

On the representativeness of plot size and location for scaling transpiration from trees to a stand

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[1] Scaling transpiration from trees to larger areas is a fundamental problem in ecohydrology. For scaling stand transpiration from sap flux sensors we asked if plot representativeness depended on plot size and location, the magnitude of environmental drivers, parameter needs for ecosystem models, and whether the goal was to estimate transpiration per unit ground area $(E_{\rm C})$, per unit leaf area $(E_{\rm L})$, or canopy stomatal conductance $(G_{\rm S})$. Sap flux data were collected in 108 trees with heat dissipation probes, and biometric properties were measured for 752 trees within a 1.44 ha Populus tremuloides stand along an upland-to-wetland gradient. $E_{\rm C}$ was estimated for the stand using eight different plot sizes spanning a radius of 2.0–12.0 m. Each estimate of $E_{\rm C}$ was derived from 200 plots placed randomly throughout the stand. We also derived leaf area index (L), canopy closure (P_{CC}), and the canopy average reference stomatal conductance (G_{Sref}), which are key parameters used in modeling transpiration and evapotranspiration. With increasing plot size, $E_{\rm C}$ declined monotonically but $E_{\rm L}$ and G_{Sref} were largely invariant. Interplot variance of E_{C} also declined with increasing plot size, at a rate that was independent of vapor pressure deficit. Plot representativeness was dependent on location within the stand. Scaling to the stand required three plots spanning the upland to wetland, with one to at most 10 trees instrumented for sap flux. Plots that were chosen to accurately reflect the spatial covariation of L, P_{CC} , and G_{Sref} were most representative of the stand.

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1. Introduction

[2] Scaling up from plots to stands and from stands to larger areas is a fundamental problem in ecohydrology. It is an essential step in combining information from multiple spatial, temporal, and organizational scales at which systems are observed, or as a pragmatic solution to the problem of having limited observational data within a larger region. No single scale is always sufficient for characterizing a system, as each scale imposes a sampling bias [*Levin*, 1992]. For example, regional scale models of canopy transpiration (E_C) typically combine vegetation classification, leaf area index (*L*), and canopy stomatal conductance (G_S) [*Bonan*, 1991; *Wood et al.*, 1992; *Foley et al.*, 1996; *Sellers et al.*, 1997; *Baldocchi et al.*, 2002; *Su et al.*, 2007]. Most of these models have the potential of blurring mechanisms when they consider

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evapotranspiration (i.e., evaporation from soils and wet canopies as well as transpiration) rather than just $E_{\rm C}$. However, a potentially bigger source of uncertainty lies with the current scaling logic where a set of representative stands is identified based on the composition of vegetation in a region, a measurement plot is identified within each stand, and flux and parameter values are obtained for the plot. The plot is assumed to be representative of its respective stand, which in turn is representative of all stands of a similar vegetation composition at the regional level. The information obtained from the plot is distributed onto the region by mapping the respective observations to their vegetation composition on the classification map [Mackay et al., 2002]. With respect to scaling to the stand, plot representativeness can be interpreted in different ways. One could consider a plot representative if its scaled flux values match those of the whole stand (i.e., absolutely representative), if fluxes of the plot and stand have similar dynamics with different scalar values (i.e., dynamically representative), or if parameters derived from the plot can be used to accurately simulate fluxes at the stand level (i.e., parametrically representative). Errors typically occur when scaling up from representative plots if the scaling process is nonlinear [Raupach and Finnigan, 1988; Baldocchi et al., 1991; Band et al., 1991; Norman, 1993]. Moreover, even scaling of

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linear quantities can introduce error if a plot is not representative in the respective interpretation employed. Recent studies using sap flux to scale-up $E_{\rm C}$ in heterogeneous forests suggest that finding representative plots may be challenging with respect to quantifying absolute values and dynamics of $E_{\rm C}$, and deriving $G_{\rm S}$ [Kumagai et al., 2005b; Tromp-van Meerveld and McDonnell, 2006; Adelman et al., 2008; Kumagai et al., 2008; Loranty et al., 2008; Traver et al., 2009]. The purpose of this paper is to examine the implications of a plot's size and location for its representativeness in scaling $E_{\rm C}$ from sap flux and quantifying L, $G_{\rm S}$, and canopy closure inputs to stand or regional models.

[3] Measurement of xylem water transport using sap flux sensors and scaling to canopy transpiration at the level of whole tree, stand, or larger areas has matured over the past few decades. For example, methods for scaling from constant-heat type sensors to whole trees have an extensive set of protocols [Granier, 1987; Phillips et al., 1996; Clearwater et al., 1999; Ewers and Oren, 2000; Meinzer et al., 2001; James et al., 2002; Ford et al., 2004; Oishi et al., 2008]. Methods for scaling from individual trees to plots and then to stands differ with respect to plot size used for sampling, replication of plots, the number of trees sampled per species, and the scalar used to convert flux per tree to flux per plot. Table 1 presents a cross section of studies that have provided sufficient information for scaling from sap flux sensors to some ground area. In these studies the ground area to which sap flux sensor observations are scaled vary from 8 m² to 60,000 m² (or 6 ha), and include circular plots [Oren et al., 1998a; Ewers et al., 2002; Ewers et al., 2005; Pataki et al., 2005; Simonin et al., 2007], noncircular plots [Hatton et al., 1990; Cermak et al., 1995; Vertessy et al., 2001; Cienciala et al., 2002; Ewers et al., 2002; Herbst et al., 2007], transects [Kurpius et al., 2003], and whole stand or hillslope boundaries [Zimmermann et al., 2000; Tromp-van Meerveld and McDonnell, 2006]. In most of the studies in Table 1 the number of plots used to represent a larger area is quite small, and in two-thirds of the studies this is limited to 1 or 2 plots. The number of trees instrumented for sap flux per species per plot varied from 1 to 19, with approximately half the studies using 8 or fewer sap flux trees per species per plot. Stand heterogeneity is included in the scaling processes by weighting each sap flux sensor by a representative scalar [Hatton and Wu, 1995], such as sapwood area, basal area, leaf area, tree circumference, or some other weighting measures. The number of scalar measurements (e.g., basal area) made typically far exceeds the number of trees instrumented for flux, as the scalars are typically easier to obtain and can be the largest source of variability in $E_{\rm C}$ [Vertessy et al., 1997; Ford et al., 2007].

[4] A number of variables contribute to the spatial heterogeneity of transpiration within a single species, including sap flux density variations [*Medhurst et al.*, 2002; *Bovard et al.*, 2005; *Ewers et al.*, 2005; *Kumagai et al.*, 2005a; *Pataki et al.*, 2005; *Ewers et al.*, 2007; *Herbst et al.*, 2007; *Adelman et al.*, 2008], the influences of a scalar such as sapwood area [*Vertessy et al.*, 1997; *Bovard et al.*, 2005; *Herbst et al.*, 2007; *Kumagai et al.*, 2007; *Loranty et al.*, 2008], and radial declines in sap flux density with increasing depth into the sapwood [*Phillips et al.*, 1996; *Pausch et al.*, 2000; *Schafer et al.*, 2000; *Ford et al.*, 2004; *Hultine et al.*, 2007]. These variables have been associated with spatial variations in edaphic properties [*Schiller et al.*, 2002; *Eberbach and Burrows*, 2006; *Tromp-van Meerveld and McDonnell*, 2006; *Wullschleger and Hanson*, 2006; *Nadezhdina et al.*, 2007], stand structure [*Cienciala et al.*, 2000; *Taylor et al.*, 2001; *Medhurst et al.*, 2002; *Giambelluca et al.*, 2003; *Simonin et al.*, 2006; *Hultine et al.*, 2007], and age [*Zimmermann et al.*, 2000; *Phillips et al.*, 2002; *Delzon and Loustau*, 2005; *Ewers et al.*, 2005], as these affect root extractable water, convective and radiative energy transfers, and plant water relations. Spatiotemporal analyses have shown temporal changes in the spatial heterogeneity of tree transpiration associated with tree-level responses to environmental drivers such as vapor pressure deficit [*Adelman et al.*, 2008; *Loranty et al.*, 2008; *Traver et al.*, 2009].

