

Review

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Commonality and variability in the structural attributes of moist temperate old-growth forests: A global review

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ABSTRACT

Temperate forests have been fundamentally altered by land use and other stressors globally; these have reduced the abundance of primary and old-growth forests in particular. Despite many regional studies, the literature lacks a global synthesis of temperate old-growth structural characteristics. In this study we compare literature derived data on mature and old-growth moist temperate forests with the aim of: (i) exploring global commonalities; (ii) investigating sources of variability among systems; and (iii) highlighting data gaps and research needs. We compiled a dataset of 147 records from 93 papers, and analyzed a set of structural indicators: basal area, stem density, large living trees, live aboveground biomass, quadratic mean diameter, and coarse woody debris volume. These indicators were contrasted between mature and old-growth age classes at a global level and across continents and broad forest types, testing for significance through Monte-Carlo permutation procedure. We also related structural indicators to age, climatic and geographical descriptors. Our results suggest that all structural indicators vary across systems in relation to geographical, compositional, and climatic influences. However oldgrowth forests showed global commonalities in structure when compared to mature forests: significantly higher densities of large living trees, higher quadratic mean diameter, and higher amounts of live aboveground biomass and coarse woody debris. Furthermore we found inconsistency in the structural variables reported by different papers; lack of studies on temperate forests in Russia, and Western and Central Asia. The findings improve our understanding of old-growth structure and function, and will help inform sustainable forest management and conservation approaches world-wide.

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Forest Ecology and Managemer

Contents

1	ntroduction	159
1. ว	Methoda	467
Ζ.	Methods	407
	2.1. Literature search	467
	2.2. Data extraction	467
	2.3. Statistical analysis	468
3.	Results	468
	3.1. Data distribution across continents, forest types and age classes	468
	3.2. Comparison between mature and old-growth forests	469
	3.3. Correlation between structural attributes and site characteristics	469
4.	Discussion	469
	4.1. Global commonalities	470
	4.2. Sources of variability	472
	4.3. Comparison against structural variability in fire-prone late-successional forests.	475
	4.4. Incomplete data and research needs	475
5.	Conclusions	475

Abbreviations: LLT, large living trees density; LAB, live aboveground biomass; QMD, quadratic mean diameter; CWD, coarse woody debris volume. * Corresponding author.

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5.1.	Management implications	/5
Acknow	vledgements	/6
Appen	lix Ā. and B. Supplementary material	<i>'</i> 6
Referen	nces	/6

1. Introduction

Temperate forests are generally more altered and reduced in extent globally than other forested biomes (Silander, 2001) due to their distribution at mid-latitudes with moderate climates, which are also some of the most heavily populated and developed regions of the world. About 35% of the world forests are considered primary (i.e. never cleared though subject to natural disturbances) but these mostly occur in tropical and boreal zones (FAO, 2010); according to one estimate less than 1% of the northern hemisphere's temperate broad-leaved forests remain in a primary condition (Silander, 2001).

Landscapes dominated by primary forests are comprised of complex patch mosaics and host a full range of stand development stages, from disturbance and cohort establishment to old-growth (Franklin et al., 2002). The latter is a late stage of stand development that typically does not occur or is rare in managed forest systems, and has been defined by many authors based on a combination of structural features, age, and sometimes human disturbance history (Wirth et al., 2009). The 'structural' approach, first developed through studies performed in the Pacific Northwest (Spies and Franklin, 1991), is useful because it can both be a good proxy of ecosystem function and is a readily measured surrogate for habitat of many taxa difficult to sample (Franklin et al., 2002; Burrascano et al., 2011).

Structural indicators, often combined with ecological and compositional information, have been proposed for temperate forests in North America (Franklin et al., 1981; Keddy and Drummond, 1996), Europe (Peterken, 1996; Nilsson et al., 2002), South America (Gutierrez et al., 2004) and China (Chen and Bradshaw, 1999). Nevertheless old-growth forests may display strong variation in their structure in different forest types (e.g. Jones, 1945; Burgman, 1996; Wells et al., 1998).

Such variation derives from the fact that the old-growth condition should be defined not only on the basis of a set of structures providing desired functions, such as habitat for late-successional biodiversity, but also of the developmental processes that produce those structures. However the lack of a global theoretical framework of temperate forest development has hindered broad application of a widely accepted process-based approach. Previous research on forest dynamics suggests considerable uncertainty in segmenting stand structural development, because many developmental processes operate continuously or episodically, often rendering discretely defined stages arbitrary (Franklin et al., 2002; Spies, 2004). Moreover, stand development may undergo several different pathways in relation to disturbance regimes and environmental controls (Donato et al., 2012). Furthermore, the age threshold to define developmental stages will differ by forest type and geographic region due to differences in stand development rates, tree longevity, disturbance regime and other factors related to these, such as climate and site conditions (Franklin et al., 1987; Motta et al., 2010). Recognizing these limitations, our review compares "mature" stands with stands classified as "old-growth". These largely age-based classes, commonly reported in the literature, are employed as surrogates for the developmental processes described in Franklin et al. (2002) and other models (e.g. Oliver and Larson, 1990). They generally correspond with the "maturation" (i.e. mature) and "vertical/horizontal diversification and pioneer cohort loss" (i.e. old-growth) stages proposed by Franklin et al. (2002).

We focus on late-successional stands since most temperate forests are either managed for wood production or reflect the longterm influence of human activities. As a consequence, old-growth forest structural features are underrepresented in most contemporary landscapes (Bauhus et al., 2009; Rhemtulla et al., 2009) and biodiversity provisioning and other functions associated with them are frequently impaired (Siitonen, 2001; Hatanaka et al., 2011). Though some extended rotation systems may approximate oldgrowth structural conditions (Keeton, 2006; Bauhus et al., 2009), most silvicultural regimes narrow the range of possible developmental processes as well as the type and spatial distribution of structures, since they do not allow the stands to develop past the maturation phase or they selectively remove certain structures, like large or dead trees (Hunter, 1999).

Our prediction was that moist temperate forests globally will share distinct differences when comparing mature to old-growth conditions, because stand structural heterogeneity (vertical and horizontal) is presumed to increase as late-successional stands develop. For example, coming out of the maturation phase a marked increase in coarse woody debris (CWD) volumes can occur (Harmon et al., 1986), and re-establishment of shade-tolerant trees in the understory initiates redevelopment of stand vertical complexity (Keeton and Franklin, 2005); while horizontal heterogeneity is primarily created by the shift from density-dependent to density-independent mortality (Franklin et al., 2002). These processes drive development of many structural characteristics associated with old-growth. Our review evaluates indirectly through a focus on structural indicators - whether these are universally operative or whether there is significant variation among systems.

Comparison between mature and old-growth forests allows identification of structures that could be promoted silviculturally in managed stands. There are a number of reasons why managers might be interested in this objective. For example, old-growth forests are known to host high plant (Aude and Lawesson, 1998; Burrascano et al., 2009), fungi (Odor et al., 2006; Persiani et al., 2010) and animal diversity (Mikusinski and Angelstam, 1998; McKenny et al., 2006; Winter and Moller, 2008). Recent studies have also shown the importance of old-growth forests in storing high quantities of carbon both as aboveground biomass (Keith et al., 2009; Keeton et al., 2010) and in soils (Zhou et al., 2006) and their ability, in some cases, to maintain positive Net Primary Productivity very late into stand development (Field and Kaduk, 2004; Luyssaert et al., 2008; Xu et al., 2012).

Based on the services they provide, old-growth forests represent an important reference point for evaluating human impacts on forest ecosystems, and for understanding forest development processes through observation of the temporal and spatial interactions between successional processes, disturbance events, and tree mortality (Peterken, 1996; Keeton, 2006; Rhemtulla et al., 2009; Hoover et al., 2012). Studying old-growth forests has long been considered the basis for developing natural disturbance-based (North and Keeton, 2008), 'close-to-nature' (Commarmot et al., 2005; Heiri et al., 2009) or 'natural dynamics' (von Oheimb et al., 2005) silvicultural systems able to emulate natural processes and fulfill socio-economic goals while maintaining a full range of

459

Table 1

Structural data and site descriptors of the stands included in the review. Age class: M – mature, OG – Old-growth. Forest types: C – conifer, DB – deciduous broadleaf, EB – evergreen broadleaf, C/DB – mixed conifer/deciduous broadleaf, EB/C – mixed evergreen broadleaf/conifer. Age values marked with asterisks represent time since last disturbance and were excluded from the analysis. LLT data marked with \dagger were approximated from stand diameter distributions, Live Aboveground Biomass values marked with \ddagger refer to stem biomass only. An asterisk (*) before the reference indicates that we averaged multiple stands data in a single record.

Continent	Region	Altitude (m a.s.l.)	Mean T (°C)	Mean P (mm)	Forest type	Dominant tree taxa	Age Class	Age	Basal Area (m² ha ⁻¹)	Stem density (n ha-1)	Large living trees (n ha ⁻¹)	Live Aboveground Biomass (Mg ha ⁻¹)	Quadratic Mean Diamater (cm)	Coarse Woody Debris (m ³ ha ⁻¹)	Reference
Asia	Iran	1475	8.0	1800	DB	Fagus orientalis	М		44.5	336	70^{\dagger}		41.1		Akhavan et al.
	Iran	1475	8.0	1800	DB	Fagus orientalis	OG		39.8	302	68^{\dagger}		40.9		(2012) Akhavan et al. (2012)
	China - Changhaishan	1600			С	Picea jezoensis, Abies nenkrolenis	OG		50.2	1316			22.0		Chen and Bradshaw (1999)
	China - Changbaishan	800	2.8	700	C/DB	Pinus koraiensis, Tilia amurensis, Quercus mongolica	OG	300	43.2	299	64^{\dagger}		42.9		Hao et al. (2007)
	China - Changbaishan	1300	4.8	612	DB	Quercus liaotungensis, Betula dahurica	М	100	15.6	729			16.5		Hou et al. (2004)
	China - Changbaishan	740	3.3	672	C/DB	Pinus koraiensis, Fraxinus mandshurica,Tilia amurensis	OG	>200	46.5	515			33.9		Dai et al. (2011)
	China - Changbaishan	1680	3.3	672	С	Picea jezoensis, Abies nenhrolenis	OG	>200	35.2	745			24.5		Dai et al. (2011)
	China - Sichuan	2930	2.3	1100	С	Abies faxoniana, Picea purpurea, Sabina saltuaria	OG	>250	45.2	404	83 [†]		37.7		*Taylor et al. (2006)
	Japan - Hokkaido		3.1	1652	DB	Quercus mongolica	OG	110		804	16^{\dagger}				Sano (1997)
	Japan - Hokkaido	245	5.7	1043	C/DB	Abies sachalinensis, Acer palmatum, Quercus mongolica	OG	>300	51.2	958			26.1		Abrams et al. (1999)
	Japan - Honshu	2200	0.0	1750	C/DB	Abies mariesii, A. veitchii, Picea jezoensis, Tsuga diversifolia	OG	>200	69.4	496.9			42.2		*Narukawa and Yamamoto (2001)
	Japan - Honshu	1319	7.8	2500	С	Chamaecyparis obtusa, C. pisifera, Thujopsis dolabrata	OG	300	64.8	1618.5			22.6		Hoshino et al. (2001)
	Japan - Honshu	900	11.0	3450	DB	Fagus crenata, Quercus mongolica, Betula grossa, Acer mono	М	105	39.9	536			30.8		*Yamamoto and Nishimura (1999)
	Japan - Honshu	650	11.0	3450	DB	Fagus crenata, Quercus mongolica, Betula grossa. Acer mono	OG	140	58.4	281.3			51.4		*Yamamoto and Nishimura (1999)
	Japan - Honshu	1100	11.0	3450	DB	Fagus crenata	OG		34.3	541			28.4		Yamamoto et al. (1995)
	Japan - Kyushu	460	14.2	3070	EB	Dystilium racemosum, Persea thunbergii, Castanopsis cuspidata	OG	>100*		696		315.7			Sato (2010)
	Japan - Kyushu	490	14.9	2370	EB	Dystilium racemosum, Persea thunbergii, Castanopsis cuspidata	OG	>100*		564		254.3			Sato (2010)
	Japan - Tsushima	340	15.3	2140	EB	Castanopsis cuspidata, Dystilium racemosum	OG		63.9	4570			13.3		Manabe et al. (2000)
	Japan - Satsunan islands	550			EB	Castanopsis cuspidata, Dystilium racemosum, Quercus salicina	М	41-60	54.7	2805		258.0	15.8		Aiba et al. (2001)

