

Title	Long-term cumulative impacts of windthrow and subsequent management on tree species composition and aboveground biomass : A simulation study considering regeneration on downed logs
Author(s)	Hotta, Wataru; Morimoto, Junko; Haga, Chihiro; Suzuki, Satoshi N.; Inoue, Takahiro; Matsui, Takanori; Owari, Toshiaki; Shibata, Hideaki; Nakamura, Futoshi
Citation	Forest ecology and management, 502, 119728 https://doi.org/10.1016/j.foreco.2021.119728
Issue Date	2021-12-15
Doc URL	http://hdl.handle.net/2115/90925
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Rights(URL)	http://creativecommons.org/licenses/by-nc-nd/4.0/
Туре	article (author version)
File Information	hotta_2021_FEM_accepted_manuscript.pdf



1	Title: Long-term cumulative impacts of windthrow and subsequent management on tree
2	species composition and aboveground biomass: A simulation study considering regeneration
3	on downed logs
4	Authors: Wataru Hotta ^{a,*} , Junko Morimoto ^a , Chihiro Haga ^b , Satoshi N. Suzuki ^c , Takahiro
5	Inoue ^d , Takanori Matsui ^b , Toshiaki Owari ^c , Hideaki Shibata ^d , Futoshi Nakamura ^a
6	
7	Affiliations:
8	a. Graduate School of Agriculture, Hokkaido University, Sapporo, Hokkaido 060-8587, Japan
9	b. Graduate School of Engineering, Osaka University, Suita, Osaka 565-0871, Japan
10	c. The University of Tokyo Hokkaido Forest, Graduate School of Agricultural and Life
11	Sciences, The University of Tokyo, Furano, Hokkaido 079-1563, Japan
12	d. Field Science Center for Northern Biosphere, Hokkaido University, Sapporo, Hokkaido
13	060-0809, Japan
14	
15	*Corresponding author: W. Hotta
16	E-mail: w-hotter97thank-you@eis.hokudai.ac.jp
17	TEL: +81-11-706-3339
18	Postal Address: Graduate School of Agriculture, Hokkaido University, Kita 9, Nishi 9, Kita ku,
19	Sapporo, Hokkaido 060-8587, Japan
20	

21 Abstract

Post-windthrow management delays forest biomass recovery by altering the situation 22of disturbance legacies and can change the species composition. Although the short-term 23effects of post-windthrow management have been well studied, we do not have enough 24knowledge about the long-term effects of post-windthrow management on species 25composition and biomass recovery. Those effects associated with an increase in the 2627windthrow frequency are also unknown. Although forest landscape models can effectively evaluate these effects, conventional models do not represent the regeneration process on 2829downed logs, which is essential for simulating forest succession. We focused on hemiboreal forests in northern Japan and aimed to (1) incorporate the regeneration process on downed 30 31logs into LANDIS-II, which is one of the most used forest landscape models; (2) evaluate the long-term effects of post-windthrow management on tree species composition and 32aboveground biomass recovery; and (3) evaluate the associated long-term effects of 33 interactions between post-windthrow management and increased windthrow frequency. We 34incorporated the regeneration process on downed logs into LANDIS-II by regulating the 35probability of the establishment of species that depend on dead wood, such as spruce, 36 according to the availability of well-decayed dead wood. The incorporation of this process 37 resulted in simulations of trends in species composition and aboveground biomass recovery 38after post-windthrow management that were more accurate than those produced by the 39original model. In the modified LANDIS-II simulation, reductions in dead wood and 40