[5] Given such sources of variability we asked the following questions: (1) How representative are plots of forest transpiration for a stand; (2) does plot representativeness change with the magnitude of environmental drivers and G_S ; and (3) does plot representativeness differ for estimates of E_C and canopy transpiration per unit leaf area (E_L)?

2. Methods and Materials

2.1. Study Site

[6] The study was conducted in the Chequamegon-Nicolet National Forest near the town of Park Falls (45.95°N, 90.27°W), Wisconsin, United States. The area represents the interface between northern temperate and southern boreal ecosystems and is part of the northern highlands physiographic province. The bedrock consists of Precambrian igneous and metamorphic rock, overlain by 8-90 m of glacial and glaciofluvial material. Geomorphic features in the area are outwash, pitted outwash, and moraines resulting in gently rolling topography. Climate is characterized by long winters and a short growing season with mean January and July temperatures of -12°C and 19°C, respectively [Fassnacht and Gower, 1997]. Data for the study were collected at a site located approximately 800 m southeast of the WLEF eddy flux tower [Bakwin et al., 1998; Davis et al., 2003]. Field observations for the study were collected in a 20 year old regenerating *P. tremuloides* stand during the summer of 2005. We selected P. tremuloides because it had the highest transpiration per unit leaf area of any species measured in northern Wisconsin [Ewers et al., 2002]. We chose the particular stand because it was spatially heterogeneous with respect to topography and basal area. The stand comprised a transition from forested wetland to upland forest over a 2-3 m variation in elevation, with the wetland dominated by speckled alder (Alnus incana (DuRoi) Spreng) and white cedar (Thuja occidentalis). The upland positions also had balsam fir (Abies balsamea (L.) Mill) in the overstory, and Sphagnum spp. covered all on hummocks in the wetland areas.

2.2. Measurements

[7] The study was conducted in 16 25×25 m substand blocks within a 1.7 ha stand (Figure 1). These blocks were a sampling construct, and there were no substantial differences in basal area between the sample areas and areas between the blocks. Two of the blocks had no *P. tremuloides*, as they were in a *T. occidentalis* wetland, and so

Table 1.	Summary c	of Sap Flux S	studies in Which Plo	t Size and Sap Flux Scaling	g Parameters Are Reported ^a				
Plot Area (m ²)	Number of Plots	Number of Species	Trees/Species (per Plot)	$A_{ m B}~({ m m}^2~{ m ha}^{-1})$	$A_{ m S}~({ m cm}^2~{ m m}^{-2})$	Scalar ^b	$IGBP^{c}$	Biome ^d	Reference
~	9	-	1			Т	EN-P	TE	Hatton et al. [1990]
6	16	1	1	27.7, 13.9	19.9, 9.7	$A_{\rm S}$	EB-P	TE	Taylor et al. [2001]
20	144	С	$79, 41, 9^{e}$. 1	5.9, 3.5, 19.7	$A_{\rm S}$	MF	TE	Loranty et al. [2008]
20	144	7	93, 41 ^e	I	5.3-11.8, 2.7-4.8	$A_{\rm S}$	MF	TE	Traver et al. [2009]
25	4	1	8	10.1 - 24.9	9.0-21.2	$A_{ m S}$	EN-P	TE	Ewers et al. $[2000]$
25	4	1	8	10.1 - 24.9	4.7–23.7	$A_{\rm S}$	EN-P	TE	Ewers et al. [2001]
25	1	1	10	I	-	$A_{\rm S}$	DB	ΤΕ	Roberts et al. [2005]
30	ŝ	1	1	20.0		C	EN-P	BO	Mellander et al. [2006]
34	144	2	99, 41°	I	33.6-47.4, 86.9-92.5	$A_{ m S}$	MF	ΤE	Traver et al. [2009]
50	48	4,	54, 37, 11, 27 ^e	24.7, 7.4, 2.6, 4.9	11.0, 1.2, 2.1, 3.5	$A_{\rm S}$	MF	Ξ	Adelman et al. [2008]
53 60 80			10	14.9	12.7	$^{\rm As}$	EN-P	HE C	<i>Oren et al.</i> [1998a, 1998b]
60-80	4 0		01 0	9.4-20.0	4. /-23./	$A_{ m S}$	EN-P	Da F	Philips et al. $[2002]$ and Ewers et al. $[2001]$
00	ז ע	-	70	150300	I	' -	7-8U	1 E	Aang et at. [2002]
0/	<u>א</u> ר ב	1 -	9, J 8 or 10			$^{\rm AB}$		3 8	Cauting et al. $[2006]$
0/	7 -	- c	0 0F 10 0	20.0.2.7	7.01-0.1	$^{\rm VS}$	EIN	3	$E_{\text{LIVERS}} = \frac{1}{2} \left[\frac{1}{2} 000, \frac{1}{2} 000 \right]$
100	- (1 -	0 6	38 0 63 0	10.4 32 1	AS AS	JNR DR	a Ca	EWETS Et ut. [2002, 2007] Hore of al [1007]
100	10		1, u	20.0, 03.0 20 62 0	10.4 201	SU			Hoad and Hundle [1007]
100	16	- c	12, 12 2 or 12	Jo.U, 0J.U	19.4, 32.1	S _V	93	C E	Togg and Hurale [1997] Cont of al [2006]
111	0 -	7 -	CT 10 C	15.0	12.0	۹ -			0.0011 et al. [2000]
111			0 0	8.CI 3.13	15.8	$^{\rm AS}$	EN D	1E	<i>Dren et al.</i> [1998a, 1998b] E
C11 211		- ,	o <u>5</u>	0.10	5.65 5.75 1 8 1 5 0 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5	SV VS			$E_{Wers} \in ut. [2002, 2007]$
011 0112		n -	12	0.4, 0.9, 1.1 40.0	0.2-1.8, 1.8-2.7, 0.3 2.0.24.0	$^{\rm VS}$	EN		[2002] <i>In 19 Stand</i>
113 113	- c		12	49.9 66.7 33.7	0.0-24.0	$^{\rm AS}$		C H	Databi at al. [2002] Databi at al. [2005]
211 711	- 1		10	16.02, 235.7	- 11	$^{\rm AS}$		1	<i>Tatakt et al.</i> [2002] <i>Draw at al</i> [1008a 1008h]
11/		- c	10			SN AS			Ureht et al. [1990a, 19900] Evicans of al [7005]
166	12	1 -	- 10 r	2.4, 2.2 15 0-30 0	0.2, 0.0	SP PD	DB-P	G E	Livers et al. [2002] [Iddling et al. [2008]
198	-	. –	9	56.4	ı	N/A	EN	E	Oliveras and Llorens [2001]
203	1	1	14	54.2	ı	A_{L}	EN-P	TE	Ford et al. $[2007]$
207	1	1	14	3.9		$A_{\rm S}$	EN-P	TE	Kumagai et al. [2005a]
289	1	1	12	68.1		A_{L}	EN-P	TE	Ford et al. $[2007]$
292	1	1	14	77.2		$A_{ m L}$	EN-P	ΤE	Ford et al. [2007]
314	1	ς	8	24.4, 16.1, 1.1	6.1, 16.1, 0.6	$A_{ m S}$	ΡW	ΤE	Ewers et al. [2002, 2007]
314	1	9	1-4	21.7 - 34.9	1 .	$A_{ m S}$	EN-P	ΞI	Wilson et al. [2001]
318	_ ,	,	23	71.7	36.3	$A_{\rm S}$	EN-P	IE	Kumagai et al. [2007, 2008]
321			<u>0</u> r	118.7	46.0	$A_{\rm S}$	EN-P	E E	Kumagai et al. [2007, 2008]
200 200	- c		- (573 645	1	AB UL AC W		3 8	Fharkach and Duranus [2007]
000	1 -	- t	7 5	04.0 15 0		\$ -			Evervach and burrows $[2000]$
000	- (0 10	C.C4 ∆N	0.24-0.0	$^{\rm AS}_{\rm f}$	EN FR	Dd Dd	Ewers et al. [2002] Hatton and Wi [1005]
000	1 -		, 10 L	2.7.1 2.7.2	I	SU 1		TE	[COUL] hy him hount
000 625		- (, s 8 s	32:2 110 fo 16 2 (As)	1 1	AB ULAC	DB	I F	Vaueznaina et al. [2007] Vertesm et al. [2001]
020 633		1 —	с., С. Г.	up w 10.2 (113)	1 5-16 7	<u>م</u> ر	EN E	HT H	Simonin of al $[7006, 2007]$
707	- 4	+ m	× ∞	8.2. 7.5. 5.4	0.9. 2.9. 2.7. 0.4–1.6	Su Ve	MF	BO	Ewers et al. [2005]
750	ŝ	1	11–13	14.0-68.0		$A_{\rm L}$	DB-P	TE	Medhurst et al. [2002]
800	1	9	6, 6, 2, 1, 1, 1	1.4 - 11.3	1.0-5.1	$A_{\rm S}$	DB	TE	Oren et al. $[2001]$
800	1	2	8	18.8, 9.7	12.6, 4.2	$A_{ m S}$	DB	TE	Ewers et al. [2002, 2007, 2008]
869		-	4	7.77	1.5 - 16.7	$A_{\rm S}$	EN	ΤE	Simonin et al. [2006, 2007]
900 000	4,	4,	4~ ;	35.6-47.6	I	$A_{\rm S}$	DB	E	Herbst et al. [2007]
960	-	_	14	24.8		A_{S}	MF	ΠE	Tromp-van Meerveld and McDonnell [2000]