Continent	Region	Altitude (m a.s.l.)	Mean T (°C)	Mean P (mm)	Forest type	Dominant tree taxa	Age Class	Age	Basal Area (m² ha ⁻¹)	Stem density (n ha-1)	Large living trees (n ha ⁻¹)	Live Aboveground Biomass (Mg ha ⁻¹)	Quadratic Mean Diamater (cm)	Coarse Woody Debris (m ³ ha ⁻¹)	Reference
	Japan - Satsunan islands	550			EB	Castanopsis cuspidata, Dystilium racemosum, Quercus salicina	OG	>150*	61.5	1795		279.0	20.9		Aiba et al. (2001)
Australia	Australia (Eastern)	850		2000	EB	Nothofagus moorei, Eucalyptus sp.pl.	OG		69.2						*Bale et al. (1998)
	New Zealand	100	10.9	4500	EB/C	Nothofagus menziesii, Weinmannia racemosa, Dacrydium cupressinum	OG	>800*	55.9	445	55 [†]		40.0		Lusk and Smith (1998)
	New Zealand	550	11.0	2000	EB/C	Dacrydium cupressinum, Weinmannia racemosa	М	252	144.5	6015			17.5		Smale et al. (1997)
	New Zealand	50	11.3	3400	EB/C	Dacrydium cupressinum, Weinmannia racemosa	М	197	61.0	1660			21.6		Walcroft et al. (2005)
	New Zealand	1136	8.0	1447	EB	Nothofagus solandri	М	>150	68.2	1216		240.4	26.7		Davis et al. (2003)
Europe	Czech Republic	1220	4.0	1400	С	Picea abies	OG	300	32.0	131	80^{\dagger}		55.8	311	Svoboda and Pouska (2008)
	Czech Republic	780	4.3	875	C/DB	Fagus sylvatica, Picea abies, Abies alba	OG	169*	40.6	205	76		50.3	201.9	Král et al. (2010)
	Czech Republic	765	5.4	1144	C/DB	Fagus sylvatica, Picea abies, Abies alba	OG	57*	38.5	306	68		40.0	149.8	Král et al. (2010)
	Czech Republic	770	6.1	740	C/DB	Fagus sylvatica, Picea abies, Abies alba	OG	66*	26.5	124	47		52.2	300.8	Král et al. (2010)
	France - Alps	1425	5.8	1138	C/DB	Fagus sylvatica, Abies alba	М	109	42.2	748			26.8		Marage and Lemperiere (2005)
	France - Alps	1425	5.8	1138	C/DB	Fagus sylvatica, Abies alba	OG	141	27.1	558			24.9		Marage and Lemperiere (2005)
	France - Fontainebleau	140	10.2	897	DB	Fagus sylvatica, Quercus petraea	OG	100-300	24.4	445			26.4	214.2	Wijdeven (2004)
	France - Vosges	200		850	DB	Fagus sylvatica, Quercus petraea	М	100-150	54.0	320			46.4	91.4	Schnitzler and Borlea (1998)
	Germany (NE)	100	7.8	593	DB	Fagus sylvatica	OG	200-230	32.7	263	75.1		39.8	94	Von Oheimb et al. (2005)
	Poland - Bialowietza				C/DB	Picea abies, Carpinus betulus, Tilia cordata	OG		36.3	665			26.4		Bernadzki et al. (1998)
	Poland - Carpathians	1300	3.0	1635	C	Picea abies	OG		41.2†	314	69 [†]		40.9	192	Zielonka (2006)
	Poland - Carpathians	900	4.5	1050	C/DB	Fagus sylvatica, Abies alba	OG		36.3		73 ⁺			383	Paluch (2007)
	Slovakia	890	5.0	1000	C/DB	Fagus sylvatica, Abies alba, Picea abies	OG		41.9	255	81'		45.8	306	Holeksa et al. (2009)
	Slovakia	1350	2.8	1105	C	Picea abies	OG		41.0	290			42.4	144	Holeksa et al. (2007)
	Slovakia	878	6.0	975	DB	Fagus sylvatica	OG		37.4	303	*		39.6	469	Kucbel et al. (2012)
	Switzerland Switzerland	470 500	8.9 8.9	976 910	DB DB	Fagus sylvatica Fagus sylvatica, Quercus	M M	70* 70*	39.0 40.8	422 400	15.5' 24.2 [†]		34.3 36.1		Heiri et al. (2009) Heiri et al. (2009)
	Switzerland	650	7.5	1250	DB	Fagus sylvatica	М	150	30.7	259	83^{\dagger}		38.9	8	Commarmot et al.
	Ukraine -	750	7.0	950	DB	Fagus sylvatica	OG		38.5	219	81†		47.3	111	Commarmot et al.

Table 1 (continued))														
Continent	Region	Altitude (m a.s.l.)	Mean T (°C)	Mean P (mm)	Forest type	Dominant tree taxa	Age Class	Age	Basal Area (m² ha ⁻¹)	Stem density (n ha-1)	Large living trees (n ha ⁻¹)	Live Aboveground Biomass (Mg ha ⁻¹)	Quadratic Mean Diamater (cm)	Coarse Woody Debris (m ³ ha ⁻¹)	Reference
	Carpathians														(2005)
	Ukraine - Carpathians	1035	7.0	790	С	Picea abies, Abies alba	М	100-150	45.3	1541	4.2	181.3	19.3	77	Keeton et al. (2010)
	Ukraine - Carpathians	1035	7.0	790	С	Picea abies, Abies alba	OG	250	47.7	725	59.8	255.5	31.9	176	Keeton et al. (2010)
	Ukraine - Carnathians	1035	7.0	790	С	Picea abies, Abies alba	OG	250	54.0	339	93.1	303.0	38.3	157	Keeton et al.
	Romania	1250	4.9	755	С	Picea abies	OG		42.1	348			39.2		Lamedica et al.
	Romania	485	8.5	838	DB	Fagus sylvatica	OG		37.3	499	72.2^{\dagger}		30.9	75	Petritan et al.
	Romania	485	8.5	838	DB	Fagus sylvatica, Quercus	OG		38.3	487	80.6^{\dagger}		31.6	135	Petritan et al.
	Romania	485	8.5	838	DB	Quercus petraea	OG		39.5	713	120.4^{\dagger}		26.6	134	(2012) Petritan et al.
	Bosnia-	1386	7.8	1600	C/DB	Fagus sylvatica, Abies	OG		47.1	489	117.0		35.0	327	(2012) Motta et al. (2011)
	Herzegovina Bosnia-	1150	5.9	1837	C/DB	alba, Picea abies Fagus sylvatica, Abies	OG		56.8	466			39.4		*Nagel et al.
	Herzegovina Italy - Apennines	1725	10.6	1035	DB	alba Fagus sylvatica	М		39.9	2182	22.0		15.3	12	(2010) Burrascano et al.
	Italy - Apennines	1725	10.6	1035	DB	Fagus sylvatica	OG		30.7	1835	71.0		14.6	45	(2008) Burrascano et al.
	Italy - Apennines	975	9	1750	C/DB	Fagus sylvatica, Abies	OG	>140	57.4	177	122^{\dagger}		64.3	65.2	(2008) Travaglini et al.
	Italy - Alps	1695			С	alba Picea abies	М	200	63.3	466			41.6	28	(2012) Motta et al. (2010)
	Italy - Alps	1815			С	Picea abies	М	220	69.2	513			41.5	80	Motta et al. (2010)
	Slovenia	900	8.2	1400	C/DB	Fagus sylvatica, Abies alba	OG		41.6	341	119†		39.4		Nagel et al. (2006)
	Slovenia	905	8.3	1500	C/DB	Fagus sylvatica, Abies alba	OG			265				138	Boncina (2000)
	Slovenia	905	8.3	1500	C/DB	Fagus sylvatica, Abies alba	М			393				10	Boncina (2000)
	Spain (NW)	140	14.0	1200	DB	Fagus sylvatica, Quercus robur	OG	220	31.9	1112	36.5^{\dagger}		19.1		*Rozas (2003)
	Spain (NW)	780	9.8	1600	DB	Fagus sylvatica	OG		48.0	569	54^{\dagger}	410.0	32.8		Merino et al. (2007)
North America	US - Michigan				C/DB	Acer saccharum, Betula alleghaniensis, Tsuga canadensis	OG	200-300	33.0	1130	60 [†]	262.0	19.3		Fisk et al. (2002)
	US - Michigan				C/DB	Acer saccharum, Betula alleghaniensis, Tsuga	М	60-80	31.0	970	10^{\dagger}	230.0	20.2		Fisk et al. (2002)
	US - Michigan	390	5.5	935	C/DB	Acer saccharum, Betula alleghaniensis, Tsuga	OG		45.6				30.4		*Dahir and Lorimer (1996)
	US - Michigan	390	5.5	935	C/DB	Acer saccharum, Betula alleghaniensis, Tsuga	М	100-175	49.4				35.2		*Dahir and Lorimer (1996)
	US - Michigan	525	3.2	850	DB	Acer saccharum	М	65-75	32.3	828			22.3	39	Goodburn and
	US - Michigan	525	3.2	850	C/DB	Acer saccharum, Betula alleghaniensis, Tsuga	М		28.0	469			27.6	61	Goodburn and Lorimer (1998)

462

Continent	Region	Altitude (m a.s.l.)	Mean T (°C)	Mean P (mm)	Forest type	Dominant tree taxa	Age Class	Age	Basal Area (m² ha ⁻¹)	Stem density (n ha-1)	Large living trees (n ha ⁻¹)	Live Aboveground Biomass (Mg ha ⁻¹)	Quadratic Mean Diamater (cm)	Coarse Woody Debris (m ³ ha ⁻¹)	Reference
						canadensis									
	US - Michigan/ Wisconsin	525	3.2	850	DB	Acer saccharum	М		25.5	446			27.0	74	Goodburn and Lorimer (1998)
	US - Michigan/ Wisconsin	525	3.2	850	DB	Acer saccharum	OG	~350	34.4	313			37.4	127	Goodburn and Lorimer (1998)
	US - Michigan/ Wisconsin	525	3.2	850	C/DB	Acer saccharum, Betula alleghaniensis, Tsuga canadensis	OG	~350	37.5	388			35.1	151	Goodburn and Lorimer (1998)
	US - Michigan/ Wisconsin	1940		1100	С	Abies magnifica, Abies concolor	OG	>200	85.2	825	125		22.1		*Taylor and Halpern (1991)
	US - Michigan/ Wisconsin				C/DB	Tsuga canadensis, Betula alleghaniensis, Acer	OG	185	48.6		2.5			84.5	*Tyrrell and Crow (1994a,b)
	US - Michigan/ Wisconsin				C/DB	Tsuga canadensis, Betula alleghaniensis, Acer	OG	231	49.9		16.2			104.2	*Tyrrell and Crow (1994a,b)
	US - Michigan/ Wisconsin				C/DB	Tsuga canadensis, Betula alleghaniensis, Acer	OG	261	46.7		35.5			121.3	*Tyrrell and Crow (1994a,b)
	US - Michigan/ Wisconsin				C/DB	saccharum Tsuga canadensis, Betula alleghaniensis, Acer saccharum	OG	336	43.3		45			151.3	*Tyrrell and Crow (1994a,b)
	US- Indiana	295			DB	Quercus rubra, Fraxinus americana, Q. alba, Q. macrocarpa	OG	>350	31	321	199		35.1		Parker et al. (1985)
	Canada - Quebec		6.1	1102	DB	Acer saccharum, Fagus grandifolia	OG	>300*	29.9	382			31.6		Beaudet et al. (2007)
	US - Massachussets	466		1120	DB	Fagus grandifolia, Quercus rubra, Acer sp pl	OG	>250	35.0	509	41^{\dagger}		29.6		*Orwig et al. (2001)
	US - Massachussets	473			C/DB	Tsuga canadensis, Fagus grandifolia, Betula sp.pl.	OG	177-246	44.8	531	58^{\dagger}		32.8	156.4	*D'Amato et al. (2008)
	US - Massachussets	473			C/DB	Tsuga canadensis, Fagus grandifolia, Betula sp.pl.	М	108-136	41.2	1000	20^{\dagger}		22.9	43.90	*D'Amato et al. (2008)
	US - New York				C/DB	Betula alleghaniensis, Fagus grandifolia, Picea rubens, Tsuga canadensis	OG	319	33.0	1005	37	221.0	20.5	206.8	*Keeton et al. (2007)
	US - New York				C/DB	Betula alleghaniensis, Fagus grandifolia, Picea rubens, Tsuga canadensis	Μ	118	26.0	981	12	148.0	18.4	177.8	*Keeton et al. (2007)
	US - New York				C/DB	Betula alleghaniensis, Fagus grandifolia, Picea rubens, Tsuga canadensis	Μ	119	24.0	777	11	144.0	19.8	110.4	*Keeton et al. (2007)
	US - New York	595			DB	Acer saccharum, Fagus grandifolia	OG	149-259	33.7	392	65^{\dagger}		33.1		McGee et al. (1999)
	US - New York	630			DB	Acer saccharum, Fagus grandifolia	М	90-100	29.1	508	17^{\dagger}		27.0		McGee et al. (1999)
	US - Vermont	481	7.6	993	DB	Acer saccharum, Fraxinus americana	OG	150-400	34.5	97		278	28.7		*Hoover et al. (2012)
	US - Vermont	464	7.6	993	С	Tsuga canadensis, Picea	OG	150-400	35.7	103		222	22.4		*Hoover et al.

Continent	Region	Altitude	Mean	Mean P	Forest	Dominant tree taxa	Age	Age	Basal	Stem	Large	Live	Quadratic	Coarse	Reference
		(m a.s.l.)	T (°C)	(mm)	type		Class		Area (m ² ha ⁻¹)	density (n ha-1)	living trees (n ha ⁻¹)	Aboveground Biomass (Mg ha ⁻¹)	Mean Diamater (cm)	Woody Debris (m ³ ha ⁻¹)	
		50.4	. -	1101	5.5	sp., Abies sp.		450 400	20 5			200	10.0		(2012)
	US - Maine/New Hampshire	534	6.5	1194	DB	Acer saccharum, Fagus grandifolia, Betula sp.pl.	OG	150-400	28.7	111		208	19.0		*Hoover et al. (2012)
	US - Maine/New Hampshire	534	6.1	1194	С	Tsuga canadensis, Picea sp., Abies sp.	OG	150-400	47.7	162		263	15.7		*Hoover et al. (2012)
	US - Ohio	1300	10.7	1012	DB	Quercus alba, Fagus grandifolia, Acer saccharum	OG	300-350	32.9	234			42.3		*McCarthy et al. (2001)
	US - Ohio	260	10.5	980	DB	Quercus alba, Acer saccharum, Quercus velutina Acer rubrum	OG	>150	30.8	365	33^{\dagger}		32.8	48.3	Goebel and Hix (1996)
	US - Ohio	240	10.5	980	DB	Quercus alba, Acer saccharum, Quercus veluting Acer rubrum	М	130-149	26.4	265	15^{\dagger}		35.6	5.3	Goebel and Hix (1996)
	US - Ohio	265	12.6	10019	DB	Quercus alba, Acer saccharum, Liriodendron tulipifera, O. prinus	OG		29.6	371.4			31.9		McCarthy et al. (1987)
	US - Pennsylvania	350	18.9	1000	C/DB	Tsuga canadensis	OG		39.1	218			47.8		Williams and Cook
	US -	350	18.9	1000	C/DB	Pinus strobus, Tsuga	М	70	27.1	266			36		Williams and Cook
	US -	530	6.0	1090	C/DB	Tsuga canadensis, Fagus	OG	~300	48.8	414	82^{\dagger}		38.7		*Ruffner and
	US -	470	8.0	1170	С	Pinus strobus, Tsuga	OG	250-300	78.6	347	179^{\dagger}		53.7		Abrams and Orwig
	US - Tennessee	930			C/DB	Tsuga canadensis, Fagus grandifolia, Acer saccharum	OG		47.6	802		398.0	26.9		(1996) *Busing (1998)
	US - Tennessee		16.2	1320	DB	Quercus sp.pl., Carya sp.pl., Acer saccharum, Fagus grandifolia	OG		39.2	363	26		37.1		Kupfer and Kirsch (1998)
	US - Tennessee	1049			C/DB	Tsuga canadensis, Halesia carolina, Fagus grandifolia, Acer saccharum	OG		52.4	877		440.0	27.6		Busing and White (1993)
	US - Virginia	195			DB	Liriodendron tulipifera, Ouercus prinus	OG		43,1	1720		379.1	17.8		*Druckenbrod et al. (2005)
	US - Virginia	1066.5	14.0	1070	DB	Quercus rubra, Betula lenta Prunus serotina	OG	300	43.9	433	83 [†]		35.9		Abrams et al.
	US - Virginia	884	14.0	1070	DB	Quercus prinus, Q. rubra, Acer rubrum	OG	300	35.8	288	73^{\dagger}		39.8		Abrams et al. (1997)
	US - West Virginia	994	8.8	1320	DB	Acer saccharum, Tilia americana, Quercus rubra	OG	~150	40.9	444	87^{\dagger}				Abrams et al. (1998)
	US - West Virginia	1155	8.1	1340	С	Picea rubens	OG	164-201	37.9	475	26.2		31.9		*Adams and Stephenson (1989)
	US - Washington				С	Abies procera	OG	400	147.0			1562 [‡]			Waring and Franklin (1979)
	US - Washington				С	Sequoia sempervirens	OG		247.0			3200.0			Waring and
	US - Washington	355			С	Tsuga heterophylla,	OG	450	94.0	520	129^{\dagger}		48.0		North et al. (2004)