41	advanced seedlings due to salvage logging had little effect on the tree species composition or
42	aboveground biomass recovery; however, the complete destruction of advanced seedlings by
43	scarification induced a delay in aboveground biomass recovery and a shift to birch-dominated
44	forests that continued for 100 years. In addition, the reduction in dead wood due to salvage
45	logging decreased the number of seedlings, especially of dead wood-dependent species, that
46	established after windthrow. When the windthrow frequency doubled, this decrease in
47	seedlings induced a delay in aboveground biomass recovery, and a substantial decrease in
48	dead wood-dependent species biomass occurred after a subsequent windthrow event.
49	However, after the second windthrow event and following scarification, the forest recovered
50	in the same way as after the first windthrow because the destruction of advanced seedlings
51	and understory plants, namely, dwarf bamboo (Sasa spp.), by scarification reset the site
52	conditions. To conserve the species composition and aboveground biomass of hemiboreal
53	forests under climate change, which is expected to increase windthrow frequency, salvage
54	logging and scarification should be avoided.
~ ~	(208/400 - monds)

- 55 (**398/400 words**)
- 56

57 Keywords:

Salvage logging; Scarification; Decayed downed logs; Regeneration; LANDIS-II; Forest
landscape model

60 1. Introduction

Windthrow is a major natural disturbance in temperate and boreal forest ecosystems 61in East Asia (Nakashizuka 1989; Yamamoto 1989) that causes mass deaths of live trees and 62temporary decreases in forest biomass. Post-windthrow managements delay the recovery of 63 biomass by altering the situation of disturbance legacies and is likely to change species 64 composition. Post-windthrow managements, such as salvage logging and scarification, affect 6566 forest recovery by creating additional disturbances in forests that have been damaged by windthrow (Leverkus et al. 2018). Salvage logging is a conventional forest management 67 68 strategy used after natural disturbances and removes dead wood to compensate for economic losses and to prevent additional wildfire and insect outbreaks (Lindenmayer et al. 2008). 69 70 Additionally, salvaged sites are sometimes scarified to destroy grasses and shrubs preventing tree regenerations and to remove the organic-rich surface soil layer, which contains pathogens 7172(Yoshida et al. 2005). All fallen trees are salvaged after severe windthrows in Japanese national forests, and sites are scarified as necessary (Director-General of Japanese forest 73agency 2011). However, post-windthrow management has significant impacts on the species 74composition and recovery of aboveground biomass in forests developing after windthrow by 75altering situation of disturbance legacies, such as advanced regeneration and dead wood (Kurz 76et al. 2008; Taeroe et al. 2019). 77

The long-term effects of post-windthrow management on species composition and
biomass recovery are unknown, although its short- and mid-term effects have been well

80	studied. A short- and mid-term study that measured the effects of salvage logging and
81	scarification on forests 3-40 years after windthrow reported a delay in forest regeneration
82	(Donato et al. 2006; Greene et al. 2006; Morimoto et al. 2011) and an increase in the
83	proportions of early successional species such as Betula spp. (Ilisson et al. 2007; Fischer and
84	Fischer 2012) due to a lack of advanced regeneration. Specifically, even-aged forests of
85	Betula spp. often develop after scarification (Yoshida et al. 2005; Prévost et al. 2010; Aoyama
86	et al. 2011; Suzuki 2020). Long-term (over 50 years after windthrow) studies also reported an
87	increase in broadleaved trees in salvaged stands (Morimoto et al. 2019a; Hotta et al. 2020).
88	On the other hand, some studies reported that the species composition of the canopy layer did
89	not differ according to whether salvage logging was performed (Sass et al. 2018). In addition,
90	Fischer et al. (2002) simulated the long-term effects of salvage logging on tree species
91	composition in a simplified manner and reported that <i>Betula</i> spp. are dominant in the first 30
92	years after windthrow and that afterward, the dominant species are gradually replaced by
93	Picea spp. These results indicate that we still do not have a consensus about the effects of
94	post-windthrow management on the long-term changes in species composition.
95	The effects of multiple windthrows should be considered to understand the long-term
96	effects of post-windthrow management on forests. This is because post-windthrow
97	management can substantially affect the quantity and species composition of advanced
98	regeneration that develops after windthrow by altering microsite conditions and the post-
99	windthrow canopy species composition (Waldron et al. 2014). In particular, salvage logging

