	-100	-100	-100	
D	93.7	61.7	-34.2	-31.6	-38.5	-31.5		6.13	2.56	-58.1	-49.4	-66.5	-57.0	
E	93.4	90.8	-2.6	-1.3	-2.7	-4.0		6.10	5.80	-4.7	-0.6	-6.3	-7.4	
F	75.7	96.0	+27.6	+28.5	+31.5	+20.7		4.12	6.44	+57.7	+60.8	+65.7	+43.6	
G	82.1	33.2	-61.4	-57.5	-69.1	-55.6		4.83	0.36	-93.1	-87.8	-97.7	-93.1	
Η	87.0	33.2	-62.9	-59.1	-70.4	-57.1		5.37	0.36	-93.6	-88.6	-97.9	-93.6	
J	95.4	61.7	-35.2	-32.4	-39.6	-32.7		6.34	2.56	-59.4	-50.6	-67.7	-58.4	

 P_{00} and P_{30} – 1961–1990 and 2030–2039 periods, respectively; NT, CT and ST – northern, central and southern regions, respectively; other codes see *Table 1*.

they are consistent with the present natural distribution of the species in Alberta. From the Alberta Climate Model (Alberta Environment, 2005), the Northern Fescue Plains seed zone (ASRD, 2005) immediately south of the limit of the white spruce distribution in Alberta has an average AMI of 3.9. This moisture index is close to the AMI of 3.7 predicted for site B in northern Alberta for the 2030–2039 period. At this AMI, site B is predicted to have an average S24 of 9.8% and may not sustain meaningful height growth (Table 6). Table 1 shows that the populations tested in this trial originated from locations with the AMI of 1.2–3.5. Therefore, the upper threshold of AMI for white spruce in Alberta could be in the range between 3.5 and 4.0.

Conclusions

Provenance trials are essential for matching tree populations to planting sites to maximise production while safeguarding adaptation. Based on our analysis of the Alberta series of white spruce provenance trials, we have concluded that: (1) The level of genetic variation in growth potential among white spruce populations is moderate but slightly higher than that observed in other areas of North America. (2) The pattern of variation is clinal whereby growth potential declines from central Alberta northward in response to a decrease in winter temperatures and precipitation, and an increase in continentality of seed origin. (3) Although many climate variables may have a role in determining survival, a balance between GDD and MAP had the strongest effect on both survival and height growth on Alberta test sites. Therefore, AMI provided a simple measure of drought potential, which was used to describe survival and growth of white spruce in Alberta in the present climate and predict the impact of climate change on forest productivity in the future. Using AMI, we determined that populations at the cold and dry areas of northern Alberta and those at high elevations in the foothills and Rocky Mountain regions inhabit climates with significantly different AMI than their potential optimum climate. Populations at mid-latitudes and elevation in central Alberta are close to their optimum AMI. (4) The predicted changes in Alberta climate would seriously reduce survival and growth of white spruce in central and northern Alberta. Because of the precipitation pattern, northern and central Alberta would become increasingly dry making it difficult for white spruce to attain significant survival and growth. On the contrary, survival and growth would increase in the Upper Foothills and Rocky Mountains regions in southwestern Alberta as warming extends the growing season, increases GDD and boosts AMI toward optimal values.

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Gene Flow Between Introduced and Native *Eucalyptus* Species: Morphological Analysis of Tri-Species and Backcross Hybrids Involving *E. nitens*

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Summary

Morphometric analyses were conducted on second-generation tri-species and backcross hybrids in Eucalyptus. These hybrids were all produced using pollen from two $E.\ nitens\ x\ cordata\ F_1$ hybrids and controlled pollination techniques. Tri-species hybrids were created with $E.\ gunnii,\ E.\ ovata$ and $E.\ viminalis$ as females, while backcrosses were produced with $E.\ cordata$. Multivari-

ate analysis of seedling characteristics indicated that eighty percent of the backcross hybrids fell within the morphological range of $E.\ cordata$. All three cross combinations of the tri-species hybrids were biased away from $E.\ nitens$ and towards their maternal parent and $E.\ cordata$. The inclusion of data for first-generation (F_1) hybrids between the pure parental species in the current work showed the F_1 's to be easily distinguishable from pure species, compared to second-generation hybrids. The use of morphology for detecting second-generation hybridisation involving exotic plantation species and native eucalypt populations will therefore be

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