S. Burrascano et al./Forest Ecology and Management 291 (2013) 458-479

464

Continent	Region	Altitude (m a.s.l.)	Mean T (°C)	Mean P (mm)	Forest type	Dominant tree taxa	Age Class	Age	Basal Area (m² ha ⁻¹)	Stem density (n ha-1)	Large living trees (n ha ⁻¹)	Live Aboveground Biomass (Mg ha ⁻¹)	Quadratic Mean Diamater (cm)	Coarse Woody Debris (m ³ ha ⁻¹)	Reference
	US - Washington-			1900	С	Pseudotsuga menziesii Pseudotsuga menziesii	OG	>195	69.0	448			44.3	425	Spies and Franklin (1991)
	US - Washington-			1900	С	Pseudotsuga menziesii	М	80-195	59.0	452			40.8	209	Spies and Franklin (1991)
	US - Oregon				С	Pseudotsuga menziesii, Tsuga heterophylla	OG	>400	127.0			1590 [‡]			Waring and Franklin (1979)
	US - Oregon				С	Sequoia sempervirens	OG	>1000	338.0			3461 [‡]			Waring and
	US - California	56		1792	С	Sequoia sempervirens	OG		379.0	164		4277.0	171.6		Sillett and Van Pelt (2007)
	US - California	75			С	Sequoia sempervirens	OG	>1000	330.0	380		4248.0		797	Busing and Fujimori (2005)
	US - California	2000		1250	С	Abies concolor, Pinus lambertiana, Calocedrus decurrens	OG		68.0	537	91 [†]		40.2		North et al. (2004)
	US - Colorado	192		1200	С	Picea engelmanni, Pinus contorta	OG	200-450	63.0	1140	111^{+}	253.0	26.5		Binkley et al. (2003)
South America	Chile - Andes	395		3100	EB	Aextoxicon punctatum, Laureliopsis philippiana	OG	380-645	242.8	1865			40.7		*Pollmann and Veblen (2004)
	Chile - Andes	890		3100	EB	Nothofagus dombeyi, Laurelionsis philippiana	OG	570-785	224.7	1321			46.5		*Pollmann and Veblen (2004)
	Chile - Andes	1110		6800	EB	Nothofagus dombeyi, Nothofagus pumilio	OG	430-590	126.1	1185			36.8		*Pollmann and Veblen (2004)
	Chile - Andes	1320		6800	DB	Nothofagus betuloides	OG	410-560	98.3	1893			25.7		*Pollmann and Voblen (2004)
	Chile - Andes	775			EB/C	Nothofagus dombeyi, Laureliopsis philippiana,	OG	>300	125.6	502		877.3	57.3	88.8	Schlegel and Donoso (2008)
	Chile - Andes	775			EB/C	Saxegothaea conspicua Laureliopsis philippiana, Saxegothaea conspicua, Dasyphyllum diagonthoidog	OG	>300	109.4	652		664.5	48.2	59.6	Schlegel and Donoso (2008)
	Chile (S-Central)	800		5000	EB/C	Nothofagus dombeyi, Laureliopsis philippiana,	OG		126.0	501	127^{\dagger}		56.6		Donoso and Lusk (2007)
	Chile (S-Central)	480	11.5		EB	Eucryphia cordifolia, Weinmannia trichosperma	OG		81.0	719			37.9		*Donoso and Nyland (2005)
	Chile (S-Central) Chile (S-Central)	376 400			EB DB	Aextoxicon punctatum Nothofagus obliqua, N. alnina, N. dombevi	OG M	260-460 86	95.1 55.0	661 1006	206^{\dagger}		42.8 26.4		Salas et al. (2006) Salas and Garcia (2006)
	Chile - Chiloe island	170	9.1	2157.8	EB/C	Podocarpus nubigena, Saxegothaea conspicua, Nothofagus nitida	М	112-134	97.9						*Gutierrez et al. (2009)
	Chile - Chiloe island	310	9.1	2157.8	EB/C	Podocarpus nubigena, Saxegothaea conspicua, Nothofagus nitida	OG	173-309	104.0	1513	177^{\dagger}		29.6		Gutierrez et al. (2009)
	Chile - Chiloe	350	11.0	2516	EB	Amomyrtus luma,	OG		78.4	1326			27.4		Armesto and
	Chile - Chiloe	50	11.0	2516	EB	Aextoxicon punctatum,	OG		32.2	1436			1639		Armesto and

(continued on next page) 65^{40}_{55}

Continent	Region	Altitude (m a.s.l.)	Mean T (°C)	Mean P (mm)	Forest type	Dominant tree taxa	Age Class	Age	Basal Area (m² ha ⁻¹)	Stem density (n ha-1)	Large living trees (n ha ⁻¹)	Live Aboveground Biomass (Mg ha ⁻¹)	Quadratic Mean Diamater (cm)	Coarse Woody Debris (m ³ ha ⁻¹)	Reference
	island Chile - Chiloe island	150	11.0	2516	EB	Luma apiculata Drimys winteri	OG		43.6	750			27.2		Figueroa (1987) Armesto and Figueroa (1987)
	Chile - Chiloe island	30	11.0	2516	EB	Drimys winteri, Amomyrtus luma, Nothofagus dombeyi	OG		61.2	949			28.7		Armesto and Figueroa, 1987
	Chile - Chiloe island	680	6.0	4500	EB/C	Fitzroya cupressoides, Pilgerodendron uviferum, Nothofagus nitida	OG	400-425	138.2	9680	36 [†]	576.0	13.5		Battles et al. (2002)
	Chile - Chiloe island	75			EB/C	Nothofagus nitida, Drimys winteri, Podocarpus nubigena	М	97-112	47.4	3600			13.0	215	*Carmona et al. (2002)
	Chile - Chiloe island	75			EB/C	Nothofagus nitida, Drimys winteri, Podocarpus nubigena	OG	133	75.7	2400			20	219	*Carmona et al. (2002)
	Chile - Chiloe island	550	9.5		EB/C	Nothofagus nitida, Drimys winteri, Podocarpus nubigena, Laureliansis nhilinniana	OG	191	89.2	3114			19.1	782	*Carmona et al. (2002)
	Chile (Southern)	30			EB/C	Nothofagus nitida, Podocarpus nubigena	М		94.1	688			41.8		Innes (1992)
	Chile (Southern)	220	4.3	690	DB	Nothofagus pumilio	Μ	180	80.0	360		492.5	41.7		Barrera et al. (2000)
	Chile (Southern)	440	3.1	954	DB	Nothofagus pumilio	М	158	67.0	780		349.8	31.7		Barrera et al. (2000)
	Chile (Southern)	1200		2750	DB	Nothofagus alpina, Nothofagus dombeyi	OG	137	89.5	451	150 [†]		50.3		Pollmann (2003)
	Chile (Southern)	1500		1800	DB	Nothofagus pumilio	OG		68.0	499			52.2		Fajardo and de Graaf (2004)
	Chile (Southern)	900		1350	DB	Nothofagus pumilio	OG		61.9	782			33.8		Fajardo and de Graaf (2004)
	Argentina				DB	Nothofagus pumilio	OG	125-250	69.2	584			38.9		*Martínez Pastur et al. (2000)
	Argentina	150		850	EB	Nothofagus betuloides, Drimys winteri	OG	150-161	104.0	1906	25^{\dagger}		26.4		*Rebertus and Veblen (1993)
	Argentina	350			DB	Nothofagus pumilio	OG	100	95.0	717	32 [†]		41.1		Rebertus and Veblen (1993)

late-successional ecosystem services (Franklin et al., 2002; Bauhus et al., 2009). However, the direct contributions from the study of old-growth forests to sustainable silvicultural systems has been partially hampered by context-dependent results from studies focused on individual forest types and by the scarcity of late-successional forest remnants for many forest types (Oliver and Larson, 1990; Brang, 2005).

Even if the interest in old-growth forests arose long ago (Jones, 1945) and these systems have been studied in many temperate regions of the world, only recently have researchers begun to work on assessments of old-growth ecosystem structure and functions at the global scale (Luyssaert et al., 2008; Keith et al., 2009). Yet there is still much to learn from old-growth forest remnants. In particular, identifying the degree to which structural characteristics co-vary among temperate old-growth forests would provide an indication of how a range of ecosystem services could respond to choices societies make about how to manage or conserve forests in the temperate zone. With old-growth forests rapidly disappearing and under pressure in many regions (e.g. Gutierrez et al., 2009; Knorn et al., in press), there is a compelling need to aggregate available information on old-growth structure. In this study we review the existing literature reporting data on structural attributes of moist temperate old-growth forests with a special focus on their distinctive features when compared to mature forests. Our research objectives are to (i) identify shared attributes among oldgrowth stands across different continents and forest types, (ii) investigate variability among these, and (iii) explore gaps in the data globally in order to inform future research.

2. Methods

2.1. Literature search

We assembled a dataset from a literature search performed in December 2011, in which we looked for studies reporting the structural attributes of old-growth temperate forests. Studies comparing data between old-growth and mature forest sites were particularly valued. Relevant publications were identified by searching the online database of Web of Science using the following search words: 'old-growth and structur* AND (volume or biomass)'. An additional search for data on mature sites was performed for those continents (Asia, Australia and South America) where age comparison studies were scarce. For these we used the additional search words: '(mature OR managed) AND forest* AND structur*'.

The initial search was then further refined within the ISI website. We retained a paper in the dataset when it was associated with one or more categories of interest. These included subject area (forestry, ecology, environmental sciences, plant sciences, biodiversity conservation), and document type (article, proceeding paper, review). We considered only data published in English language and peer reviewed journals. We included data from any sites defined in their respective papers as 'temperate'. Where no direct reference to biome occurred we followed the geographic boundaries reported for the temperate forest biomes (Olson et al., 2001). We focused only on moist temperate forests, i.e. those systems where climate seasonal variation is determined more by temperature than by precipitation. The search did not include forests prone to high to moderate frequency ground-fire in regions with low precipitation during the growing season due to fundamentally different structural development processes in these systems. The Web of Science search missed several papers reporting data on old-growth forests because terms other than 'old-growth' were used in the title and abstract. Terms like 'virgin', 'ancient', 'primeval', 'near-natural', and 'untouched' were often used interchangeably with the concept of 'old-growth', sometimes without clear distinction of whether the reference was to an old-growth condition or a primary (never cleared) forest. Therefore to achieve a comprehensive dataset, we added studies cited in other papers included in our dataset, or based on the authors' knowledge of the field, or suggested by the reviewers. Each paper was checked to confirm that data were reported for late-successional/old-growth stands, rather than early to mid-successional (natural disturbance originated) stands existing within a primary forest matrix.

This process produced a robust dataset that, nevertheless, is not meant to be a full representation of all papers ever published on this topic. Indeed, during the selection process we evaluated the return in terms of information provided by each additional paper, taking into account regional gaps, and discontinued the research when there was sufficient representation of major forest types for each region.

2.2. Data extraction

The full set of papers identified by our search was filtered to remove those not reporting values for biometrics of direct relevance to our analysis. We excluded those papers not reporting at least three of the stand structural or compositional variables included in a preliminary list comprising: basal area, stem density, large living tree density, live aboveground biomass, mean stand and quadratic mean diameter, volumes of different components of coarse woody debris, canopy height and closure, gap metrics, vertical heterogeneity indices, and tree species richness. Furthermore, in cases where more than one paper reported data for the same site(s), we selected the study reporting the largest number of variables. Since several papers reported data from numerous different stands, these structural data were averaged only if those stands were similar in terms of geographic location, canopy composition and age classes.

The final dataset consisted of 147 records derived from 93 papers (Table 1) describing late-successional/old-growth temperate forests from many different regions and forest types around the world, all continents having temperate zones were represented (Fig. 1). Each record was classified in terms of age class and broad forest type.

Age classes (e.g. mature vs. old-growth) were assigned as reported in the original paper, and not from the analysis of age or structural attributes. The criteria employed in the original papers resulted in wide age ranges, both for mature and old-growth age classes, accounting for intrinsic variability across regions and forest types. In two cases (Wijdeven, 2004; Merino et al., 2007) stands described as near-natural were included in the old-growth group. A stand attributed to the 'degradation phase' was included in the old-growth group (Marage and Lemperiere, 2005) while stands attributed to the 'optimal' and 'decay' stages were included respectively in the mature and old-growth group (Akhavan et al., 2012; Heiri et al., 2009). In few comparative studies, stands defined by authors as 'secondary' or 'second growth' (Aiba et al., 2001; Fisk et al., 2002) or in 'biostatic phase' (Marage and Lemperiere, 2005), were classified in our analysis as 'mature'.

On the basis of the main traits of dominant species we defined five broad forest types: C – conifers; C/DB – mixed conifer/deciduous broadleaf; DB – deciduous broadleaf; EB – evergreen broadleaf; and EB/C mixed evergreen broadleaf/conifer. Since very few examples of mixed evergreen/deciduous broadleaf forests occurred in our dataset (see Pollmann and Veblen, 2004; Salas and Garcia, 2006), these stands were included either in the EB or DB type according to the dominant species. In the final table, we maintained only those structural variables reported for at least 25% of the records. These were: basal area, stem density, large living tree density (LLT – i.e. trees with DBH > 50 cm per hectare), live aboveground biomass (LAB), quadratic mean diameter (QMD) and coarse woody debris volume (CWD).



Fig. 1. World distribution of temperate forests (dark grey) within the temperate latitudinal zone according to Olson et al. (2001), and location of stands (triangles) included in the review for which detailed geographic information was available.

Inevitably structural attributes were derived from field measurements that followed different sampling designs. Only in a minority of papers, plots were randomly or systematically located. Most of the studies, instead, located plots on the basis of either subjective or qualitative assessment (closeness to the core area, no sign of alteration, etc.), also in relation to the small extent of old-growth remnants. Plot sizes and layouts varied greatly; indeed two different approaches were generally used: (i) one or few big plots (\sim 1 ha), or (ii) several relatively small plots (100–400 m²). However the sampling of old-growth spatial heterogeneity was generally ensured by a trade-off between plot size and number.