100	decreases the amount of dead wood in forests after windthrow (Suzuki et al. 2019; Morimoto
101	et al. 2019a; Hotta et al. 2020); therefore, tree species that establish only on well-decayed
102	dead wood could decrease in the long term. Because advanced seedlings greatly contribute to
103	forest recovery after windthrow, differences in seedling species composition could also affect
104	the recovery process of forests after a subsequent windthrow event.
105	Furthermore, hemiboreal forests are projected to experience multiple windthrows at
106	shorter return intervals than in the past due to the increase in windthrow frequency under
107	climate change (Usbeck et al. 2010; Donat et al. 2011; Gregow et al. 2017; Laapas et al.
108	2019). The effects of post-windthrow management on forests would diminish with time
109	(Taeroe et al. 2019). However, when windthrow occurs before the effects of previous post-
110	windthrow management do not completely faded, the effects of post-windthrow management
111	could accumulate; in the worst case, this could result in the conversion of forests to
112	completely different forest types. Thus, understanding the effects of multiple windthrows at
113	shorter return intervals and following management is essential in considering appropriate
114	post-windthrow management under climate change.
115	It is difficult to empirically evaluate the effects of forest management and the
116	changes in the frequency of windthrow which occurs only once every few decades on forest
117	ecosystems because it needs continuous monitoring extending over several hundred years.
118	Only Pontailler et al. (1997) reported the effects of multiple windthrows and their cumulative
119	effects on forest succession; however, the effects of management after multiple windthrows

120	have not been assessed. Forest landscape simulations are an effective tool for evaluating the
121	effects of various post-windthrow management and windthrow regimes on long-term forest
122	dynamics. However, no modeling study has yet revealed the effects of multiple windthrows
123	and subsequent management on species composition and aboveground biomass.
124	To evaluate the effects of post-windthrow management on species composition and
125	the recovery of forest biomass using forest landscape models, it is essential to refine the
126	representation of regeneration processes. The lack of detailed representation of regeneration
127	processes is one of the problems with existing forest landscape models (Albrich et al. 2020).
128	Among the various regeneration processes, tree-grass competition is represented in some
129	models (Thrippleton et al. 2016; Scheller et al. 2021), but regeneration on downed logs has
130	not been represented in any models. In forests in North America and Japan, many studies have
131	reported that seedlings of certain tree species distribute only on downed logs or stumps
132	(Harmon and Franklin 1989; Takahashi et al. 2000; Nakagawa et al. 2001). In particular,
133	more than 90% of seedlings of Picea spp. and Tsuga spp. distribute on well-decayed downed
134	logs or stumps (Weaver et al. 2009). Furthermore, downed logs and stumps function as places
135	where seedlings can escape shading by dwarf bamboo (Sasa spp., hereafter referred to as
136	Sasa); Sasa dominates the understory of forests in regions covered with snow in winter in east
137	Asia, and downed logs and stumps are therefore essential for forest regeneration in such
138	forests (Hiura et al. 1996). Thus, the incorporation of the regeneration process on downed

140	management using these models.
141	In this study, we focused on hemiboreal mixed forests in northern Japan and
142	approached the following three objectives using the process-based forest landscape model
143	LANDIS-II (Scheller et al. 2007):
144	(1) To incorporate the regeneration process on downed logs into LANDIS-II.
145	(2) To reveal the long-term effects of post-windthrow management on species composition
146	and the recovery of aboveground biomass.
147	(3) To reveal the long-term effects of the interaction between post-windthrow management
148	and windthrow frequency on species composition and aboveground biomass recovery.
149	LANDIS-II is a well-known powerful model that simulates forest dynamics and has been
150	tested and validated by many previous studies in various regions (e.g., Scheller et al. 2011;
151	Shifley et al. 2017; Lucash et al. 2019; Haga et al. 2020; Petter et al. 2020).
152	
153	2. Materials & Methods
154	2.1 Study area
155	The simulated landscape was a 12,169 ha forested area in the eastern half of the
156	University of Tokyo Hokkaido Forest (UTHF: 43°10'-21'N, 142°23'-41'E), located at 350-
157	1,000 m a.s.l. in central Hokkaido, northern Japan (Fig. 1). The annual mean temperature and

logs into forest landscape models is essential for evaluating the effects of post-windthrow