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tumber of sap flux trees per species per plot, the	re made, the n	trements we	ap flux measu	le number of species in which se	in are the number of plots employed, th	thest plot size. Show	n lowest to hig	re sorted from	^a Studies a
Roupsard et al. [2006]	TR	EB-P	$A_{\rm S}$	1	8.3	10	1	1	60,000
Unsworth et al. [2004]	TE	EN	$A_{\rm S}$	4.6, 11.1, 1.4	I	9	ę	1	40,000
Williams et al. [2004]	TE	DB-P	$A_{\rm S}$. 1	N/A	8	1	1	40,000
Oguntunde and van de Giesen [2005]	TR	EB-P	$A_{\rm S}$	2.3, 3.9	3.3, 6.0	, , ,	1	1	20,000
Wullschleger et al. [2001]	TE	DB	$\tilde{A}_{ m S}$	11.9	22.5	9, 1, 1, 1, 1, 1	9	1	19,200
<i>lida et al.</i> [2006]	TE	MF	$A_{\rm S}$	9.7, 28.4	18.1, 32.6	9 total trees	4	1	17,000
Zimmermann et al. [2000]	BO	EN	$A_{\rm S}$	4.7, 11.4, 5.7, 7.7, 6.8, 3.7	6.6, 22.0, 28.0, 30.4, 42.8, 10.6	6-12	1	9	15,000
Bovard et al. [2005]	TE	DB	$A_{ m S}$	0.5-7.7	2.1–12.2	4	4	1	11,000
Delzon and Loustau [2005]	TE	EN	$A_{ m S}$	5.6 - 13.1	up to 40.0	6 or 7	1	4	$10,000^{h}$
Phillips et al. [2002]	TE	EN	A_{L}	23.9, 12.7, 4.5	46.9, 38.4, 35.6	9, 9, 6	1	б	$10,000^{h}$
Granier [1987]	TE	EN-P	$A_{\rm S}$	25.2, 17.3	N/A	S	1	7	10,000
Cermak et al. [1995]	TE	EN	$A_{ m B}$	I	27.9, 6.8	12	7	1	10,000
Tang et al. [2006]	TE	MF	$A_{\rm S}$	13.8, 2.9, 2.9	23.4, 4.9, 5.0	14, 15, or 19	ę	1	10,000
Wullschleger and Hanson [2006]	TE	DB	$A_{\rm S}$	11.8–12.4	23.1 - 24.6	1 or 2	7	б	6,400
Davi et al. [2005]	TE	DB	C	I	15.6	10	1	1	6,000
Vertessy et al. [1997]	TE	DB	$A_{\rm B}$ or $A_{\rm L}$	3.2	50.0	10	1	1	4,900
Cienciala et al. [2000]	TR	EB-P	U U	10.2, 12.4	18.6, 40.4	10, 6		0	4,000
Cienciala et al. [2002]	BO	EN	°U	0	20.0	10		. –	4.000
Tesken and Sheriff [1006]	TH T	EN-P	4~	85.0	101 6	50	o –	1	4,000
Giambelluca et al. [2003]	1K TF	EB	$^{A_{\rm S}}$	C.01	7:17	- رو م	n v	- (2,000 2,000
Yue et al. [2008]	TE	SO	A_{L}	1	N/A	9	-	1	2,500
Bowman et al. [2005]	TE	MF	M	I	N/A	5	1	1	2,500
Hultine et al. [2007]	TE	DB	$A_{\rm S}$	x	45.0 (As)	10-17	4	6 ^g	2,500
Vertessy et al. [1995]	TE	DB	$A_{\rm S}$	8.0, 4.2	21.4, 9.5	19, 7	7	1	2,500
Dunn and Connor [1993]	TE	EB	$A_{\rm S}$	6.7, 6.1, 4.0	56.6, 63.1, 78.6	19, 5, 8		ŝ	2,500
Giamhelluca et al. [2003]	TR	EB	₹. 7°	10.6	21.4	"	"	_	1,965
Oren et al. $[2001]$	TE	PW-P	$A_{\rm S}$	31.7	73.6	10°	1		1.963
Pataki and Oren [2003]	TF	DR	5 V	0 6-6 1		2-6	. 9		1 963
Kurpius et al. [2003]	TE	EN-P	As	8.0	9.4	4	ı —	. –	1.800
Lundblad and Lindroth [2002]	BO	EN	As	2.3. 2.3. 0.8. 2.3	29.9. 44.0. 26.9. 39.5	12 per stand	5	4	1.600
Dunn and Connor [1993]	TE	EB	As	4.2	73.2	ŝ			1.600
Ewers et al. [2008]	TE	DB	As	12.2	15.4	ŝ	ŝ		1.256
Pataki et al. [2000]	TE	MF	An An	24.1-48.8	43.0 - 91.1	Ś	4	-	1.256
Ewers et al. [2008]	TE	DB	$A_{\rm s}$	13.8	87.5	19	б	1	1.212
Lundblad and Lindroth [2002]	BO	EN	$A_{\rm S}$	1.5, 1.4	17.8, 28.0	12 per stand	7	7	1,200
Bernier et al. [2002]	TE	DB	$A_{\rm S}$. 1	28.2	5	1	1	1,100
Engel et al. $[2002]$	TE	DB	$A_{\rm S}$	0.9, 1.1	4.8, 10.3	9	1	2	1,000
Schiller et al. [2002]	TE	SO	A_{L}	I	7.5	8	1	4	1,000
Schiller et al. [2007]	TE	DB-P	C	I	8.2–8.9	8	1	2	1,000
Reference	Biome ^d	IGBP ^c	Scalar ^b	$A_{\rm S}~({ m cm}^2~{ m m}^{-2})$	$A_{\mathbf{B}} \ (\mathbf{m}^2 \ \mathbf{ha}^{-1})$	(per Plot)	of Species	of Plots	(m ²)
c \$	ہ ب	34401	م - ت	2 -2,	2 - 1	Trees/Species	Number	Number	Plot Area