We also found some discrepancies among structural data collection and reporting protocols in different studies, such as minimum live tree and CWD size thresholds, and LAB estimation methods. Before the analysis, we tested for significance the difference in live tree DBH thresholds between age classes (Kruskall–Wallis: H = 1.655, p = 0.198), since a systematic difference could have biased our results. Live trees and CWD thresholds were reported in Appendix B, Table B1. The lack of LAB data reported for Europe is noteworthy and is linked to the tradition of focusing on wood volumes more than aboveground biomass. Volume data was available only for 27 European records (Appendix A, Table A1, Fig. A1). For a thorough description of the variables considered in this paper and on the way the other discrepancies were taken into account see Appendix B.

Beside these structural indicators, we recorded also data to be used as site descriptors, such as stand age and environmental variables, namely geographic location (latitude and longitude), altitude, and climate (mean annual temperature and annual precipitation). In particular, stand age in multi-aged forests was assessed in several different ways in the papers included in our review. We reported as age the value given in the original papers. Most authors used the mean age of the overstory trees. In some studies these estimates were derived from partial observations, such as the mean age of the dominant species (Sano, 1997), or the age of the illegally cut stumps at the periphery of the area (Keeton et al., 2010) or of fallen individuals (Parker et al., 1985). In other cases (Battles et al., 2002; Pollmann, 2003) we derived age from the most common class of the age histogram. Finally, in 10 stands (Lusk and Smith, 1998; Aiba et al., 2001; Beaudet et al., 2007; Heiri et al., 2009; Král et al., 2010; Sato, 2010) we reported the time since last disturbance given by the authors; however we did not use these data as age in the analysis because they were reported only in a small percentage of papers and had a range substantially different from the other age values.

2.3. Statistical analysis

Age and structural variables were compared between mature and old-growth stands, globally and across different continents and broad forest types. For each contrast Monte-Carlo simulations were used to test whether the median of each variable was significantly different between age classes: we randomly reassigned each record 4999 times to an age class (mature or old-growth), while maintaining the sample size of each group. The actual difference between medians was considered significant when it was not within the range encompassing the 2.5–97.5 percentiles of the distribution of the permuted differences. When the sample size was not large enough to allow at least 4999 different combinations, we compared the actual difference to the complete distribution of the permuted differences, based on each possible combination of reassignments.

To explore sources of variability within the dataset, nonparametric correlation coefficients (Spearman's ρ) were calculated for structural variables in relation to site descriptors (latitude, altitude, mean annual temperature and precipitation, stand age), the significance level was set at $\alpha = 0.05$. All the statistical calculations were performed using R 2.12.0 (R Development Core Team, 2010).

3. Results

3.1. Data distribution across continents, forest types and age classes

North America had the largest number of records (37%), reporting data predominantly from the Midwest (13%), and U.S. Northeast (12%), although a great body of research on late-successional forest dynamics (e.g. disturbance regimes, development processes, etc.) came from the Pacific Northwest region. Europe and South America were well represented (26% and 20% of records respectively). Asia and Australia accounted only for 14% and 3% of the records respectively (Fig. 2), also in relation to the lower representation of temperate biome in these continents. The literature spanned a wealth of forest types. The majority of the papers investigated deciduous broadleaf (33%), mixed conifer/deciduous broadleaf



Fig. 2. Distribution of forest types across continents. C – conifer; DB – deciduous broadleaf; EB – evergreen broadleaf; C/DB – mixed conifer/deciduous broadleaf; EB/C – mixed evergreen broadleaf/conifer.

(26%), and conifer forests (20%). Less numerous were papers describing evergreen broadleaf forests (12%) and mixed evergreen broadleaf/conifer forests (9%).

Papers on late-successional stands by forest type were unevenly distributed among continents. In Europe and North America three forest types were represented (C, C/DB, DB) with very similar proportion of mixed conifer/deciduous broadleaf and deciduous broadleaf forests, which accounted respectively for 36% and 34% of the records in North America, and 37% and 39% in Europe (Fig. 2). The most common systems in our dataset were northern hardwood (*Acer saccharum, Fagus grandifolia, Tsuga canadensis, Betula* sp.) in North-America, and European beech-Norway spruce-silver fir (*Fagus sylvatica, Picea abies* and *Abies alba*) in Europe (Table 1).

At the global level median age values differed significantly between age classes (250 years for old-growth and 125 for mature stands, p < 0.001). When single contrasts were considered all continents displayed significant differences but Australia, for which no sufficient data were available; all forest types but deciduous broadleaf and mixed evergreen broadleaf/conifer forests significantly differed in age values (Fig. 3). The analysis of age ranges (51– 252 years for mature stands and 100–1000 for old-growth) further confirms the great variability in stand development rates across forest types and geographic regions.

3.2. Comparison between mature and old-growth forests

The comparison between mature and old-growth stands identified common patterns for most of the structural attributes, but also a certain degree of variability related to management history, dominant species traits, and site characteristics. In the global dataset, old-growth forests showed higher basal area values than mature forests although the difference was not significant neither at the global level (p = 0.207), nor for single forest types. The only significant differences we found were for North and South America (p = 0.028 and 0.007 respectively). An opposite non-significant result was observed in Europe and Australia (Fig. 4; Table 2) and in deciduous broadleaf forests globally (Fig. 5; Table 3).

Stem density values recorded globally were slightly lower for old-growth forests when compared to mature stands (Table 2). This result was consistent across all continents and forest types. The data for LLT showed distinct differences between age classes, with significantly higher densities reported for old-growth (median 72.6 trees ha⁻¹) compared to mature stands (16.3 trees ha⁻¹; p < 0.001). However, we only had data to contrast LLT for Europe and North America, and for the forest types surveyed in these continents. For North America the difference between the two age classes was significant (p = 0.031), with mature forests having far lower LLT densities (10–20 trees ha⁻¹) than old-growth (2.5– 199 trees ha⁻¹, Table 2). For Europe, the density of LLT ranged from 36.5 to 122 trees ha⁻¹ in the old-growth stands and from 4.2 to 83 trees ha⁻¹ in mature stands (p = 0.031).

LAB data were available only for few studies (n = 33) and this allowed us to contrast age classes only for a subset of regions, limiting our ability to discern age related trends. At the global level we observed significantly higher LAB in old-growth than in mature stands (medians equal 379.1 and 235.2 Mg ha⁻¹ respectively, p = 0.011, Table 2). The only individual contrast giving a significant result was North America, (p = 0.008). This result may have been influenced by the lack of data for mature stands dominated by *Sequoia sempervirens* and *Pseudotsuga menziesii*, that are species characterized by very high LAB values, although also excluding these stands the result did not change (not shown).

Our data demonstrated slightly higher QMD values in oldgrowth forests when compared to mature stands in all continents and forest types but this difference was significant only at the global level (p = 0.026).

CWD volumes exhibited consistent age class contrasts for all the continents and forest types. The median value calculated on the global dataset was significantly different, with 67.8 m³ ha⁻¹ for mature and 151.4 m³ ha⁻¹ for old-growth stands (p = 0.002). While the difference between median values was high, CWD was highly variable and the range overlapped considerably among age classes (Table 2). The only contrasts giving significant results were Europe, (p = 0.002), and conifer forests (p = 0.036). No contrasts were possible for Asia and Australia, and for evergreen broadleaf stands due to scant availability of data on CWD.

3.3. Correlation between structural attributes and site characteristics

Many site descriptors showed significant correlations with structural variables at global scale (Table 4). With regards to



Fig. 3. Boxplots of the age data distribution for the global dataset and for each continent and forest type. The median, first and third quartiles, and whiskers (corresponding to 1.5 times the inter-quartile distance) are reported. Empty circles represent outliers. Sample size is reported at the bottom of each box. Monte-Carlo two-sided significance levels are at the top (n/a: no test was possible; ns: not significant; *0.05 ; <math>**p < 0.01).

climatic and geographic variables, basal area and stem density were positively correlated with mean annual precipitation and negatively correlated with latitude. CWD displayed a negative association with mean annual temperature, and LLT was positively correlated with precipitation. Among the site descriptors, forest age was positively and significantly correlated with the highest number of structural indicators, including basal area, LLT, LAB and QMD.

4. Discussion

With some exceptions, old-growth forests show many structural similarities at the global scale.

Contrasts against mature stands hold true across different continents and forest types despite the variability in age ranges used to differentiate between mature and old-growth stands in different regions. Indeed old-growth are characterized globally by significantly higher LLT densities, higher QMD, and greater LAB and CWD volume. We also observed somewhat higher basal areas and lower stem densities in comparison to mature stands, although these differences were not statistically significant due to substantial variability among forest types.

4.1. Global commonalities

Our review showed old-growth forests to host significantly higher amounts of biomass than mature forests, stored both in LAB and in CWD pools. Furthermore, the global data showed a significant positive correlation between LAB and stand age. This is consistent with recent models describing carbon dynamics in boreal and temperate forests, supporting the hypothesis of a potential for continued carbon accumulation late into stand development (Keeton et al., 2011), both in broadleaf and conifer stands (Luyssaert et al., 2008), often mediated by variation in climate, ecoregion and forest type (Keith et al., 2009; Hudiburg et al., 2009; Stegen et al., 2011). The analysis of trends in aboveground biomass in late-successional systems is of primary importance because it correlates also with carbon allocation to belowground biomass (Cheng et al., 2007) and to total ecosystem carbon (Houghton, 2005; Hoover et al., 2012) that is often difficult to quantify. LAB is the quickest pool to recover after disturbance, and likely a full accounting of all pools, especially C stored in soil, would reveal stronger

differences between old-growth and mature forests than comparing LAB alone (Zhou et al., 2006; Rhemtulla et al., 2009).

Higher LAB and associated carbon storage values in old-growth forests relate to structural attributes typical of this developmental stage. Living biomass reaches very high levels because of: (i) the vertically continuous and multispecies canopy typically associated with old-growth forests (Keith et al., 2009; Wirth et al., 2009), allowing for efficient allocation of growing space within threedimensions and across trees of different age, size and shade tolerances; and (ii) the possibility of LLT to maintain net positive growth and accumulate biomass for centuries (Xu et al., 2012). Indeed the literature reports significantly a general trend of higher LLT densities in old-growth when compared to mature forests, and despite variability attributed to species composition, region, disturbance history and site productivity, our results suggest a significant correlation between stand age and LLT at global scale.

Our results at the global scale were consistent with previous studies showing LLT to be a strong predictor of late-successional structure at regional scales (e.g. Whitman and Hagan, 2007). A minimum LLT density of 30 trees with DBH > 70 cm per hectare was reported for European old-growth forests (Nilsson et al., 2002). Such a threshold seems too high and not characteristic of old-growth systems at global level; it would exclude several oldgrowth stands in our dataset. Indeed about 90% of the old-growth stands in our dataset contained more than 30 LLT per ha, therefore, we believe a more inclusive minimum density of 30 trees with DBH > 50 cm per hectare may be useful both at global and at European level. LLT density is directly related to CWD recruitment, particularly for the largest size classes, and contributes to the pool of long-lasting deadwood, as deadwood residence time is strongly influenced, together with wood density and decay resistance, by tree size at the time of death (Mackensen et al., 2003; Cornwell et al., 2009).

Our review showed significantly higher CWD quantities in oldgrowth than in mature forests and confirmed the importance of CWD as an indicator of old-growth condition (Peterken, 1996; Bobiec, 2002), at least for those continents for which we had sufficient data. Despite the significant differences between medians at the global level, ranges in CWD widely overlapped across most continents and forest types, probably due to a strong variability among and within systems that may be related to two main factors. First, our results clearly showed a climatic influence, with the warmest regions characterized by the lowest volumes of deadwood. Accordingly, the lowest values for the old-growth stands were recorded in



Fig. 4. Boxplots of the data distribution for the global dataset and for each continent. The median, first and third quartiles, and whiskers (corresponding to 1.5 times the interquartile distance) are reported. Empty circles represent outliers. Sample size is reported at the bottom of each box. Monte-Carlo two-sided significance levels are at the top (n/ a: no test was possible; ns: not significant; *0.05 < p < 0.01; **p < 0.001, **p < 0.001).

a mixed oak forest in southern Ohio (48.3 m³ ha⁻¹, Goebel and Hix, 1996); in Italy (45 m³ ha⁻¹, Burrascano et al., 2008; and 65.2 m³ ha⁻¹, Travaglini et al., 2012); and in the northern-most Chilean old-growth stands (59.6 m³ ha⁻¹, Schlegel and Donoso, 2008). The correlation we observed between CWD and temperature gradient at the global level are consistent with the results of a global analysis that highlighted mean annual temperature as a main driver of CWD decay rates (Mackensen et al., 2003). Second, the high degree of variability in CWD is also related to differences in plant traits. Especially wood decomposition rates are species-specific (Zell et al., 2009), and are far slower in gymnosperms than angio-sperms due to a higher C:N ratio, higher lignin and resin content,

and fine-scale anatomical traits that slow the spread of fungi (Cornwell et al., 2009).

Understanding the sources of variability in CWD volume has important consequences for quantifying the carbon stored in deadwood, given that CWD is estimated to contain 3.3 billion tons of carbon in temperate forests globally corresponding to about 7% of total living carbon (Pan et al., 2011). Focusing on CWD/live wood ratios allows us to eliminate productivity related differences between forest types (Wirth et al., 2009). CWD/live wood volume ratios in the literature ranges from 13% to 37% in European beech forest reserves (Christensen et al., 2005), and between 6% and 23% in old-growth broadleaved forests of North America (Peterken,

Table 2

Median and range of age and structural variables, calculated for the whole dataset and for each continent. When only one value was available the median alone was reported. Bold values represent significantly different contrasts (*p* < 0.05).