139

precipitation at the meteorological observatory at Rokugo (43°18′6′′N, 142°31′18″E, 315 m

159	a.s.l.; Japanese Meteorological Agency 2012, an average of 1981-2010) are 5.5 °C and 972.6
160	mm, respectively. The forested area is dominated by Abies sachalinensis (F. Schmidt) Mast.,
161	Picea jezoensis (Siebold et Zucc.) Carrière var. jesoensis, Betula ermanii Cham., and Tilia
162	japonica (Miq.) Simonk. The dominant soil types at the study site are Cambisols (parent
163	material: rhyolite or dacite) and Andosols (parent material: andesite, rhyolite, or dacite) (IUSS
164	Working Group WRB 2015). The natural disturbance regime of this area is stand-replacement
165	windthrow, and there are many records of damage by windthrow (e.g., Typhoon Marie in 1954
166	and Typhoon Thad in 1981; Watanabe et al. 1990). In particular, the damage by Typhoon Thad
167	in 1981 accounted for 8,735 ha (38.9% of the UTHF) in the area and 807,000 m ³ in timber
168	volume (Takada et al. 1986).

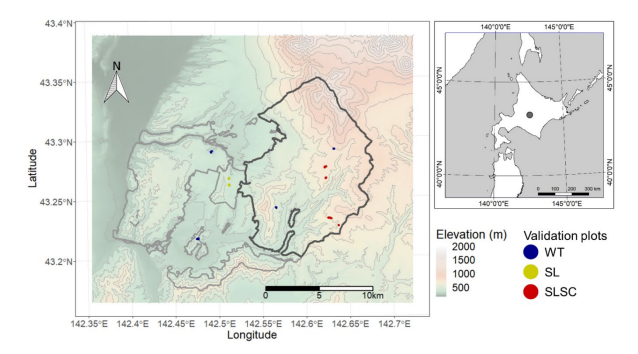


Fig. 1 Right: Map of northern Japan and its surroundings. The gray point indicates the location of the study area. Left: Detail of the study landscape. The University of Tokyo Hokkaido Forest is indicated by the light and dark gray lines, and the simulated landscape is indicated by the dark gray lines. The points indicate the locations of the plots used for validation; WT: stands where dead wood generated by the windthrow in 1981 was left intact; SL: stands where dead wood generated by the windthrow in 1981 was salvaged; SLSC: stands where dead wood generated by the windthrow in 1981 was salvaged, followed by scarification.

LANDIS-II, one of the forest landscape simulation models, can simulate forest 179dynamics and biogeochemical cycling at the landscape scale (Scheller et al. 2007). The 180 181 landscape is represented as a collection of interconnecting uniformly sized sites (cells) that contain information about vegetation, climate, soil nutrients, and so on. In this study, the size 182of a site was defined as 1 ha. Vegetation is represented in the model as species-age cohorts, 183184 and any number of cohorts can be contained in each site. The vegetation and environment at each site influence other surrounding sites through seed dispersal. The seed dispersal 185186 algorithm proceeds by the following steps: (1) the conditions at each site, such as light and nutrients, are checked to determine whether seeds of each species can germinate and 187188 establish; (2) for each site, neighboring sites are searched for seed sources; (3) a neighboring site can serve as a seed source if (a) at least one cohort of the same species at the neighboring 189 190 site is older than the age of maturity and (b) the distance of the neighboring site from the examined site is equal to or less than (maximum distance + cell size); (4) the probability of 191192 seed arrival is calculated based on the effective and maximum distances, which are userdefined parameters; and (5) neighboring sites, as defined by the maximum distance, are 193searched until the probability of seed arrival exceeds a random number. At this point, all three 194 criteria for reproduction (light, establishment, and seeds) are met, and seed dispersal ends. 195Alternatively, seed dispersal ends when all neighboring sites have been checked (Scheller et 196 al. 2007). LANDIS-II requires a single "succession extension" and can include optional 197