general biome, and cited work.

^bScalars: B, branch area; C, circumference; D, canopy diameter; T, Derichlet tessellation; W, weighted averaging; A_s, sapwood area; A₁, heaf area. ^cIGBP global land cover classes: EN, evergreen needleleaf; EB, evergreen broadleaf; DB, deciduous broadleaf; MF, mixed forest; OS, open shrub; PW, permanent wetland; "–P" indicates plantation. ^dBiome types: BO, boreal; TE, temperate; TR, topical. ^eShows total number of trees per species among all plots, selecting the dominant tree of each species in each plot. ^fFive scalars tested were leaf area, sapwood area, diameter, tessellation, and field theory. ^sSix plots established for sapwood area calculation.

Table 1. (continued)



Figure 1. Map of study site showing distribution of *Populus tremuloides* (trembing aspen) sap flux trees (white circles) and all other *P. tremuloides* trees measured (black circles). Superimposed are 11 randomly located 2-12 m radius plots (gray circles). Shown also are 16 25 m × 25 m blocks used in the cyclic sampling design. Blocks a–n contained one or more *P. tremuloides* trees.

the total effective stand area was 0.875 ha. The remaining 14 blocks consisted of forested wetland (blocks b, e, f, and i) and upland forest. In each block, 9 cyclic sampling plots with a 2.5 m radius were arranged using lags of 5 and 10 m in two dimensions [Loranty et al., 2008]. The cyclic sampling scheme maximizes the efficiency with which a fixed number of samples is distributed in space, and optimizes the sampling for spatial analysis [Burrows et al., 2002]. However, in each sampling plot we recorded the absolute (Easting, Northing) locations of the dominant P. tremuloides trees rather than sampling plot centers, and so measured fluxes were related to their absolute location in space. Dominant trees were selected for flux measurements as these were expected to be most representative of edaphic and other environmental characteristics limiting growth in the respective sampling plots. This is justified because tree size impacts on sap flux could still be investigated when many trees were sampled across the site [Adelman et al., 2008; Loranty et al., 2008; Traver et al., 2009]. Moreover, recent work has shown that G_{Sref} as well as differences in $E_{\rm C}$ among individual trees can be explained by competition for light [Loranty, 2009; Loranty et al., 2010; M. M. Loranty et al., Competition for light between individual trees lowers reference canopy stomatal conductance: Results from a model, submitted to Journal of Geophysical Research, 2010], and so even the bias introduced by measuring sap flux in dominant individuals could be corrected with rela-

tively simple intertree competition data. We also recorded height and DBH for each dominant tree. In addition, we recorded the location, height, and DBH of every P. tremuloides individual within 5 m of each dominant tree. This resulted in an inventory of 109 dominant and 644 additional P. tremuloides trees, for a total of 752 trees distributed among the 14 0.06 ha blocks. One dominant tree was eliminated from the analysis, as it was a much older remnant of the previously clear-cut stand. Mean basal areas, on a per-tree basis, for the dominant and other P. tremuloides trees were 80.3 (\pm 5.3 standard error) cm² and 70.5 (\pm 2.1 standard error) cm², respectively. Selection of the dominant tree in each sampling plot slightly biased the flux measurements toward higher mean basal area. However, this was not considered a problem for the scales in this study, as basal areas were spatially autocorrelated at up to 70 m lag distance [Loranty et al., 2008] and the primary source of variability of reference $G_{\rm S}$ was competition among trees for light [Loranty, 2009; Loranty et al., 2010, also submitted manuscript, 2010].

[8] In each dominant *P. tremuloides* we installed Graniertype 20 mm length constant-heat sap flux sensors [*Granier*, 1987]. One sensor pair was placed on the north side of each tree at breast height (1.3 m). Intratree scaling followed *Ewers et al.* [2002], which accounted for bark thickness, sapwood depth, and radial and circumferential trends in sap flux. We assumed the same relationships in this study. To

Radius (m)	Trees (per Plot)	A_{plot} (m ²)	$(m^2 m^{-2})$	$\begin{array}{c} G_{\rm Sref}^{\ \ b}\\ (\rm mmol\ m^{-2}\ s^{-1})\end{array}$	$(\text{mmol } \text{m}^{-2} \text{ s}^{-1})$	$\begin{array}{c} P_{\rm CC}{}^{\rm c}\\ ({\rm m}^2~{\rm m}^{-2})\end{array}$
2.0	1–3	12.6	5.96 (0.23)	87.3 (6.1)	51.4	1.08
2.5	1-4	19.6	4.92 (0.25)	82.5 (6.0)	49.4	0.84
3.0	1-4	28.3	3.83 (0.15)	86.4 (5.2)	52.0	0.75
4.0	1-5	50.3	3.28 (0.17)	81.1 (5.0)	46.6	0.58
5.0	1-5	78.5	2.81 (0.14)	82.8 (5.1)	48.3	0.49
7.0	1-7	153.9	2.13 (0.12)	77.7 (4.8)	42.6	0.39
9.0	2-8	254.5	1.86 (0.10)	79.9 (4.7)	47.4	0.34
12.0	2-10	452.4	1.48 (0.09)	95.9 (5.6)	56.2	0.25

 Table 2.
 Number of Sap Flux Trees per Plot and Mean Parameters for Plots at Each Plot Size and Respective Plot

 Area^a

^aHere A_{plot} is plot area. Leaf area index (L) and canopy closure (P_{CC}) were derived allometrically from site-specific data. Reference canopy average stomatal conductance (G_{Sref}) and sensitivity of stomatal closure to vapor pressure deficit (m) were derived from boundary line analysis of canopy average stomatal conductance versus vapor pressure deficit.

^bShows mean at each plot size with standard error in parentheses.