	Global		Asia		Australia		Europe		North An	ierica	South An	nerica
	М	OG	М	OG	М	OG	М	OG	М	OG	М	OG
Age (yrs) Min. Median Max.	51 125 252	100 250 1000	51 100 105	110 200 300	150 197 252		109 137.5 220	140 217.5 300	70 118.5 140	150 300 1000	105 123 180	100 300 678
BA (m ² ha ⁻¹) Min. Median Max.) 15.6 41.7 144.5	24.4 47.3 379	15.6 42.2 54.7	34.3 50.2 69.4	61 68.2 144.5	55.9 62.6 69.2	30.7 42.2 69.2	24.4 38.5 57.4	24 29.1 59	28.7 44.4 379	47.4 67 97.9	32.2 94.6 242.8
Stem density Min. Median Max.	(n ha ⁻¹) 217.5 536 6015	96.7 501 9680	336 632.5 2805	281.3 630 4570	1216 1660 6015	- 444.7 -	259 444 2182	124 341 1835	217.5 488.5 1000	96.7 392 1720	360 893 3600	451 865.5 9680
LLT (n ha ⁻¹) Min. Median Max.	4.2 16.3 83	2.5 72.6 206	- 70 -	70 66 83	- - -	- 55 -	4.2 22 83	36.5 75.5 122	10 13.5 20	2.5 62.5 199	- -	25 127 206
LAB (n ha ⁻¹) Min. Median Max.	144 235.2 492.5	208 379.1 4277	- 258 -	254.3 279 315.7	- 240.4 -	- - -	- 181.2 -	255.5 303 510	144 148 230	208 388.6 4277	349.8 421.2 492.5	576 664.5 877.3
QMD (cm) Min. Median Max.	13 27 53.2	13.4 34.6 171.6	15.8 23.7 41.1	13.4 28.4 51.4	17.5 21.6 26.7	- 40 -	15.3 36.1 46.4	14.6 39.3 64.3	18.4 27 47.8	15.7 32.8 171.6	13 29.8 53.2	13.5 37.4 57.3
CWD (m ³ ha' Min. Median Max.	⁻¹) 5.3 67.8 214.5	45 151.4 797	- - -	- -	- -	- -	8 28 91.4	45 157.3 469	5.3 67.8 209	48.3 151.3 797	- 214.5 -	59.6 153.6 781.5

1996). The ratios based on our data on CWD and live wood volume in Europe ranged from 6% to 89% in old-growth and from 2% to 9% in mature forests (see Table Al). Indeed, Europe was the only contrast returning significant differences between CWD volumes in mature and old-growth stands. Very low CWD values in European mature forests may be due to the long history of forest exploitation that probably limited the ability of European studies to find or assess unmanaged mature reference stands (Commarmot et al., 2005; Merino et al., 2007; Burrascano et al., 2008; Motta et al., 2010). It may relate also to higher amounts of deadwood collected for fuel by local populations in mature forests, which are usually more accessible than old-growth remnants located in reserves or in remote areas and high altitudes.

In conclusion, these results showed ratios for late successional forests to be much higher than the global average of temperate forests, suggesting that the CWD carbon pool is currently understocked in secondary and managed temperate forests, especially in Europe. The CWD pool, though transient, can represent an important net C sink as long as input rates (mortality) equal or exceed outputs (decomposition), and thus should be carefully accounted for in terrestrial carbon models (Stephenson et al., 2011). Management practices aimed at maintaining, or recreating, old-growth forest attributes (Keeton, 2006; Bauhus et al., 2009; Rhemtulla et al., 2009) may increase the amount of carbon stocked in temperate forests, both in live biomass and CWD (Keeton, 2006; Bauhus et al., 2009; Rhemtulla et al., 2009). This is especially relevant given the growing interest in extending mechanisms like REDD (Reduced Emissions from Deforestation and Degradation) to temperate regions, to maintain terrestrial carbon sinks.

Furthermore high CWD volumes and LLT densities in oldgrowth forests, which were evident in our data, also influence the provision of habitat for late-successional biodiversity. CWD is a key ecosystem feature supporting high levels of biodiversity, for instance by providing diverse microhabitats for numerous specialized and saproxylic organisms (Grove, 2002; Odor et al., 2006; Mckenny et al., 2006). The long-term availability of deadwood may be related to the persistence of many saproxylic organisms (Lassauce et al., 2011) with low dispersal capabilities that may be driven to local or regional extinction in the absence of a temporal continuity in the deadwood substrate, due to intensive management (Siitonen, 2001). For this reason, besides being a robust indicator of old-growth condition (Whitman and Hagan, 2007), CWD has become one of the primary indicators of sustainable forest management in Europe (MCPFE, 2003; Lassauce et al., 2011) and a structural indicator in the context of Biodiversity Evaluation Tools (Larsson, 2001) and international certification systems (Foster et al., 2008).

Also LLT, besides storing a large amount of carbon, provide habitat functions for a number of threatened or ecologically important temperate forest species, ranging from fungi to lichens, saproxylic beetles, birds and bats (Mikusinski and Angelstam, 1998; Ruffell et al., 2007; Persiani et al., 2010; Hatanaka et al., 2011). These functions relate to the great variety of microhabitats large trees offer, including rough bark, trunk hollows, exposed deadwood, sapflows, dead branches, and dead tops (Nilsson et al., 2002; Winter and Moller, 2008).

4.2. Sources of variability

Although our review found distinct commonality in old-growth structure, several attributes showed a high degree of variation precluding overly simplistic generalizations at the global scale. Indeed basal area and stem density did not show statistically significant differences between age classes in our dataset, but instead were more strongly related to geographic location, species composition, and climatic influences. QMD on the other hand, showed a signifi-



Fig. 5. Boxplots of data distribution for forest types (C – conifer, DB – deciduous broadleaf, EB – evergreen broadleaf, C/DB – mixed conifer/deciduous broadleaf, EB/C – mixed evergreen broadleaf/conifer). The median, first and third quartiles, and whiskers (corresponding to 1.5 times the inter-quartile distance) are reported. Empty circles represent outliers. Sample size is reported at the bottom of each box. Monte-Carlo two-sided significance levels are at the top (n/a: no test was possible; ns: not significant; *0.05 < p < 0.01; **p < 0.001, ***p < 0.001).

cant difference between age classes only at the global level with slight, non-significant differences in the other contrasts.

Variation like this raises interesting questions regarding differences among forest systems, for instance related to environmental controls, disturbance regimes, and management history. Previous researchers have found old-growth to generally have higher basal areas compared to mature forests (Keddy and Drummond, 1996; Gutierrez et al., 2009). Nevertheless, despite the significant positive correlation between basal area and age, our contrasts highlighted a large amount of variation between forest types and continents: old-growth showed significantly higher values in North and South America and non-significantly lower basal area in Australia and Europe when compared to mature stands. For Australia, very few basal area data were available and the high value reported for mature stands may derive from a single mature site with a relatively high age (252 years) (Smale et al., 1997). In Europe old-growth showed lower basal area values than mature stands, either depending on the confusion in terminology for late-successional stands, especially pervasive in the European literature, or because the old-growth remnants spared from exploitation are mostly located in remote, unproductive areas at relatively high altitude, where a lower basal area is naturally expected to occur (Nagel et al., 2006; Burrascano et al., 2008; Svoboda and Pouska, 2008; Holeksa et al., 2009). In some cases, anthropogenic influence may explain the high basal area values observed in European mature stands. Indeed timber production

Table 3

Median and range of age and structural variables calculated for each forest type. When only one value was available the median alone was reported. Bold values represent significantly different contrasts (p < 0.05).

	Conifer		Mixed conit deciduous b	fer/ proadleaf	Deciduou: broadleaf	5	Evergreen broadleaf		Mixed conit evergreen b	fer/ proadleaf
	М	OG	М	OG	М	OG	М	OG	М	OG
Age (yrs) Min. Median Max.	125 168.5 220	190 287.5 1000	70 118 135	140 250 350	70 115 180	100 209.5 350	51 100.5 150	155.5 520.5 678	105 160 252	133 270.5 413
BA (m ² ha ⁻¹) Min. Median Max.	45.3 61.2 69.2	32 63 379	24 31 49.4	26.5 44.8 69.4	15.6 39.9 80	24.4 37.3 95	54.7 61.5 68.2	32.2 79.7 242.8	47.4 79.45 144.5	55.9 104 138.2
Stem density (Min. Median Max.	(n ha ⁻¹) 452 489.5 1541	103 404 1618	217.5 748 1000	124 477.5 1130	259 446 2182	96.7 438.2 1835	1216 2010 2805	564 1321 4570	1660 3600 6015	444.7 687.5 9680
LLT (n ha ⁻¹) Min. Median Max.	- 4.2 -	26.2 91 179	10 11.5 20	2.5 64 122	15 22 83	16 71.6 199	- -	25 115.5 206	- - -	36 91 177
LAB (n ha ⁻¹) Min. Median Max.	- 181.2 -	222 1562 4277	144 148 230	221 330 440	349.8 421.2 492.5	208 328.6 510	240.4 249.2 258	254.3 279 315.7	- - -	576 664.5 877.3
QMD (cm) Min. Median Max.	19.2 41.2 41.6	15.7 38.3 171.6	18.4 26.8 47.8	19.3 35.1 64.3	15.3 33.1 53.2	14.6 33.1 51.4	15.8 21.3 26.7	13.4 27.4 46.5	13 17.5 21.6	13.5 40 57.3
CWD (m ³ ha ⁻ Min. Median Max.	¹) 28 78.4 209	144 191.5 797	10 61.1 177.8	65.2 151.4 383	5.3 25.6 91.4	45 119 469	- - -	- - -	- 214.5 -	59.6 153.6 781.5

Table 4

Spearman correlation coefficients (*ρ*), significance and sample size (*n*) for each pair of structural/explanatory variables. Bold values represent significant correlations (*p* < 0.05).

	Latitude	_	Altitude		Temperatur	e	Precipitatio	n	Age	
	ρ	n	ρ	n	ρ	n	$\overline{ ho}$	n	ρ	n
Basal area	- 0.25 **	120	-0.04	121	0.08	81	0.52***	105	0.39***	83
Stem density	- 0.25 **	120	-0.08	120	0.19	82	0.33***	105	-0.07	75
Large trees	-0.26	52	0.14	53	-0.17	38	0.29*	46	0.60***	40
LAB	-0.27	23	-0.20	22	-0.07	11	0.43	15	0.74***	18
QMD	0.10	117	0.12	118	-0.18	79	0.04	102	0.33**	75
CWD	0.25	38	-0.05	41	- 0.39 *	30	0.18	38	0.25	31

* 0.01 < *p* < 0.05.

** 0.001 < *p* < 0.01.

^{****} *p* < 0.001.

management both controls species composition towards more productive species and efficiently allocates growing space, and results in optimized tree production and high stand basal area.

The global variability among continents and forest types in the relationship between basal area and age class, may also relate to differences in the density of shade-tolerant species and productivity-related climatic controls. Indeed in a comprehensive study of old-growth forests on Chiloe island (Chile) the density of shade-tolerant stems was positively correlated with basal area, vertical heterogeneity and age, with high variability in relation to changes in temperature and in distance to coastline (Gutierrez et al., 2009). At the global scale our results showed precipitation to have a significant positive relation with basal area. We recommend further research on the relationships among these variables to assess the consistency of high basal area as a characteristic of old-growth conditions.

The relative abundance of shade-tolerant and intolerant species, together with differences in gap dynamics and pattern, and the rapidity of the self-thinning process, may also influence variation in stem densities observed among late-successional systems globally. Although stem density was generally lower in old-growth than in mature forests across most forest types, this difference was not statistically significant, nor did we find any significant correlation between stem density and age. This is consistent with the forest development model proposed by Franklin et al. (2002), wherein tree mortality in late-successional forests becomes increasingly influenced by density-independent factors (Das et al., 2011), such as natural disturbances, which can be highly variable across regions and forest types (Franklin et al., 1987; Allen et al., 2010). The death of overstory dominants allows more light to reach the forest floor, encouraging re-establishment of the understory community, depending on previous stand density dynamics. Other causes of variability in stem density from the mature to the old-growth condition may be environmental conditions (e.g. precipitation) together with factors affecting seed germination and establishment, such as availability of seed

sources, presence of suitable seed beds, and competition with herbaceous shrubs (Franklin et al., 1987; Donato et al., 2012).

In the comparative studies included in our review, stem density was generally lower in old-growth stands. Nevertheless, an increase in stem density with age was found in two comparative studies performed in U.S. Northeast forests dominated by *T. canadensis* and *Betula alleghaniensis* (Fisk et al., 2002; Keeton et al., 2007. These exceptions may be attributed to dense regeneration of intermediate to shade tolerant species within canopy gaps, at gap edges, or in the suppressed layers.

Variability in stem density was also related to latitudinal and precipitation gradients. Indeed, the stands with the highest stem density, with values up to four times greater than the other forest types, were evergreen broadleaf and mixed conifer/evergreen broadleaf stands that were located at lowest latitudes and displayed the highest precipitation values. Accordingly, at lower latitudes the more vertical sun angles may allow the evolution of vertically complex canopies with emerging large trees, and shade-tolerant trees in both the upper and lower canopy layers and thus very high stem densities (Armesto and Figueroa, 1987; Stephenson et al., 2011).

The slightly higher QMD values reported for old-growth when compared to mature stands were consistent with the results reported for basal area and stem density. At the global level the general higher basal areas and lower stem densities in the old-growth stands when compared to mature ones result in a significant higher QMD, that is by nature correlated with the variables from which it is derived. However QMD helps elucidate how basal area is distributed across sizes classes, in this case indicating an allocation weighted more heavily towards larger trees in old-growth.

4.3. Comparison against structural variability in fire-prone latesuccessional forests

Our review was restricted to moist temperate forest ecosystems and therefore excluded forest types classified as temperate, but characterized by lower growing season precipitation, warmer temperatures and, most significantly, high to moderate frequency ground-fire regimes. These systems share some structural characteristics common in moist temperate old-growth, such as abundant large living and dead trees (Covington et al., 1997; Mitchell et al., 2009). However, low to moderate severity fire disturbances result in fundamentally different vertical development, spatial patterning, and CWD accumulation processes (North et al., 2004; Larson and Churchill, 2012).

For example, fire maintained old-growth Ponderosa pine forests (*Pinus ponderosa*) historically had low vertical complexity and open canopies, and widely spaced over-stories (Cooper, 1960). Moderate severity mixed-conifer forests, for example in the Sierra Nevada mountains of California have complex spatial structure (i.e. patchiness) driven by variable burn intensity and associated understory mortality effects (North et al., 2007). This produces a mosaic of both multi- and single-layered patch structures (Kane et al., 2013). As another example, variable vertical structure and multiaged demographics in *Eucalyptus regnans* forests of southeastern Australia have been attributed to fire effects (Lindenmayer et al., 2000). Clearly an expansion of scope to include all temperate forests, particularly drier systems prone to fire, would substantially increase the range of variability in old-growth structure and development described in this review.

4.4. Incomplete data and research needs

Through our literature review we identified gaps in knowledge both in terms of seldom reported relevant structural attributes and of poorly investigated regions. Indeed papers focused on oldgrowth forests often do not report a comprehensive set of structural attributes, including those deemed particularly useful for understanding old-growth forest dynamics and structure. These include, for example, volume of deadwood components, such as logs and snags, vertical and horizontal heterogeneity indices, gap metrics, and tree species diversity indices (Nilsson et al., 2002; Keith et al., 2009; Wirth et al., 2009).

Data on these old-growth indicators are often available in studies focused on a single specific attribute (e.g. deadwood, vertical structure, gaps, etc.), that often do not provide a holistic sense of structure or dynamics (Woldendorp et al., 2004; Zenner, 2004; Schliemann and Bockheim, 2011). We recommend that future studies of old-growth forests provide as wide set of meaningful structural attributes as is possible in order to contribute to a thorough description of these ecosystems.