198	additional extensions such as disturbance or output extensions. These features enable the
199	effects of forest fire, windthrow, or harvest on forest ecosystems to be evaluated.
200	To simulate forest succession, we used the improved version of the Net Ecosystem
201	Carbon and Nitrogen Succession extension v 6.3 (Scheller et al. 2011; NECN succession) in
202	LANDIS-II. The improvements to NECN succession are explained in detail in section 2.3. In
203	NECN succession, the growth of biomass and the establishment of seedlings are calculated
204	based on the environmental conditions at each site. The growth of the biomass of each cohort
205	is represented as the difference between the monthly aboveground net primary production and
206	the monthly mortality. The monthly aboveground net primary production is calculated by
207	multiplying the maximum value of that (maxANPP; user-defined parameter) by coefficients
208	related to environmental limiting factors as follows: (1) limitTemperature: the coefficient
209	calculated with soil temperature; (2) limitN: the coefficient calculated with nitrogen
210	availability; (3) limitH ₂ O: the coefficient calculated with soil water content; (4) limitLAI: the
211	coefficient calculated with the Leaf Area Index (LAI) of the cohort itself; and (5)
212	limitCompetition: the coefficient calculated with the LAIs of the other cohorts within the site.
213	The establishment of cohorts of each tree species at each site is determined by two criteria: (1)
214	temperature and moisture conditions and (2) light availability. The light availability at each
215	site is defined as one of five shade classes (ShadeClass) determined by the total LAI at the
216	site. When these two requirements were satisfied, cohorts established at the site. The tree

- species parameters and their calibrations are explained in detail in supplementary materials S1
- 218 and S2.

2.3 Improvements to LANDIS-II 219

220	We added the regeneration process on downed logs to LANDIS-II NECN succession
221	v 6.3 with tree-grass competition implemented (Scheller et al. 2021) as follows.
222	(1) The identification of dead wood decay classes and the calculation of the amount of dead
223	wood in each decay class.
224	To calculate the amount of well-decayed dead wood at each site, NECN succession
225	extension was modified to compute the amount of dead wood in each decay class (five
226	levels). It is noted that "dead wood" in LANDIS-II includes dead wood of all sizes, such as
227	fine and coarse woody debris. The decay class is an indicator of the dead wood decay
228	advancement level, and the characteristics of dead wood in each decay class are as follows:
229	"decay class 1 - boles with no decay, fine twigs remaining, and complete bark coverage;
230	decay class 2 - slightly decayed boles with most of the bark present but no fine twigs; decay
231	class 3 - moderately decayed boles with some bark present but only stubs of branches
232	remaining; decay class 4 - boles that can no longer support themselves, with all bark gone;
233	and decay class 5 - boles detected only by their moss outlines on the forest floor" (Graham
234	and Cromack 1982). First, we modified the model to track dead wood and to calculate the
235	retention rate of dead wood (the ratio of the amount of dead wood in each year
236	(currentDeadWood) to that in the year in which each piece of dead wood was produced
237	(originalDeadWood)). Then, we determined the decay class of the dead wood by comparing

238	the calculated retention rate with empirical data on the retention rate of each decay class. The
239	empirical data on the retention rate of each decay class was calculated by Eq. 2.3.1.
240	$RR_{decayClassX} = WD_{decayClassX}/WD_{decayClass1}$ 2.3.1
241	* $RR_{decayClass X}$: the mass retention ratio of dead wood in decay class X relative to decay class 1;
242	$WD_{decayClass X}$: the wood density of dead wood in decay class X (the empirical data provided in
243	Ugawa <i>et al.</i> (2012)); and <i>X</i> : decay class (1, 2, 3, 4, 5).
244	Finally, NECN succession extension was modified to compute the amount of dead
245	wood in each decay class in each year (see also, supplementary materials S3).
246	
247	(2) The calculation of the area occupied by downed logs available as regeneration sites
248	The downed logs available as regeneration sites were those in decay classes 3, 4, and
249	5. In these downed logs, decay had progressed throughout the wood, the bark had been
250	removed, and moss was attached to the surface (Takahashi et al. 2000). Thus, the area
251	occupied by downed logs in decay classes 3, 4, and 5 at each site was made computable by
252	the model. The shape of downed logs was assumed to be an elliptical column, and the area
253	occupied by downed logs in decay classes 3, 4, and 5 was calculated by Eq. 2.3.2 using the
254	calculated amount of downed logs in decay classes 3, 4, and 5. Finally, the ratio of the total
255	area occupied by downed logs in decay classes 3, 4, and 5 to the site area
256	(nurseryLogAreaRatio) was calculated.
257	nurseryLogArea _{decayClassX} = 4 * 2 * $C_{decayClassX}/(\pi * h * WD_{decayClassX})$ 2.3.2