^cCanopy closure exceeding 1.0 occurred when total crown cross-sectional area at canopy base exceeded plot area.

include nighttime transpiration, baselines were determined from the maximum temperature difference over a week long period [Oishi et al., 2008]. Sap flux measurements for the study were made from mid-June to mid-July 2005. The period 1-7 July was used for the present analyses, as it was the one contiguous period in which sensors in no trees were affected by power outages. Concurrently, air temperature (T_A) and relative humidity (R_H) (Vaisala HMP 45C, Vaisala Oyj, Helsinki, Finland) were measured at two-thirds canopy height (~7 m). Vapor pressure deficit (D) was calculated from T_A and R_H based on equations adapted from [Goff and Gratch, 1946]. Ewers et al. [2007, 2008] showed that P. tremuloides and other stands in this area are well coupled to the atmosphere because D measurements at 2/3 canopy height from six different stands are the same as D measurements at 30 m on the WLEF tower (regression analysis indicated that the slope of the two measurements was not different from 1 and the intercept was not different from zero). This also justifies an assumption of D being spatially unaffected by varying transpiration rates in each stand. Sap flux, temperature, and relative humidity measurements were recorded every 30 s (CR10X, Campbell Scientific, Logan, UT, United States) and aggregated to 30 min values. Wind speed, photosynthetically active radiation (Q_0) , and precipitation measurements from the nearby WLEF (~1 km) [Davis et al., 2003] and Lost Creek (~10 km) [Cook et al., 2004] flux towers were used as model inputs.

[9] Total height was measured for each tree using a laser rangefinder and clinometer with triangulation methods outlined by the Eastern Native Tree Society [Bolzan, 2004]. Leaf area $(A_{\rm L})$ (m²) and projected crown area $(A_{\rm C})$ (m²) were determined for each tree through allometric relationships with tree basal area $(A_{\rm BT})$ (cm²). These relationships were developed through destructive harvesting of 13 P. tremuloides trees spanning the wetland to upland. All harvested trees fit allometric relations, $A_{\rm L} = 10.2 \ln A_{\rm BT} - 24.9$ (r² = 0.97; N = 13) and $A_{\rm C} = 1.74 \exp(A_{\rm BT}*0.0111)$ (r² = 0.61; N = 13). For each tree we used the allometric relationship for sapwood depth, $d_{\rm S} = -1.91 + 7.159 \text{*DBH}^{0.7501}$, where $d_{\rm S}$ is given in mm and DBH in cm, derived from a *P. tre*muloides stand within 10 km of our site [Ewers et al., 2002, 2007], from which heartwood area was calculated and then subtracted from $A_{\rm BT}$ to obtain sapwood area ($A_{\rm S}$) (cm²).

2.3. Scaling From Plots to the Stand

[10] Circular scaling plots (hereafter called plots) were generated by randomly sampling, with replacement, using plot centers (Easting, Northings) from a uniform distribution bounded within the domain (Figure 1). Plot selection was subject to three constraints: (1) no part of a plot was allowed to fall outside one of the 0.06 ha blocks, (2) every plot was required to have a minimum of one sap flux tree, and (3) every plot had to have at least three total trees. The second constraint ensured that every plot met the minimum sap flux sensor sample size employed in previous studies (Table 1). The third constraint ensured that no plot represented a single, isolated tree. These constraints were intended to mimic sap flux plot selection. We note that for small plots the latter two constraints would ignore gaps in the canopy and overestimate canopy density. However, such bias is not inconsistent with sap flux studies in which plots are selected randomly [e.g., Kang et al., 2002; Mellander et al., 2006], based on site characteristics [e.g., Hogg and Hurdle, 1997; Roberts et al., 2005], or using aggregate measures such as stand basal area or leaf area [e.g., Ewers et al., 2002; Simonin et al., 2006].

[11] To examine the effects of plot size on scaling $E_{\rm C}$ and $G_{\rm S}$ we generated 8 sets of 200 plots using uniform radii of 2 m, 2.5 m, 3 m, 4 m, 5 m, 7 m, 9 m, or 12 m, giving a range of plot areas from 12 to 450 m², respectively (Table 2). This bootstrapping method produced overlap of some plots, which meant that total variance declined and variance shifted from between plots to within plots with increasing plot size. In each plot we calculated L (m² leaf m⁻² ground area) by summing $A_{\rm L}$ of all *P. tremuloides* trees within the plot and dividing by plot area (A_{plot}) (m²). We note that these calculations include trees whose crowns extend beyond the plot boundary. This is necessary to ensure that total leaf area used to calculate $E_{\rm L}$ is consistent with the total sapwood area used to calculate $E_{\rm C}$. While it could lead to overestimates of L and canopy cover, particularly for small plots, these parameters would also be consistent with the plot total $E_{\rm C}$. The proportion of canopy cover (P_{CC}) was estimated by summing $A_{\rm C}$ for all trees in the plot, subtracting overlapping crown areas that were counted multiple times, and dividing by A_{plot} . Overlapping crown areas were quantified using a simple algorithm that estimates the area of the asymmetric

lens produced by the intersection of two circles [*Weisstein*, 2009].

[12] Canopy transpiration on a per unit ground area basis, $E_{\rm C} \,({\rm mm \ H_2O \ s^{-1}})$, was calculated following the approach of *Oren et al.* [1998b]:

$$E_{C} = \frac{1}{A_{plot}} \sum_{i=1}^{k} \sum_{j=1}^{l} J_{i,j} \cdot A_{S_{i,j}}$$
(1)

where k is the number of bins of sapwood area $(A_{\rm S})$ (m²), *l* is the number of individuals within each bin, and *J* is sap flux velocity $(m^3 s^{-1})$. To include nighttime transpiration, baselines were determined from the maximum temperature difference over a weeklong period using the approach of Oishi et al. [2008]. Radial and circumferential trend information used data from *Ewers et al.* [2002] to set up ratios between the outer xylem measurements and inner or southern xylem. Sapwood area bins were obtained by sorting from lowest to highest $A_{\rm S}$, all sap flux and the *P. tre*muloides not instrumented for sap flux trees (or nonflux trees) within the plot. Nonflux trees were then binned with flux trees with bins spanning from the midpoints between successive sap flux tree $A_{\rm S}$ values. In the degenerative case where a plot had only a single flux tree all trees in the plot were put in a single bin. We note that this method can lead to a systematic bias in plots with only one sap flux tree, if the chosen sap flux tree is always a dominant tree.

[13] First and second moments of the distribution of plot canopy transpiration were calculated at every half-hourly time step, with the mean and variance respectively calculated as

$$E[E_C(t)] = \frac{1}{n} \sum_{i=1}^{n} E_C(t)_i$$
(2)

and

$$\operatorname{VAR}[E_C(t)] = \frac{1}{n} \sum_{i=1}^{n} \left\{ E_C(t)_i - E[E_C(t)] \right\}^2$$
(3)

where t is the time (30 min step) and n is the number of plots. In general, relationships between variability and areal sample size follow some form of power law [Levin, 1992]. Power law functions yield straight lines on log-log plots, and so we plotted variance of $E_{\rm C}$ versus plot area with log-log axes.

[14] $G_{\rm S}$ was calculated by inverting a form of the Penman-Monteith equation [*Monteith and Unsworth*, 1990]:

$$G_S(t) = \frac{K_G[T(t)]}{D(t)} \frac{\mathrm{E}[E_C(t)]}{L}$$
(4)

where *D* is vapor pressure deficit (kPa), *L* is mean leaf area index (m² leaf m⁻² ground), and $K_G(T) = 115.8 + 0.4236T$ (kPa m³ kg⁻¹) is the conductance coefficient at average canopy temperature *T* (°C). $K_G(T)$ accounts for temperature effects on the psychrometric constant, latent heat of vaporization, specific heat of air at constant pressure, and density of air [*Phillips and Oren*, 1998].