Through our review we also identified a considerable unevenness of research studies among geographic regions. Old-growth ecosystems in some regions, in particular, seem to be poorly investigated. For instance, though Russian temperate forests are widespread (Silander, 2001), to the best of our knowledge data on old-growth forests are almost completely lacking in English language peer-reviewed journals. Furthermore, in our search we found no or scarce data concerning old-growth temperate forests in Western Asia (Anatolian, Colchic and Hyrcanic region) and for Central Asia in general (e.g. Altai-Sayan and Himalayan regions). We have no data on the occurrence of old-growth remnants in these regions, however, even if few old-growth forests remain, it is important to identify and study forests that are in the most advanced developmental condition, regardless of the successional status of species, and to use them as long-term monitoring areas. We also observed a scarcity of studies comparing age classes in Asia, Australia, and South America that provide a comprehensive characterization of old-growth forest ecosystems. For example, temperate forests in Australia have been extensively investigated (see, for example, Keith et al., 2009, and references therein), yet published structural data were difficult to find. Several studies are rather focused on more specific issues, such as spatial structure, explored extensively in Asia for example (Chen and Bradshaw, 1999; Manabe et al., 2000; Hou et al., 2004; Salas et al., 2006; Hao et al., 2007); regeneration, studied especially in South America (Pollmann and Veblen, 2004; Donoso and Nyland, 2005); deadwood (Schlegel and Donoso, 2008); and forest response to topographical factors (Bale et al., 1998; Barrera et al., 2000).

Based on these gaps in knowledge, future research on old-growth systems needs, above all, to integrate analyses of different ecosystem components and processes. For example, interdisciplinary investigations could better relate old-growth forest structural and compositional dynamics to their carbon storage capacity and biodiversity (e.g.Hatanaka et al., 2011). While a comprehensive body of knowledge is available for some systems, such as *P. menziesii* forests of Pacific Northwest (Franklin et al., 2002; Field and Kaduk, 2004) and northern hardwood forest in U.S. Northeast (Keeton et al., 2007; D'Amato et al., 2008, 2009), it is noticeably incomplete for others.

5. Conclusions

Based on the existing data, our review identified a number of stand structure attributes strongly associated with temperate old-growth forest globally. The published data support a contention that some stand development processes, such as LLT, LAB, and CWD accumulation, have the potential to continue very late into stand development.

Given the commonality in structure among late-successional temperate systems, a consistent set of indicators can be used to describe, map, and understand old-growth globally. However, this set must recognize that structural indicators can be highly dynamic, reflecting varied disturbance processes and successional pathways, as well as climatic and edaphic differences among sites and regions. Consequently, a context-specific analysis, for instance employing a comparative approach involving multiple age classes, is necessary to accommodate the high levels of variability we observed globally.

5.1. Management implications

Our review suggests that temperate old-growth forests globally share characteristics associated with a range of important ecosystem services. As a consequence, conservation of the world's remaining oldgrowth forests and, where appropriate, management to restore oldgrowth to landscapes where they are poorly represented, are required to maintain provisioning of these unique services. Management for 'old-growthedness' (Bauhus et al., 2009) can be incorporated into a variety of forest management scenarios, both commercial and restorative (Keeton, 2006; Gronewold et al., 2010). However, to operationalize this objective, forest managers will need quantitative, regional targets for many of the structural attributes identified in this review. Our review highlighted a minimum LLT density threshold of at least 30 trees with DBH > 50 cm per hectare. This threshold should be emploved as a minimum target for promotion by disturbance-based or 'close-to-nature' silviculture in managed forests. It would not be appropriate as a fixed level to be maintained in stands already displaying old-growth structural attributes, and therefore capable of recruiting higher LLT densities. Crown release is an example of a silvicultural technique that can be used to increase large tree densities (Singer and Lorimer, 1997).

Another key attribute of old-growth forests, both in terms of long-residence carbon accumulation and habitat provisioning, is the amount of CWD. The latter could be increased through silvicultural techniques, such as girdling of selected medium to large sized, low vigor trees, felling or pulling over and leaving some dead or dying trees, where microhabitat requirements related to LLT are already met. These techniques can be employed to varying degrees and in different combinations depending on stand conditions and management objectives. This approach would allow forest managers to build some degree of old-growth associated structure into actively managed stands, while maintaining timber management flexibility (Choi et al., 2007).

Our results contribute to a growing understanding of oldgrowth temperate forests structure across continents and systems. This is critical to enhance forestry practices that maintain both biodiversity associated with structural complexity (e.g. management for microhabitats), and overall favorable conditions for increasing carbon stocks in forest pools, such as large living trees, CWD and soils. Our results will also help inform discussions of the importance of conserving forests with large carbon stocks and providing late-successional habitats.

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Appendix A and B. Supplementary material

Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.foreco.2012.11. 020.

References

- Abrams, M.D., Orwig, D.A., 1996. A 300-year history of disturbance and canopy recruitment for co-occurring white pine and hemlock on the Allegheny Plateau, USA. Journal of Ecology 84, 353–363.
- Abrams, M.D., Orwig, D.A., Dockry, M.J., 1997. Dendroecology and successional status of two contrasting old-growth oak forests in the Blue Ridge mountains, USA. Canadian Journal of Forest Research – Revue Canadienne De Recherche Forestiere 27, 994–1002.
- Abrams, M.D., Ruffner, C.M., DeMeo, T.E., 1998. Dendroecology and species coexistence in an old-growth *Quercus-Acer-Tilia* talus slope forest in the central Appalachians, USA. Forest Ecology and Management 106, 9–18.
- Abrams, M.D., Copenheaver, C.A., Terazawa, K., Umeki, K., Takiya, M., Akashi, N., 1999. A 370-year dendroecological history of an old-growth *Abies-Acer-Quercus* forest in Hokkaido, northern Japan. Canadian Journal of Forest Research – Revue Canadienne De Recherche Forestiere 29, 1891–1899.
- Adams, H.S., Stephenson, S.L., 1989. Old-growth red spruce communities in the mid-appalachians. Vegetatio 85, 45–56.
- Aiba, S., Hill, D.A., Agetsuma, N., 2001. Comparison between old-growth stands and secondary stands regenerating after clear-felling in warm-temperate forests of Yakushima, southern Japan. Forest Ecology and Management 140, 163–175.
- Akhavan, R., Sagheb-Talebi, K., Zenner, E., Safavimanesh, F., 2012. Spatial patterns in different forest development stages of an intact old-growth Oriental beech forest in the Caspian region of Iran. European Journal of Forest Research 1–12. http://dx.doi.org/10.1007/s10342-10012-10603-z.
- Allen, C.D., Macalady, A.K., Chenchouni, H., Bachelet, D., McDowell, N., Vennetier, M., Kitzberger, T., Rigling, A., Breshears, D.D., Hogg, E.H., Gonzalez, P., Fensham, R., Zhang, Z., Castro, J., Demidova, N., Lim, J.H., Allard, G., Running, S.W., Semerci, A., Cobb, N., 2010. A global overview of drought and heat-induced tree mortality reveals emerging climate change risks for forests. Forest Ecology and Management 259, 660–684.
- Armesto, J.J., Figueroa, J., 1987. Stand structure and dynamics in the temperate rainforests of Chiloe Archipelago, Chile. Journal of Biogeography 14, 367–376.
- Aude, E., Lawesson, J.E., 1998. Vegetation in Danish Beech forests: the importance of soil, microclimate and management factors, evaluated by variation partitioning. Plant Ecology 134, 53–65.
- Bale, C.L., Williams, J.B., Charley, J.L., 1998. The impact of aspect on forest structure and floristics in some Eastern Australian sites. Forest Ecology and Management 110, 363–377.
- Barrera, M.D., Frangi, J.L., Richter, L.L., Perdomo, M.H., Pinedo, L.B., 2000. Structural and functional changes in *Nothofagus pumilio* forests along an altitudinal gradient in Tierra del Fuego, Argentina. Journal of Vegetation Science 11, 179– 188.
- Battles, J.J., Armesto, J.J., Vann, D.R., Zarin, D.J., Aravena, J.C., Perez, C., Johnson, A.H., 2002. Vegetation composition, structure, and biomass of two unpolluted watersheds in the Cordillera de Piuchue, Chiloe Island, Chile. Plant Ecology 158, 5–19.
- Bauhus, J., Puettmann, K., Messier, C., 2009. Silviculture for old-growth attributes. Forest Ecology and Management 258, 525–537.
- Beaudet, M., Brisson, J., Gravel, D., Messier, C., 2007. Effect of a major canopy disturbance on the coexistence of *Acer saccharum* and *Fagus grandifolia* in the understorey of an old-growth forest. Journal of Ecology 95, 458–467.
- Bernadzki, E., Bolibok, L., Brzeziecki, B., Zajaczkowski, J., Zybura, H., 1998. Compositional dynamics of natural forests in the Bialowieza National Park, northeastern Poland. Journal of Vegetation Science 9, 229–238.
- Binkley, D., Olsson, U., Rochelle, R., Stohlgren, T., Nikolov, N., 2003. Structure, production and resource use in some old-growth spruce/fir forests in the front range of the Rocky Mountains, USA. Forest Ecology and Management 172, 271– 279.
- Bobiec, A., 2002. Living stands and dead wood in the Bialowieza forest: suggestions for restoration management. Forest Ecology and Management 165, 125–140.
- Boncina, A., 2000. Comparison of structure and biodiversity in the Rajhenav virgin forest remnant and managed forest in the Dinaric region of Slovenia. Global Ecology and Biogeography 9, 201–211.
- Brang, P., 2005. Virgin forests as a knowledge source for central European silviculture: reality or myth? Forest Snow and Landscape Research 79, 19–32.
- Burgman, M.A., 1996. Characterisation and delineation of the eucalypt old-growth forest estate in Australia: a review. Forest Ecology and Management 83, 149– 161.
- Burrascano, S., Lombardi, F., Marchetti, M., 2008. Old-growth forest structure and deadwood: are they indicators of plant species composition? A case study from central Italy. Plant Biosystems 142, 313–323.
- Burrascano, S., Rosati, L., Blasi, C., 2009. Plant species diversity in Mediterranean old-growth forests: a case study from central Italy. Plant Biosystems 143, 190– 200.
- Burrascano, S., Sabatini, F.M., Blasi, C., 2011. Testing indicators of sustainable forest management on understorey composition and diversity in southern Italy through variation partitioning. Plant Ecology 212, 829–841.

- Busing, R.T., 1998. Composition, structure and diversity of cove forest stands in the Great Smoky Mountains: a patch dynamics perspective. Journal of Vegetation Science 9, 881–890.
- Busing, R.T., Fujimori, T., 2005. Biomass, production and woody detritus in an old coast redwood (Sequoia sempervirens) forest. Plant Ecology 177, 177–188.
- Busing, R.T., White, P.S., 1993. Effects of area on old-growth forest attributes implications for the equilibrium landscape concept. Landscape Ecology 8, 119– 126.
- Carmona, M.R., Armesto, J.J., Aravena, J.C., Perez, C.A., 2002. Coarse woody debris biomass in successional and primary temperate forests in Chiloe Island, Chile. Forest Ecology and Management 164, 265–275.
- Chen, J.Q., Bradshaw, G.A., 1999. Forest structure in space: a case study of an old growth spruce-fir forest in Changbaishan Natural Reserve, PR China. Forest Ecology and Management 120, 219–233.
- Cheng, D.L., Wang, G.X., Li, T., Tang, Q.L., Gong, C.M., 2007. Relationships among the stem, aboveground and total biomass across Chinese forests. Journal of Integrative Plant Biology 49, 1573–1579.
- Choi, J., Lorimer, C.G., Vanderwerker, J.M., 2007. A simulation of the development and restoration of old-growth structural features in northern hardwoods. Forest Ecology and Management 249, 204–220.
- Christensen, M., Hahn, K., Mountford, E.P., Odor, P., Standovar, T., Rozenbergar, D., Diaci, J., Wijdeven, S., Meyer, P., Winter, S., Vrska, T., 2005. Dead wood in European beech (*Fagus sylvatica*) forest reserves. Forest Ecology and Management 210, 267–282.
- Commarmot, B., Bachofen, H., Bundziak, Y., Bürgi, A., Shparyk, Y., Sukhariuk, D., Viter, R., Zingg, A., 2005. Structures of virgin and managed beech forests in Uholka (Ukraine) and Sihlwald (Switzerland): a comparative study. Forest Snow and Landscape Research 79, 45–56.
- Cooper, C.F., 1960. Changes in vegetation, structure, and growth of southwestern pine forests since white settlement. Ecological Monographs 30, 129–164.
- Cornwell, W.K., Cornelissen, J.H.C., Allison, S.D., Bauhus, J., Eggleton, P., Preston, C.M., Scarff, F., Weedon, J.T., Wirth, C., Zanne, A.E., 2009. Plant traits and wood fates across the globe: rotted, burned, or consumed? Global Change Biology 15, 2431–2449.
- Covington, W.W., Fule, P.Z., Moore, M.M., Hart, S.C., Kolb, T.E., Mast, J.N., Sackett, S.S., Wagner, M.R., 1997. Restoring ecosystem health in ponderosa pine forests of the southwest. Journal of Forestry 95, 23–29.
- Dahir, S.E., Lorimer, C.G., 1996. Variation in canopy gap formation among developmental stages of northern hardwood stands. Canadian Journal of Forest Research – Revue Canadienne De Recherche Forestiere 26, 1875–1892.
- Dai, L.M., Qi, L., Wang, Q.W., Su, D.K., Yu, D.P., Wang, Y., Ye, Y.J., Jiang, S.W., Zhao, W., 2011. Changes in forest structure and composition on Changbai Mountain in Northeast China. Annals of Forest Science 68, 889–897.
- D'Amato, A.W., Orwig, D.A., Foster, D.R., 2008. The influence of successional processes and disturbance on the structure of *Tsuga canadensis* forests. Ecological Applications 18, 1182–1199.
- D'Amato, A.W., Orwig, D.A., Foster, D.R., 2009. Understory vegetation in old-growth and second-growth *Tsuga canadensis* forests in western Massachusetts. Forest Ecology and Management 257, 1043–1052.
- Das, A., Battles, J., Stephenson, N.L., van Mantgem, P.J., 2011. The contribution of competition to tree mortality in old-growth coniferous forests. Forest Ecology and Management 261, 1203–1213.
- Davis, M.R., Allen, R.B., Clinton, P.W., 2003. Carbon storage along a stand development sequence in a New Zealand Nothofagus forest. Forest Ecology and Management 177, 313–321.
- Donato, D.C., Campbell, J.L., Franklin, J.F., 2012. Multiple successional pathways and precocity in forest development: can some forests be born complex? Journal of Vegetation Science 23, 576–584.
- Donoso, P.J., Lusk, C.H., 2007. Differential effects of emergent Nothofagus dombeyi on growth and basal area of canopy species in an old-growth temperate rainforest. Journal of Vegetation Science 18, 675–684.
- Donoso, P.J., Nyland, R.D., 2005. Seedling density according to structure, dominance and understory cover in old-growth forest stands of the evergreen forest type in the coastal range of Chile. Revista Chilena De Historia Natural 78, 51–63.
- Druckenbrod, D.L., Shugart, H.H., Davies, I., 2005. Spatial pattern and process in forest stands within the Virginia piedmont. Journal of Vegetation Science 16, 37–48.
- Fajardo, A., de Graaf, R., 2004. Tree dynamics in canopy gaps in old-growth forests of Nothofagus pumilio in Southern Chile. Plant Ecology 173, 95–105.
- FAO, 2010. Global Forest Resources Assessment 2010. Main Report. FAO Forestry Paper. FAO, Rome, p. 340.
- Field, C.B., Kaduk, J., 2004. The carbon balance of an old-growth forest: building across approaches. Ecosystems 7, 525–533.
- Fisk, M.C., Zak, D.R., Crow, T.R., 2002. Nitrogen storage and cycling in old- and second-growth northern hardwood forests. Ecology 83, 73–87.
- Foster, B.C., Wang, D., Keeton, W.S., 2008. An exploratory, post-harvest comparison of ecological and economic characteristics of forest stewardship council certified and uncertified northern hardwood stands. Journal of Sustainable Forestry 26, 171–191.
- Franklin, J.F., Cromack, K., Denison, W., McKee, A., Maser, C., Sedell, J., Swanson, F., Juday, G., 1981. Ecological characteristics of old-growth Douglas-fir forests. In: General Technical Report PNW-118. US Department of Agriculture, Portland, OR, USA.
- Franklin, J.F., Shugart, H.H., Harmon, M.E., 1987. Tree death as an ecological process. BioScience 37, 550–556.