258	nurseryLogArea _{decayClassX} : the area occupied by downed logs in decay class X; $C_{decayClassX}$: the
259	carbon stock of dead wood in decay class X ; h: the average height of downed logs (0.28 m:
260	data from Hotta <i>et al.</i> (2020)); $WD_{decayClassX}$: the wood density of dead wood in decay class X
261	(data from Ugawa et al. (2012)); X: the decay class (3, 4, or 5).
262	The amount of dead wood is not strictly equal to the number of downed logs because
263	dead wood includes snags, which are standing dead trees. However, almost all snags are
264	classified as decay class 1 or 2; as decay progresses, the snags break or fall and turn into
265	downed logs. Thus, we assumed that the amount of dead wood in decay classes 3, 4, or 5 was
266	equal to that of downed logs in decay classes 3, 4, or 5.
267	
268	(3) The improvements to the determination of cohort establishment
268 269	(3) The improvements to the determination of cohort establishment In NECN succession, the probability of the establishment of cohorts based on light
269	In NECN succession, the probability of the establishment of cohorts based on light
269 270	In NECN succession, the probability of the establishment of cohorts based on light availability is determined by comparing random numbers with "LightProbability", which is
269 270 271	In NECN succession, the probability of the establishment of cohorts based on light availability is determined by comparing random numbers with "LightProbability", which is determined by the shade tolerance of each species and the shade class of each site. In this
269 270 271 272	In NECN succession, the probability of the establishment of cohorts based on light availability is determined by comparing random numbers with "LightProbability", which is determined by the shade tolerance of each species and the shade class of each site. In this study, the tree species were categorized into dead wood-dependent and dead wood-
269 270 271 272 273	In NECN succession, the probability of the establishment of cohorts based on light availability is determined by comparing random numbers with "LightProbability", which is determined by the shade tolerance of each species and the shade class of each site. In this study, the tree species were categorized into dead wood-dependent and dead wood- independent species; dead wood-dependent species required well-decayed dead wood for
 269 270 271 272 273 274 	In NECN succession, the probability of the establishment of cohorts based on light availability is determined by comparing random numbers with "LightProbability", which is determined by the shade tolerance of each species and the shade class of each site. In this study, the tree species were categorized into dead wood-dependent and dead wood- independent species; dead wood-dependent species required well-decayed dead wood for regeneration, while dead wood-independent species did not require dead wood for

(A) Dead wood-dependent species 278

279	P. jezoensis and Picea glehnii (F. Schmidt) Mast. were defined as dead wood-
280	dependent species (Takahashi et al. 2000). dead wood-dependent species require dead wood
281	for regeneration; thus, we modified the calculation of the probability of establishment based
282	on LightProbability as shown in Eq. 2.3.3.
283	$LightProbability_{Modified} = LightProbability_{Original} * nurseryLogAvailability 2.3.3$
284	*LightProbability _{Modified} : LightProbability calculated by the modified way;
285	LightProbabilityOriginal: LightProbability calculated by the original way;
286	nurseryLogAvailability: the adjustment coefficients calculated by nurseryLogAreaRatio
287	$(nurseryLogAvailability = 1 - (nurseryLogAreaRatio - 1)^{nurseryLogAvailabilityModifier};$
288	nurseryLogAvailabilityModifier: a tuning parameter that is an even number. In this study, 4
289	was the most suitable nurseryLogAvailabilityModifier according to our calibrations
290	(supplementary materials S4).
291	(B) Dead wood-independent species
292	Dead wood-independent species were defined as species that can establish without
293	dead wood. All tree species except for P. jezoensis and P. glehnii were defined as dead wood-
294	independent species because previous studies suggested those species can establish on forest
295	floor (Takahashi et al. 2000; Weaver et al. 2008). In determining the establishment of dead
296	wood-independent species, in addition to the original determination process, we made it