2.4. Statistical Analysis and Evaluation of Goodness of Fit

[15] Curve fitting, regression analysis, ANOVAs, and *t* tests were performed in Sigmaplot (version 11.0 2008 Systat

Software, CA, United States). We used two criteria to evaluate the goodness of fit between individual plot $E_{\rm C}$ values and mean values for the whole stand. First, a regression analysis was conducted and the slope of the regression was used as a criterion for goodness of fit. In a subset of plots that had slopes between 0.97 and 1.03 we calculated the index of agreement (IOA) [*Willmott*, 1982] and sorted the results from highest (=1) to lowest (=0) IOA.

3. Results

3.1. Representativeness of Plots

[16] The actual number of sap flux trees per plot ranged from as low as 1 for the 13 m^2 plots to as high as 10 for some of the 452 m^2 plots (Table 2). The upper end of this range is comparable to most of the studies presented in Table 1, as 67 of the 90 studies had at most 10 sap flux trees per species per plot, and 48 had fewer than 8 trees per species per plot.

[17] $E_{\rm C}$ for any 12 m radius plot selected from block h (see Figure 1) was representative of stand $E_{\rm C}$, as indicated absolutely by a slope of regression of 0.999 and dynamically by IOA of 0.991 between the plot and whole stand fluxes evaluated using all 30 min values for the 7 days. Block h had mean canopy structural parameters ($A_{\rm B} = 5.7 \text{ m}^2 \text{ ha}^{-1}$, $P_{\rm CC}$ = 0.29, L = 1.24) that were similar to those for the whole stand at the 12 m radius scale ($A_{\rm B} = 6.3 \text{ m}^2 \text{ ha}^{-1}$, $P_{\rm CC} = 0.25, L = 1.48$). At the 12 m radius plot size the canopies in the other blocks were either denser or more open than the stand average with $A_{\rm B}$ ranging from 0.7 to 22.7 m² ha⁻¹. Using the same slope of regression and IOA criteria we also found representative plots at all sizes in different locations in the stand having similar values of L and P_{CC} as the stand. All plots shown in gray in Figure 1 had slopes between 0.97 and 1.03, and IOA values of 0.98 or higher. For instance, representative 7 and 9 m plots were located in the center of block i and representative 4 m plots were found centered at 5 m east, 50 m north in block g. No representative plots were found in areas of dense canopy, such as the west half of block k and west edge of block n, or in the low canopy density areas such as blocks a-d.

3.2. Representativeness and Environmental Drivers

[18] Figures 2–4 summarize the first two moments of $E_{\rm C}$ derived from the plots. Figure 2 shows scaled $E_{\rm C}$ for 2 m, 3 m and 12 m radius plots, along with environmental drivers. Although $E_{\rm C}$ values changed in an absolute sense as the size of the plot changed, the values showed the same dynamic responses to environmental drivers. To test whether or not this environmental driver affected the representativeness of plots we examined mean and variance of $E_{\rm C}$ in three bins of D representing low ($D \le 0.5$ kPa), intermediate (0.5 < $D \le 1.0$ kPa), and high levels (D >1.0 kPa). $E_{\rm C}$ declined monotonically with increasing plot size, and the rate of change of flux per unit increase in plot size also declined with increasing plot size (Figure 3a). Moreover, L and P_{CC} also declined monotonically (Table 2), and so $E_{\rm L}$ was less variable among plot sizes (Figure 3b). The mean responses were similar among bins of D. The $\log[VAR(E_C)]$ versus $\log(A_{plot})$ all fit straight lines (P < 0.0001) (Figure 4). The absolute $E_{\rm C}$ variances increased



Figure 2. Mean diurnal canopy transpiration (E_C) for 2, 3, and 12 m radius plots, along with diurnals for vapor pressure deficit (*D*) and incident photosynthetically active radiation (Q_0).





Figure 4. Logarithm of the variance of canopy transpira-

tion (E_C) versus the logarithm of plot area, binned into three ranges of vapor pressure deficit (*D*). The linear fits are based on y = y0 + ax, where y0 = -8.4635 and a = -0.9131 for low *D*, y0 = -7.5844 and a = -0.8423 for intermediate *D*, (A_{plot}), and y0 = -7.2952 and a = -0.8291 for high *D*. All regression lines are significant (P < 0.0001), and slopes are not significantly different (P > 0.39, R2 > 0.95).

Figure 3. (a) Mean canopy transpiration ($E_{\rm C}$) and (b) mean transpiration per unit leaf area ($E_{\rm L}$) versus plot area ($A_{\rm plot}$), into groups of low, intermediate, and high vapor pressure deficit (D).



Figure 5. Block mean (a) leaf area index (*L*) and (b) reference canopy average stomatal conductance (G_{Sref}) versus canopy closure (P_{CC}). A linear equation ($L = -0.40 + 6.94 P_{\text{CC}}$) is fit for *L* versus P_{CC} ($r^2 = 0.86$, P < 0.0001), while an exponential rise to a maximum { $G_{\text{Sref}} = 96.9[1 - \exp(-9.36P_{\text{CC}})]$ } is fit through G_{Sref} versus P_{CC} ($r^2 = 0.61$, P = 0.0016).

with *D*, but the rate of change of $\log[VAR(E_C)]$ with log (A_{plot}) was independent of *D* (P > 0.39).

3.3. Representativeness of Parameters for Ecosystem Models

[19] G_{Sref} and *m* are given in Table 2. G_{Sref} varied from 77.7 to 95.9 mmol m⁻² s⁻¹ and *m* varied in proportion to G_{Sref} , but there was no clear relationship between these parameters and plot size. However, across scales m was linearly related to G_{Sref} with a slope not statistically different from 0.6 (P = 0.367). Figure 5 shows L and G_{Sref} versus $P_{\rm CC}$ at the block level using one randomly chosen 12 m radius plot in each block. Blocks a and d (Figure 1) were excluded as their respective plots had near-zero $E_{\rm C}$, and so only 12 blocks were included in Figure 5. $P_{\rm CC}$ ranged from 0.03 to 0.58, with values less than 0.2 all occurring in forested wetland blocks. $A_{\rm B}$ ranged from 0.7 to $3.9 \text{ m}^2 \text{ ha}^{-1}$ in the wetland blocks and 4.3 to 22.7 m² ha⁻¹ in upland blocks. L varied from 0.13 to 1.03 m² m⁻² in the wetland and 0.76 to 4.39 m² m⁻² in the upland and was linearly related to $P_{\rm CC}$, as would be expected given that both parameters were derived allometrically from $A_{\rm BT}$. $G_{\rm Sref}$ also varied with $P_{\rm CC}$, albeit nonlinearly. Indeed, there was a threshold response in which G_{Sref} increased with P_{CC} in the wetland, but showed little or no variation with $P_{\rm CC}$ in upland blocks. Moreover, the nonwetland G_{Sref} values closely corresponded to the values reported for the whole stand using any plot size (Table 2).