- Franklin, J.F., Spies, T.A., Van Pelt, R., Carey, A.B., Thornburgh, D.A., Berg, D.R., Lindenmayer, D.B., Harmon, M.E., Keeton, W.S., Shaw, D.C., Bible, K., Chen, J.Q., 2002. Disturbances and structural development of natural forest ecosystems with silvicultural implications, using Douglas-Fir forests as an example. Forest Ecology and Management 155, 399–423.
- Goebel, P.C., Hix, D.M., 1996. Development of mixed-oak forests in southeastern Ohio: a comparison of second-growth and old-growth forests. Forest Ecology and Management 84, 1–21.
- Goodburn, J.M., Lorimer, C.G., 1998. Cavity trees and coarse woody debris in oldgrowth and managed northern hardwood forests in Wisconsin and Michigan. Canadian Journal of Forest Research – Revue Canadienne De Recherche Forestiere 28, 427–438.
- Gronewold, C.A., D'Amato, A.W., Palik, B.J., 2010. The influence of cutting cycle and stocking level on the structure and composition of managed old-growth northern hardwoods. Forest Ecology and Management 259, 1151–1160.
- Grove, S.J., 2002. Saproxylic insect ecology and the sustainable management of forests. Annual Review of Ecology and Systematics 33, 1–23.
- Gutierrez, A.G., Armesto, J.J., Aravena, J.C., 2004. Disturbance and regeneration dynamics of an old-growth North Patagonian rain forest in Chiloe Island, Chile. Journal of Ecology 92, 598–608.
- Gutierrez, A.G., Armesto, J.J., Aravena, J.C., Carmona, M., Carrasco, N.V., Christie, D.A., Pena, M.P., Perez, C., Huth, A., 2009. Structural and environmental characterization of old-growth temperate rainforests of northern Chiloe Island, Chile: regional and global relevance. Forest Ecology and Management 258, 376–388.
- Harmon, M.E., Franklin, J.F., Swanson, F.J., Sollins, P., Gregory, S.V., Lattin, J.D., Anderson, N.H., Cline, S.P., Aumen, N.G., Sedell, J.R., Lienkaemper, G.W., Cromack Jr., K., Cummin, K.W., 1986. Ecology of Coarse Woody Debris in Temperate Ecosystems. Advances in Ecological Research 15, 133–302.
- Hao, Z.Q., Zhang, J., Song, B., Ye, J., Li, B.H., 2007. Vertical structure and spatial associations of dominant tree species in an old-growth temperate forest. Forest Ecology and Management 252, 1–11.
- Hatanaka, N., Wright, W., Loyn, R.H., Mac Nally, R., 2011. 'Ecologically complex carbon' – linking biodiversity values, carbon storage and habitat structure in some austral temperate forests. Global Ecology and Biogeography 20, 260–271.
- Heiri, C., Wolf, A., Rohrer, L., Bugmann, H., 2009. Forty years of natural dynamics in Swiss beech forests: structure, composition, and the influence of former management. Ecological Applications 19, 1920–1934.
- Holeksa, J., Saniga, M., Szwagrzyk, J., Dziedzic, T., Ferenc, S., Wodka, M., 2007. Altitudinal variability of stand structure and regeneration in the subalpine spruce forests of the Pol'ana biosphere reserve, Central Slovakia. European Journal of Forest Research 126, 303–313.
- Holeksa, J., Saniga, M., Szwagrzyk, J., Czerniak, M., Staszynska, K., Kapusta, P., 2009. A giant tree stand in the West Carpathians – an exception or a relic of formerly widespread mountain European forests? Forest Ecology and Management 257, 1577–1585.
- Hoover, C.M., Leak, W.B., Keel, B.G., 2012. Benchmark carbon stocks from oldgrowth forests in northern New England, USA. Forest Ecology and Management 266, 108–114.
- Hoshino, D., Nishimura, N., Yamamoto, S., 2001. Age, size structure and spatial pattern of major tree species in an old-growth *Chamaecyparis obtusa* forest, Central Japan. Forest Ecology and Management 152, 31–43.
- Hou, J.H., Mi, X.C., Liu, C.R., Ma, K.P., 2004. Spatial patterns and associations in a Quercus-Betula forest in northern China. Journal of Vegetation Science 15, 407– 414.
- Houghton, R.A., 2005. Aboveground forest biomass and the global carbon balance. Global Change Biology 11, 945–958.
- Hudiburg, T., Law, B., Turner, D.P., Campbell, J., Donato, D.C., Duane, M., 2009. Carbon dynamics of Oregon and Northern California forests and potential landbased carbon storage. Ecological Applications 19, 163–180.
- Hunter Jr., M.L., 1999. Maintaining biodiversity in forest ecosystems. Maintaining biodiversity in forest ecosystems, vol. i–xiv, pp. 1–698.
- Innes, J.L., 1992. Structure of evergreen temperate rain forest on the Taitao Peninsula, southern Chile. Journal of Biogeography 19, 555–562.
- Jones, E.W., 1945. The structure and reproduction of the virgin forest of the north temperate zone. New Phytologist 44, 130–148.
- Kane, V.R., Lutz, J.A., Roberts, S.L., Smith, D.F., McGaughey, R.J., Povak, N.A., Brooks, M.L., 2013. Landscape-scale effects of fire severity on mixed-conifer and red fir forest structure in Yosemite National Park. Forest Ecology and Management 287, 17–31.
- Keddy, P.A., Drummond, C.G., 1996. Ecological properties for the evaluation, management, and restoration of temperate deciduous forest ecosystems. Ecological Applications 6, 748–762.
- Keeton, W.S., 2006. Managing for late-successional/old-growth characteristics in northern hardwood-conifer forests. Forest Ecology and Management 235, 129– 142.
- Keeton, W.S., Franklin, J.F., 2005. Do remnant old-growth trees accelerate rates of succession in mature Douglas-fir forests? Ecological Monographs 75, 103–118.
- Keeton, W.S., Kraft, C.E., Warren, D.R., 2007. Mature and old-growth riparian forests: structure, dynamics, and effects on adirondack stream habitats. Ecological Applications 17, 852–868.
- Keeton, W.S., Chernyavskyy, M., Gratzer, G., Main-Knorn, M., Shpylchak, M., Bihun, Y., 2010. Structural characteristics and aboveground biomass of old-growth spruce-fir stands in the eastern Carpathian mountains, Ukraine. Plant Biosystems 144, 148–159.

- Keeton, W.S., Whitman, A.A., McGee, G.C., Goodale, C.L., 2011. Late-successional biomass development in Northern Hardwood-Conifer forests of the Northeastern United States. Forest Science 57, 489–505.
- Keith, H., Mackey, B.G., Lindenmayer, D.B., 2009. Re-evaluation of forest biomass carbon stocks and lessons from the world's most carbon-dense forests. Proceedings of the National academy of Sciences of the United States of America 106, 11635–11640.
- Knorn, J., Kuemmerle, T., Radeloff, V.C., Keeton, W.S., Gancz, V., Biris, I.A., Svoboda, M., Griffiths, P., Hagatis, A., Hostert, P., in press. Continued loss of temperate old-growth forests in the Romanian Carpathians despite an increasing protected area network. Environmental Conservation. http://dx.doi.org/ 10.1017.S0376892912000355.
- Král, K., Janik, D., Vrska, T., Adam, D., Hort, L., Unar, P., Samonil, P., 2010. Local variability of stand structural features in beech dominated natural forests of Central Europe: implications for sampling. Forest Ecology and Management 260, 2196–2203.
- Kucbel, S., Saniga, M., Jaloviar, P., Vencurik, J., 2012. Stand structure and temporal variability in old-growth beech-dominated forests of the northwestern Carpathians: a 40-years perspective. Forest Ecology and Management 264, 125–133.
- Kupfer, J.A., Kirsch, S.W., 1998. Heterogeneity of forest characteristics in primary and secondary forest stands on the third Chickasaw loess bluff, Tennessee. Physical Geography 19, 35–54.
- Lamedica, S., Lingua, E., Popa, I., Motta, R., Carrer, M., 2011. Spatial structure in four Norway spruce stands with different management history in the Alps and Carpathians. Silva Fennica 45, 865–873.
- Larson, A.J., Churchill, D., 2012. Tree spatial patterns in fire-frequent forests of western North America, including mechanisms of pattern formation and implications for designing fuel reduction and restoration treatments. Forest Ecology and Management 267, 74–92.
- Larsson, T.B., 2001. Biodiversity evaluation tools for European forests. In: Franc, A., Laroussinie, O., Karjalainen, T. (Eds.), Criteria and Indicators for Sustainable Forest Management at the Forest Management Unit Level, pp. 75–81.
- Lassauce, A., Paillet, Y., Jactel, H., Bouget, C., 2011. Deadwood as a surrogate for forest biodiversity: meta-analysis of correlations between deadwood volume and species richness of saproxylic organisms. Ecological Indicators 11, 1027– 1039.
- Lindenmayer, D.B., Cunningham, R.B., Donnelly, C.F., Franklin, J.F., 2000. Structural features of old-growth Australian montane ash forests. Forest Ecology and Management 134, 189–204.
- Lusk, C.H., Smith, B., 1998. Life history differences and tree species coexistence in an old-growth New Zealand rain forest. Ecology 79, 795–806.
- Luyssaert, S., Schulze, E.D., Borner, A., Knohl, A., Hessenmoller, D., Law, B.E., Ciais, P., Grace, J., 2008. Old-growth forests as global carbon sinks. Nature 455, 213–215.
- Mackensen, J., Bauhus, J., Webber, E., 2003. Decomposition rates of coarse woody debris – a review with particular emphasis on Australian tree species. Australian Journal of Botany 51, 27–37.
- Manabe, T., Nishimura, N., Miura, M., Yamamoto, S., 2000. Population structure and spatial patterns for trees in a temperate old-growth evergreen broad-leaved forest in Japan. Plant Ecology 151, 181–197.
- Marage, D., Lemperiere, G., 2005. The management of snags: a comparison in managed and unmanaged ancient forests of the Southern French Alps. Annals of Forest Science 62, 135–142.
- Martínez Pastur, G., Cellini, J.M., Peri, P.L., Vukasovic, R.F., Fernández, M.C., 2000. Timber production of *Nothofagus pumilio* forests by a shelterwood system in Tierra del Fuego (Argentina). Forest Ecology and Management 134, 153–162.
- McCarthy, B.C., Hammer, C.A., Kauffman, G.L., Cantino, P.D., 1987. Vegetation patterns and structure of an old-growth forest in southeastern Ohio. Bulletin of the Torrey Botanical Club 114, 33–45.
- McCarthy, B.C., Small, C.J., Rubino, D.L., 2001. Composition, structure and dynamics of Dysart Woods, an old-growth mixed mesophytic forest of southeastern Ohio. Forest Ecology and Management 140, 193–213.
- McGee, G.G., Leopold, D.J., Nyland, R.D., 1999. Structural characteristics of oldgrowth, maturing, and partially cut northern hardwood forests. Ecological Applications 9, 1316–1329.
- McKenny, H.C., Keeton, W.S., Donovan, T.M., 2006. Effects of structural complexity enhancement on eastern red-backed salamander (*Plethodon cinereus*) populations in northern hardwood forests. Forest Ecology and Management 230, 186–196.
- MCPFE, 2003. Improved pan-European indicators for sustainable forest management as adopted by the MCPFE Expert Level Meeting. In: Ministerial Conference on the Protection of Forests in Europe, Vienna.
- Merino, A., Real, C., Alvarez-Gonzalez, J.G., Rodriguez-Guitian, M.A., 2007. Forest structure and C stocks in natural *Fagus sylvatica* forest in southern Europe: the effects of past management. Forest Ecology and Management 250, 206–214.
- Mikusinski, G., Angelstam, P., 1998. Economic geography, forest distribution, and woodpecker diversity in central Europe. Conservation Biology 12, 200–208.
- Mitchell, R., Engstrom, T., Sharitz, R.R., De Steven, D., Hiers, K., Cooper, R., Kirkman, L.K., 2009. Old forests and endangered woodpeckers: old-growth in the southern coastal plain. Natural Areas Journal 29, 301–310.
- Motta, R., Berretti, R., Castagneri, D., Lingua, E., Nola, P., Vacchiano, G., 2010. Stand and coarse woody debris dynamics in subalpine Norway spruce forests withdrawn from regular management. Annals of Forest Science 67.
- Motta, R., Berretti, R., Castagneri, D., Dukic, V., Garbarino, M., Govedar, Z., Lingua, E., Maunaga, Z., Meloni, F., 2011. Toward a definition of the range of variability of

central European mixed *Fagus-Abies-Picea* forests: the nearly steady-state forest of Lom (Bosnia and Herzegovina). Canadian Journal of Forest Research – Revue Canadienne De Recherche Forestiere 41, 1871–1884.