- possible to determine the possibility of establishment on downed logs in case the site was toodark for the species to establish, as follows (Fig. 2).
- 299 (a) In the original determination method, the cohort can establish when LightProbability_{Original}
- 300 exceeds a random number generated for each site. If a cohort can establish in stage (a), the
- 301 location of its establishment is the forest floor.
- 302 (b) If cohorts cannot establish in stage (a) and the shade class of the site is darker than the
- 303 most suitable shade class for the species, the model checks whether the cohorts can establish
- if they are not shaded by grasses, such as *Sasa* dwarf bamboo. Specifically, the model checks
- 305 whether the LightProbability calculated according to the tree species LAI exceeds the random
- 306 number at each site. In stage (b), the model checks only the possibility of establishment and
- 307 does not determine the establishment of the cohorts.
- 308 (c) When the LightProbability calculated according to the tree species LAI exceeds the
- 309 random number in stage (b), cohorts can establish if there are sufficient amounts of downed
- 310 logs in decay classes 3, 4, and 5. Specifically, cohorts can establish on downed logs when
- 311 nurseryLogAvailability exceeds a random number generated for each site.
- 312

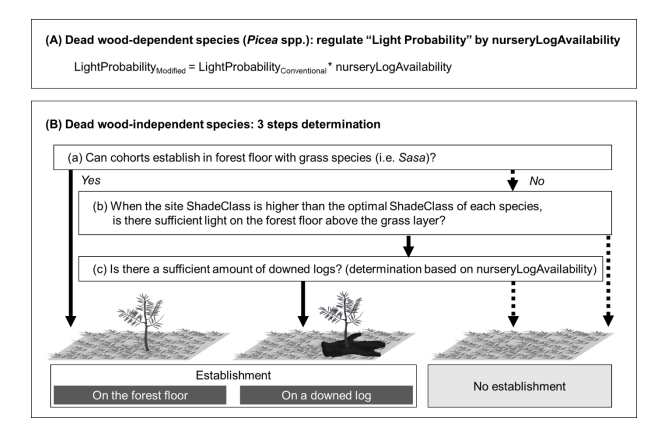


Fig. 2 Flowchart showing the determination of the establishment of seedlings in LANDIS-II NECN succession in which the process of regeneration on downed logs was implemented. A: in the case of dead wood-dependent species; B: in the case of dead wood-independent species. LightProbability: the probability of establishment determined by the shade tolerance of each species and the shade class of each site; nurseryLogAvailability: the amount of dead wood available as regeneration sites; and ShadeClass: the degree of darkness of the site, as classified into five levels.

2.4 *Input data*

2.4.1 *Initial communities*

323	We focused on the tree species that accounted for 90% of the total biomass and on the
324	dominant grass species: A. sachalinensis, P. jezoensis, P. glehnii, T. japonica, Acer pictum
325	Thunb., B. ermanii, Betula maximowicziana Regel, Quercus crispula Blume var. crispula,
326	Kalopanax septemlobus (Thunb.) Koidz., Fraxinus mandshurica Rupr., Ulmus laciniata
327	(Trautv.) Mayr ex Schwapp., and Sasa senanensis (Franch. et Sav.) Rehder.
328	The initial communities of trees were created based on forest inventory data from the
329	UTHF. The data were acquired in three different periods in the study landscape: period 11:
330	1996-2005; period 12: 2006-2010; and period 13: 2011-2020. We summarized the inventory
331	data by period, and we treated the data collected in periods 11, 12, and 13 as data collected in
332	2000, 2008, and 2015, respectively. Additionally, the data collected in periods 11 and 12 were
333	simulated for 15 years and 7 years, respectively, to align all data to the same year (2015).
334	Then, we constructed the initial communities in the study landscape in 2015.
335	The initial aboveground biomass of S. senanensis at each site was determined
336	through an estimation from tree biomass at each site using the Sasa distribution model
337	developed by Tatsumi and Owari (2013).
338	