4. Discussion

4.1. Plot Representativeness

[20] The representativeness of plots depended upon on whether total stand average canopy transpiration, $E_{\rm C}$, or transpiration per unit leaf area, $E_{\rm L}$, was considered. Although mean $E_{\rm C}$ decreased when going from the smallest to largest plot sizes, when scaled to $E_{\rm L}$ there was little variation among plot sizes. All plots were required to have at least one flux tree and two additional trees for scaling from sap flux to $E_{\rm C}$. This forced small plots to sample in relatively dense parts of the stand, which excluded gaps in the small plots. As plot size increased more gap areas were included. We note that the decline in $E_{\rm C}$, L, and $P_{\rm CC}$ would not occur if randomly selected plots were permitted to occur in gaps where there were no flux measurements. However, including plots with no flux trees would be an unrealistic representation of traditional sap flux plot sampling methods. Moreover, both mean and variance of $E_{\rm C}$ showed a monotonic decline with plot size, supporting a predictable scaling behavior even at the canopy scale. It should be noted that the scaling behavior for the variance (Figure 4) would be expected to change among studies as a function of the amount of spatial autocorrelation in $E_{\rm C}$. For example, a relatively homogeneous stand with respect to $E_{\rm C}$ would have less steep slopes of variance versus plot size. The slopes would be steepest in a stand where the individual fluxes were spatially independent. Alternatively, if one partitions a study area into nonoverlapping areal units then beyond the correlation length the variance is expected to fall off as the inverse of the number of areal units [Levin, 1992]. Within the correlation length recent evidence that spatial heterogeneity of $E_{\rm C}$ is a function of the rate of water loss, as expressed through changes in D [Adelman et al., 2008; Loranty et al., 2008; Traver et al., 2009], suggest that through plant hydraulics one can increase the predictability and interpretation of $E_{\rm C}$ variance between studies.

[21] Plot representativeness was also independent of the magnitude of environmental drivers, as the rate of change in variance of $E_{\rm C}$ with plot size did not change with *D*. We suggest that it was time-invariant stand structural characteristics (i.e., *L* and $P_{\rm CC}$), and by implication $A_{\rm B}$ and $A_{\rm S}$, not spatial dynamics of flux rates that was key to scaling fluxes to the stand and extracting parameters such as *L* and $G_{\rm Sref}$, as has been suggested by other studies [*Ford et al.*, 2007; *Adelman et al.*, 2008; *Loranty et al.*, 2008].

4.2. Random or Biased Plot Selection

[22] Given our plot sampling criteria, smaller plot sizes were more likely to systematically overestimate $E_{\rm C}$ than larger plots. Similarly, systematic error could potentially have occurred in any of the studies shown in Table 1 had they not chosen plots representative of the stand density. Our plot sizes (13 to 452 m²) spanned a range representing 40 percent of the sap flux studies presented in Table 1, and bounded all plot sizes of those studies conducted in *P. tremuloides* stands [*Hogg et al.*, 1997; *Hogg and Hurdle*, 1997; *Pataki et al.*, 2000; *Ewers et al.*, 2002; *Uddling et al.*, 2008]. The rate of change in mean stand values for $E_{\rm C}$, *L*, and $P_{\rm CC}$

declined with increasing plot size and was smallest for plots between 154 and 452 m². Randomly located plots at the 7-12 m radius plot sizes would therefore be expected to yield more representative canopy parameters than smaller plots. However, these plot areas are larger than those previously employed in sap flux studies conducted in P. tremuloides [Hogg et al., 1997; Hogg and Hurdle, 1997; Pataki et al., 2000; Ewers et al., 2002; Adelman et al., 2008], with the exception of a plantation study [Uddling et al., 2008]. Clearly either biased plot selection is preferred, our stand is anomalous, or the prior studies undersampled their respective stands. We considered the possibility that our stand was unusual in terms of structure or canopy transpiration. Evidence from the literature on P. tremuloides suggests that this is not the case. Mean basal area $(A_{\rm B})$ (m² ha⁻¹) of our 20 year old stand was 6.3 m² ha⁻¹, but varied from 0.7 to 22.7 m^2 ha⁻¹ among 12 m radius plots. This range bounded $A_{\rm B}$ values reported for 30 year old Boreal (3.4 m² ha⁻¹) [*Ewers et al.*, 2005] and subalpine Wyoming (4.9 $\text{m}^2 \text{ ha}^{-1}$) [Adelman et al., 2008] stands, but our basal areas were smaller than values reported in northern Wisconsin (30.0 m² ha^{-1}) [Ewers et al., 2002; Uddling et al., 2008], 20 year old Boreal (38.0 to 63.0 m² ha⁻¹) [Hogg and Hurdle, 1997], and another subalpine stand in Wyoming (43.0 m² ha⁻¹) [Pataki et al., 2000].

[23] The range of L, 0.13 to 4.39 m² m⁻² among blocks, spanned the range of values among prior sap flux studies conducted on P. tremuloides stands in Boreal [Hogg et al., 1997; Hogg and Hurdle, 1997; Ewers et al., 2005], subalpine [Pataki et al., 2000; Adelman et al., 2008], and northern temperate [Ewers et al., 2002, 2007; Uddling et al., 2008] biomes. Our mean stand L of 1.48 m² m⁻² using 12 m radius plots was similar to the values reported by Hogg and Hurdle [1997], less than half the values reported for stands with higher $A_{\rm B}$ [Pataki et al., 2000; Ewers et al., 2002; Uddling et al., 2008], and higher than the values reported in studies that had smaller $A_{\rm B}$ [Ewers et al., 2005; Adelman et al., 2008]. Thus, the allometry of our stand (i.e., relationship between L and $A_{\rm B}$) appears to be consistent with other P. tremuloides studies.

[24] The scale independence of $E_{\rm L}$ is important since ecosystem models typically require some parameter for $G_{\rm S}$ (or its reference parameter value, G_{Sref}), which is expressed on a per unit leaf area basis. While plot location had a large effect on L and P_{CC} , the effect was minimal for G_{Sref} . Plots located in the wetland ($P_{\rm CC} < 0.2$) had low and variable $G_{\rm Sref}$ values, while other plots showed a limited range of G_{Sref} . Moreover, we derived a proportionality between m and G_{Sref} of 0.59, which is consistent with previous studies involving P. tremuloides [Hogg et al., 2000; Ewers et al., 2005, 2007; Uddling et al., 2008] and other species [Oren et al., 1999; Ewers et al., 2000; Ogle and Reynolds, 2002; Mackay et al., 2003; Addington et al., 2004; Fisher et al., 2006; David et al., 2007; Ewers et al., 2007]. Our estimates of G_{Sref} varied between 77.7 and 95.9 mmol m⁻² s⁻¹ among plot sizes, which were somewhat higher than the maximum value of 65 mmol $m^{-2} s^{-1}$ reported by *Ewers et al.* [2005], but smaller than the values reported in other P. tremuloides studies [Hogg and Hurdle, 1997; Ewers et al., 2007]. Moreover, the range of G_{Sref} (77.7 to 95.9 mmol $m^{-2} s^{-1}$) is quite small compared to intertree variability of G_{Sref} attributed to competition for light at our study site

[25] If our stand is not an anomaly then the more likely alternative explanation for the apparent effectiveness of relatively smaller plot sizes employed in previous P. tremuloides studies is that they relied on biased selection of plot locations. This is clearly evidenced in the earlier Boreal studies [Hogg et al., 1997; Hogg and Hurdle, 1997] in which plots were chosen to distinguish among clones, or plantation plots [Uddling et al., 2008] that had experimental control on factors affecting growth rates. *Pataki et al.* [2000] chose 3 m radius plots centered on each sap flux tree on the assumption that these plots contained all the flux trees' competitors. Ewers et al. [2002, 2007] measured L optically and allometrically in an 804 m² stand. A 79 m² sap flux plot within the stand was chosen to closely match the mean $L (=3.5 \text{ m}^2 \text{ m}^{-2})$ and to capture the range of DBH within the stand. Adelman et al. [2008] and Loranty et al. [2008] used initial surveys of $A_{\rm B}$ to determine the spatial extent and sample design for their plots. The logic behind this was that sapwood area would be related to $A_{\rm B}$, and $E_{\rm C}$ would scale with sapwood area. The unit lag distance needed to adequately sample $A_{\rm B}$ at a stand scale while providing sufficient information to quantify spatial autocorrelation of tree transpiration determined the respective plot sizes. Our results from selecting the best fit plots at each of the radii, and these other P. tremuloides studies, support a scaling logic that exploits stand structural properties to find representative plots.