- Nagel, T.A., Svoboda, M., Diaci, J., 2006. Regeneration patterns after intermediate wind disturbance in an old-growth *Fagus-Abies* forest in southeastern Slovenia. Forest Ecology and Management 226, 268–278.
- Nagel, T.A., Svoboda, M., Rugani, T., Diaci, J., 2010. Gap regeneration and replacement patterns in an old-growth *Fagus-Abies* forest of Bosnia-Herzegovina. Plant Ecology 208, 307–318.
- Narukawa, Y., Yamamoto, S.I., 2001. Gap formation, microsite variation and the conifer seedling occurrence in a subalpine old-growth forest, central Japan. Ecological Research 16, 617–625.
- Nilsson, S.G., Niklasson, M., Hedin, J., Aronsson, G., Gutowski, J.M., Linder, P., Ljungberg, H., Mikusinski, G., Ranius, T., 2002. Densities of large living and dead trees in old-growth temperate and boreal forests. Forest Ecology and Management 161, 189–204.
- North, M.P., Keeton, W.S., 2008. Emulating natural disturbance regimes: an emerging approach for sustainable forest management. In: Lafortezza, R., Chen, J., Sanesi, G., Crow, T.R. (Eds.), Patterns and Processes in Forest Landscapes – Multiple Use and Sustainable Management. Springer, The Netherlands, pp. 341–372.
- North, M., Chen, J.Q., Oakley, B., Song, B., Rudnicki, M., Gray, A., Innes, J., 2004. Forest stand structure and pattern of old-growth western hemlock/Douglas-fir and mixed-conifer forests. Forest Science 50, 299–311.
- North, M., Innes, J., Zald, H., 2007. Comparison of thinning and prescribed fire restoration treatments to Sierran mixed-conifer historic conditions. Canadian Journal of Forest Research – Revue Canadienne De Recherche Forestiere 37, 331–342.
- Odor, P., Heilmann-Clausen, J., Christensen, M., Aude, E., van Dort, K.W., Piltaver, A., Siller, I., Veerkamp, M.T., Walleyn, R., Standovar, T., van Hees, A.F.M., Kosec, J., Matocec, N., Kraigher, H., Grebenc, T., 2006. Diversity of dead wood inhabiting fungi and bryophytes in semi-natural beech forests in Europe. Biological Conservation 131, 58–71.
- Oliver, C.D., Larson, B.C., 1990. Forest Stand Dynamics. McGraw-Hill, New York.
- Olson, D.M., Dinerstein, E., Wikramanayake, E.D., Burgess, N.D., Powell, G.V.N., Underwood, E.C., D'Amico, J.A., Itoua, I., Strand, H.E., Morrison, J.C., Loucks, C.J., Allnutt, T.F., Ricketts, T.H., Kura, Y., Lamoreux, J.F., Wettengel, W.W., Hedao, P., Kassem, K.R., 2001. Terrestrial ecoregions of the worlds: a new map of life on Earth. Bioscience 51, 933–938.
- Orwig, D.A., Cogbill, C.V., Foster, D.R., O'Keefe, J.F., 2001. Variations in old-growth structure and definitions: forest dynamics on Wachusett Mountain, Massachusetts. Ecological Applications 11, 437–452.
- Paluch, J.G., 2007. The spatial pattern of a natural European beech (*Fagus sylvatica* L.)-silver fir (*Abies alba* Mill.) forest: a patch-mosaic perspective. Forest Ecology and Management 253, 161–170.
- Pan, Y.D., Birdsey, R.A., Fang, J.Y., Houghton, R., Kauppi, P.E., Kurz, W.A., Phillips, O.L., Shvidenko, A., Lewis, S.L., Canadell, J.G., Ciais, P., Jackson, R.B., Pacala, S.W., McGuire, A.D., Piao, S.L., Rautiainen, A., Sitch, S., Hayes, D., 2011. A large and persistent carbon sink in the world's forests. Science 333, 988–993.
- Parker, G.R., Leopold, D.J., Eichenberger, J.K., 1985. Tree dynamics in an old-growth, deciduous forest. Forest Ecology and Management 11, 31–57.
- Persiani, A.M., Audisio, P., Lunghini, D., Maggi, O., Granito, V.M., Biscaccianti, A.B., Chiavetta, U., Marchetti, M., 2010. Linking taxonomical and functional biodiversity of saproxylic fungi and beetles in broad-leaved forests in southern Italy with varying management histories. Plant Biosystems 144, 250–261.
- Peterken, G.F., 1996. Natural Woodland: Ecology and Conservation in Northern Temperate Regions. Cambridge University Press, Cambridge UK.
- Petritan, A.M., Biris, I.A., Merce, O., Turcu, D.O., Petritan, I.C., 2012. Structure and diversity of a natural temperate sessile oak (*Quercus petraea* L.) – European Beech (*Fagus sylvatica* L.) forest. Forest Ecology and Management 280, 140–149. Pollmann, W., 2003. Stand structure and dendroecology of an old-growth
- Pollmann, W., 2003. Stand structure and dendroecology of an old-growth Nothofagus forest in Conguillio National Park, south Chile. Forest Ecology and Management 176, 87–103.
- Pollmann, W., Veblen, T.T., 2004. Nothofagus regeneration dynamics in southcentral Chile: a test of a general model. Ecological Monographs 74, 615–634.
- R Development Core Team, 2010. R: a language and environment for statistical computing. R Foundation for Statistical Computing, Vienna, Austria.
- Rebertus, A.J., Veblen, T.T., 1993. Structure and tree-fall gap dynamics of old-growth nothofagus forests in tierra-del-fuego, argentina. Journal of Vegetation Science 4, 641–654.
- Rhemtulla, J.M., Mladenoff, D.J., Clayton, M.K., 2009. Historical forest baselines reveal potential for continued carbon sequestration. Proceedings of the National Academy of Sciences of the United States of America 106, 6082–6087.
- Rozas, V., 2003. Regeneration patterns, dendroecology, and forest-use history in an old-growth beech-oak lowland forest in Northern Spain. Forest Ecology and Management 182, 175–194.
- Ruffell, J., Sedgeley, J., Parsons, S., 2007. The potential availability of roosting sites for lesser short-tailed bats (*Mystacina tuberculata*) on Kapiti Island, New Zealand: implications for a translocation. New Zealand Journal of Zoology 34, 219–226.
- Ruffner, C.M., Abrams, M.D., 2003. Disturbance history and stand dynamics along a topographic gradient in old-growth hemlock-northern Hardwood Forests of the Allegheny Plateau, USA. Natural Areas Journal 23, 98–113.
- Salas, C., Garcia, O., 2006. Modelling height development of mature Nothofagus obliqua. Forest Ecology and Management 229, 1–6.

- Salas, C., Lemay, V., Nunez, P., Pacheco, P., Espinosa, A., 2006. Spatial patterns in an old-growth Nothofagus obliqua forest in south-central Chile. Forest Ecology and Management 231, 38–46.
- Sano, J., 1997. Age and size distribution in a long-term forest dynamics. Forest Ecology and Management 92, 39–44.
- Sato, T., 2010. Stocks of coarse woody debris in old-growth lucidophyllous forests in southwestern Japan. Journal of Forest Research 15, 404–410.
- Schlegel, B.C., Donoso, P.J., 2008. Effects of forest type and stand structure on coarse woody debris in old-growth rainforests in the Valdivian Andes, south-central Chile. Forest Ecology and Management 255, 1906–1914.
- Schliemann, S.A., Bockheim, J.G., 2011. Methods for studying treefall gaps: a review. Forest Ecology and Management 261, 1143–1151.
- Schnitzler, A., Borlea, F., 1998. Lessons from natural forests as keys for sustainable management and improvement of naturalness in managed broadleaved forests. Forest Ecology and Management 109, 293–303.
- Siitonen, J., 2001. Forest management, coarse woody debris and saproxylic organisms: Fennoscandian boreal forests as an example. Ecological Bulletins 49.
- Silander, J.A., 2001. Temperate forests. In: Levin, S.A. (Ed.), Encyclopedia of Biodiversity. Princeton University, New Jersey, USA, pp. 607–626.
- Sillett, S.C., Van Pelt, R., 2007. Trunk reiteration promotes epiphytes and water storage in an old-growth redwood forest canopy. Ecological Monographs 77, 335–359.
- Singer, M.T., Lorimer, C.G., 1997. Crown release as a potential old-growth restoration approach in northern hardwoods. Canadian Journal of Forest Research – Revue Canadienne De Recherche Forestiere 27, 1222–1232.
- Smale, M.C., Burns, B.R., Smale, P.N., Whaley, P.T., 1997. Dynamics of upland podocarp/broadleaved forest on Mamaku Plateau, central North Island, New Zealand. Journal of the Royal Society of New Zealand 27, 513–532.
- Spies, T.A., 2004. Ecological concepts and diversity of old-growth forests. Journal of Forestry 102, 14–20.
- Spies, T.A., Franklin, J.F., 1991. The structure of natural young, mature, and oldgrowth Douglas-fir forests in Oregon and Washington. Wildlife and Vegetation of Unmanaged Douglas-Fir Forests 285, 91–109.
- Stegen, J.C., Swenson, N.G., Enquist, B.J., White, E.P., Phillips, O.L., Jorgensen, P.M., Weiser, M.D., Mendoza, A.M., Vargas, P.N., 2011. Variation in above-ground forest biomass across broad climatic gradients. Global Ecology and Biogeography 20, 744–754.
- Stephenson, N.L., van Mantgem, P.J., Bunn, A.G., Bruner, H., Harmon, M.E., O'Connell, K.B., Urban, D.L., Franklin, J.F., 2011. Causes and implications of the correlation between forest productivity and tree mortality rates. Ecological Monographs 81, 527–555.
- Svoboda, M., Pouska, V., 2008. Structure of a Central-European mountain spruce old-growth forest with respect to historical development. Forest Ecology and Management 255, 2177–2188.
- Taylor, A.H., Halpern, C.B., 1991. The structure and dynamics of Abies magnifica forests in the Southern Cascade Range, Usa. Journal of Vegetation Science 2, 189–200.
- Taylor, A.H., Jang, S.W., Zhao, L.J., Liang, C.P., Miao, C.J., Huang, J.Y., 2006. Regeneration patterns and tree species coexistence in old-growth Abies-Picea forests in southwestern China. Forest Ecology and Management 223, 303–317.
- Travaglini, D., Paffetti, D., Bianchi, L., Bottacci, A., Bottalico, F., Giovannini, G., Maltoni, A., Nocentini, S., Vettori, C., Calamini, G., 2012. Characterization, structure and genetic dating of an old-growth beech-fir forest in the northern Apennines (Italy). Plant Biosystems 146, 175–188.

- Tyrrell, L.E., Crow, T.R., 1994a. Dynamics of dead wood in old-growth Hemlock Hardwood forests of Northern Wisconsin and Northern Michigan. Canadian Journal of Forest Research – Revue Canadienne De Recherche Forestiere 24, 1672–1683.
- Tyrrell, L.E., Crow, T.R., 1994b. Structural characteristics of old-growth Hemlock-Hardwood forests in relation to age. Ecology 75, 370–386.
- von Oheimb, G., Westphal, C., Tempel, H., Hardtle, W., 2005. Structural pattern of a near-natural beech forest (*Fagus sylvatica*) (Serrahn, North-east Germany). Forest Ecology and Management 212, 253–263.
- Walcroft, A.S., Brown, K.J., Schuster, W.S.F., Tissue, D.T., Turnbull, M.H., Griffin, K.L., Whitehead, D., 2005. Radiative transfer and carbon assimilation in relation to canopy architecture, foliage area distribution and clumping in a mature temperate rainforest canopy in New Zealand. Agricultural and Forest Meteorology 135, 326–339.
- Waring, R.H., Franklin, J.F., 1979. Evergreen Coniferous Forests of the Pacific Northwest. Science 204, 1380–1386.
- Wells, R.W., Lertzman, K.P., Saunders, S.C., 1998. Old-growth definitions for the forests of British Columbia, Canada. Natural Areas Journal 18, 279–292.
- Whitman, A.A., Hagan, J.M., 2007. An index to identify late-successional forest in temperate and boreal zones. Forest Ecology and Management 246, 144–154.
- Wijdeven, S.M.J., 2004. Stand dynamics in Fontainebleu. Dynamics in beech forest structure and composition over 17 years in La Tillaie forest reserve, Fontainebleu, France. In: Aterra-rapport 1124. Alterra, Wageningen.
- Williams, C.E., Cook, R.E., 2010. Forest Seral stage and large woody debris abundance in an Allegheny high plateau forest-stream system. Natural Areas Journal 30, 20–26.
- Winter, S., Moller, G.C., 2008. Microhabitats in lowland beech forests as monitoring tool for nature conservation. Forest Ecology and Management 255, 1251–1261.
- Wirth, C., Gleixner, G., Heimann, M. (Eds.), 2009. Old-growth Forests: Function, Fate and Value. Springer-Verlag, Berlin Heidelberg.
- Woldendorp, G., Keenan, R.J., Barry, S., Spencer, R.D., 2004. Analysis of sampling methods for coarse woody debris. Forest Ecology and Management 198, 133– 148.
- Xu, C.Y., Turnbull, M.H., Tissue, D.T., Lewis, J.D., Carson, R., Schuster, W.S.F., Whitehead, D., Walcroft, A.S., Li, J.B., Griffin, K.L., 2012. Age-related decline of stand biomass accumulation is primarily due to mortality and not to reduction in NPP associated with individual tree physiology, tree growth or stand structure in a Quercus-dominated forest. Journal of Ecology 100, 428–440.
- Yamamoto, S., Nishimura, N., 1999. Canopy gap formation and replacement pattern of major tree species among developmental stages of beech (*Fagus crenata*) stands, Japan. Plant Ecology 140, 167–176.
- Yamamoto, S., Nishimura, N., Matsui, K., 1995. Natural disturbance and tree species coexistence in an old-growth beech – Dwarf bamboo forest, southwestern Japan. Journal of Vegetation Science 6, 875–886.
- Zell, J., Kandler, G., Hanewinkel, M., 2009. Predicting constant decay rates of coarse woody debris – a meta-analysis approach with a mixed model. Ecological Modelling 220, 904–912.
- Zenner, E.K., 2004. Does old-growth condition imply high live-tree structural complexity? Forest Ecology and Management 195, 243–258.
- Zhou, G.Y., Liu, S.G., Li, Z., Zhang, D.Q., Tang, X.L., Zhou, C.Y., Yan, J.H., Mo, J.M., 2006. Old-growth forests can accumulate carbon in soils. Science 314, 1417.
- Zielonka, T., 2006. Quantity and decay stages of coarse woody debris in old-growth subalpine spruce forests of the western Carpathians, Poland. Canadian Journal of Forest Research – Revue Canadienne De Recherche Forestiere 36, 2614– 2622.