4.3. Implications for Scaling From Trees to Stands

[26] $E_{\rm C}$ scaling logic that relies on characterizing stand structure in space with relatively few flux measurement points is also generally supported by the literature spanning a range of species composition and biomes. Nonetheless, multiple sources of spatial variability in $E_{\rm C}$ have been implicated. Intertree variations in $J_{\rm S}$ have been cited as primary [Kumagai et al., 2005a; Pataki et al., 2005; Hultine et al., 2007; Nadezhdina et al., 2007] or secondary [Bovard et al., 2005; Herbst et al., 2007; Adelman et al., 2008] sources of heterogeneity in canopy transpiration. We cannot rule out the potential for spatial variability in $J_{\rm S}$ in our own study, but such heterogeneity was not expressed in such a way that it prevented us from scaling up from any plot size. Moreover, Loranty et al. [2008], using data from the same stand a year earlier, found no clear evidence of spatial autocorrelation in $J_{\rm S}$. Oren et al. [1998a, 1998b] used a coefficient of variation of 20% as a guideline and said that the sample size thus changes with stand characteristics. Alternatively, Kumagai et al. [2005a] found up to a threefold difference in sap flux velocity among trees, and so he argued that sap flux measurements should be made in as many trees per species as possible to accurately quantify scaled $E_{\rm C}$. On the one hand, if such a requirement were indeed justified it could make scaling from a practical number of sap flux measurements to regional levels questionable. On the other hand, Ford et al. [2007] found that variations in sapwood area among plots was the greatest source of variability in canopy transpiration despite the relative homogeneity in species and age of their stand. The need for a large number of sap flux sensors per species was unwarranted for our stand because (1) single plots were

representative of stand mean $E_{\rm C}$, (2) no single plot at any size had more than 10 sap flux trees, (3) G_{Sref} was insensitive to $P_{\rm CC}$ and, by implication, plot location in the upland forest areas of the stand, and (4) G_{Sref} was predictable from $P_{\rm CC}$ in the forested wetland areas of the stand. We suppose, based on Figure 5b, that three plots representing wetland, upland, and transitional between these end-members, with at most 10 sap flux trees per plot, would be sufficient for parameterizing models where each of these components of the stand were needed. Consequently, our analysis suggests a representative number of sap flux trees for our stand was somewhere between 3 and 30. However, to move toward the low end of this range, a tractable number of $J_{\rm S}$ measurements for studies requiring many species, requires mechanistic explanations for transpiration variability in space and time.

[27] Potential mechanistic explanations of such spatial heterogeneity implicate edaphic and topographic controls on vertical and lateral water flow, respectively, as well as stand structure. A number of studies have shown that spatial variability of transpiration attributed to soil moisture controls was predictable from edaphic or topographic properties [Schiller et al., 2002; Eberbach and Burrows, 2006; Trompvan Meerveld and McDonnell, 2006; Nadezhdina et al., 2007]. Tromp-van Meerveld and McDonnell [2006] showed that topographic and edaphic controls affected growth rates of individual trees, which in turn modified the spatial distribution of transpiration. In northern Wisconsin, Traver et al. [2009] showed no impact of edaphic conditions, while Loranty et al. [2010, also submitted manuscript, 2010] showed that competition for light explained a significant amount of spatial variation of $E_{\rm C}$ and $G_{\rm Sref}$. Higher flux rates have also been shown near stand edges in comparison to stand centers [Taylor et al., 2001; Cienciala et al., 2002; Giambelluca et al., 2003; Herbst et al., 2007]. Taylor et al. [2001] attributed higher transpiration rates in edge trees to advection, although considerable velocity variance along forest edges suggests they cannot be treated simply as edges [Detto et al., 2008]. Similarly, evidence of stand structure as a control on the variability of $E_{\rm C}$ among stands was shown when stand thinning led to increased soil moisture that in turn promoted greater transpiration during periods of intermediate soil moisture [Simonin et al., 2006, 2007]. Alternatively, some stand structural changes associated with age [Zimmermann et al., 2000; Phillips et al., 2002; Ewers et al., 2005], such as adjustments in leaf-to-sapwood area [Phillips et al., 2002] and root-shoot ratio [Ewers et al., 2005] appear to be more elusive qualities to obtain routinely.

[28] Most studies that account for spatial heterogeneity of $E_{\rm C}$ at the stand level employ a logic that is based on the scaling theory of *Hatton and Wu* [1995] relating tree water use to leaf area index. Numerous studies either use this approach explicitly [*Vertessy et al.*, 1997; *Medhurst et al.*, 2002; *Phillips et al.*, 2002; *Schiller et al.*, 2002] or modify it by using sapwood area as the primary scalar of $E_{\rm C}$ [*Vertessy et al.*, 1995; *Oren et al.*, 1998b; *Bovard et al.*, 2005; *Ford et al.*, 2007; *Herbst et al.*, 2007; *Kumagai et al.*, 2007]. However, this simplified scaling logic loses credibility when different scalars yield different flux values, as has been suggested by some studies [*Vertessy et al.*, 1997; *Ford et al.*, 2007]. This would seem distressing, as scalars such as sapwood area, leaf area, and stem diameter should be correlated

through allometric relationships. Our results and numerous others [Taylor et al., 2001; Cienciala et al., 2002; Giambelluca et al., 2003; Tromp-van Meerveld and McDonnell, 2006; Kumagai et al., 2007; Adelman et al., 2008] suggest that this problem would be avoided if plots were located in structurally representative areas within the stand, and not simply defined to include a wide range of DBH. So, this study and others suggest that to scale flux observations from trees to stand we should take into account the spatial arrangement of DBH and not just its moments. Our study differed from others in that we measured and directly analyzed both sap flux and transpiration scalars spatially with high density within the stand. Previous studies either examined just the sap flux trees using aggregate measures of spatial variability [e.g., Adelman et al., 2008; Loranty et al., 2008; Traver et al., 2009] or had representative measures of the scalars and a limited number of sap flux sensors [e.g., Hogg and Hurdle, 1997; Vertessy et al., 1997; Oren et al., 1998b; Wilson et al., 2001; Ewers et al., 2002; Giambelluca et al., 2003; Ewers et al., 2005; Simonin et al., 2006]. By employing a high density and spatially extensive set of both flux and scalar measurements in a heterogeneous stand our study provides a robust demonstration of how scaling transpiration from plots to stands should work. This study shows quantitatively, as other studies have suggested, that biased sampling would be preferred over random sampling for characterizing stand level water fluxes. We suggest that future scaling efforts take into consideration three spatially covarying parameters, G_{Sref} , L, and P_{CC} , which are needed to accurately scale canopy transpiration to the whole stand level or to larger spatial extents.

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