2.4.2 *Dead wood*

The amount of dead wood was uniformly input within the landscape to eliminate

341	uncertainties derived from the heterogeneity of the distribution of dead wood. We used the
342	average value of the amount of dead wood in Hotta et al. (2020), 3,100 g m ⁻² , as the input
343	amount. We assumed the decay class of dead wood to be decay class 1 because the amount of
344	dead wood in each decay class was unknown.
345	
346	2.4.3 Soil
347	The input data related to soils were based on Asahi (1963), which reported the soil
348	properties of the UTHF in detail. The following data were used as soil input data: drainage,
349	field capacity, wilting point, sand rate, clay rate, carbon content, and nitrogen content.
350	
351	2.4.4 Climate
352	The input data related to climate were configured as follows. The 1 km mesh climate
353	data (Japanese Meteorological Agency 2012, on average over 1981-2010) in the study
354	landscape were classified into 10 clusters with k-means clustering based on the monthly
355	maximum temperature, monthly minimum temperature, and monthly precipitation. Each
356	climate data was standardized before clustering. The climate data were spatially averaged and
357	input according to climate cluster.
358	

2.5 Settings for windthrow and post-windthrow management scenarios

360	The Biomass Harvest extension v. 4.3 (Gustafson et al. 2000) was used to represent
361	stand-replacing windthrow in the study landscape. We defined stand-replacing windthrow as
362	follows: windthrow area: 20% of the study landscape; windthrow intensity: all living trees
363	blown down except for advanced seedlings; windthrow frequency: interval of 50 years (with
364	windthrow occurring in years 15 and 65). The ratio of windthrow area for each windthrow
365	event was determined based on the records of windthrow by Typhoon Thad in 1981
366	(Watanabe et al. 1990). The interval of windthrow was determined based on Abe et al. (2006).
367	The sites where windthrow occurred were randomly selected from among sites where the
368	stand age was more than 50 years. Windthrow risk is influenced not only by stand age but also
369	by the wind direction, wind intensity, forest structures, topography, and so on. Previous
370	studies have reported that the risk of windthrow is greater in individual trees with larger
371	diameters at breast height or tree heights (Rich et al. 2007) and in stands that have
372	homogeneous structures (Jalkanen and Mattila, 2000; Mitchell et al. 2001; Morimoto et al.
373	2019b). On the other hand, predicting windthrow risk is very challenging because it varies
374	greatly among regions and among windthrow events (Dobor et al. 2019). Because a detailed
375	evaluation of windthrow risk is beyond the scope of the objectives of this study, the
376	windthrow risk was determined only by the stand age in this study.
377	We evaluated the following three post-windthrow management scenarios: (a) dead
378	trees left undisturbed; (b) salvage logging; and (c) salvage logging and scarification. The
379	proportion of advanced seedlings that are destroyed differs among scenarios. Whether the

380	cohorts contained advanced seedlings was determined by the cohort age, and the threshold
381	age was determined by the tree species based on data from the windthrow sites in the UTHF
382	caused by Typhoon Thad in 1981 (supplementary materials S5).
383	(a) Dead trees left undisturbed (WT)
384	In the WT scenario, the dead wood generated by the windthrow was left intact, and
385	20% of advanced seedlings were destroyed (Table 1). The advanced seedling destruction ratio
386	was determined based on data from the windthrow sites in the UTHF caused by Typhoon
387	Thad in 1981 (Kurahashi et al. 1983).
388	(b) Salvage logging (SL)
389	In the SL scenario, dead wood generated by windthrow was salvaged, and 60% of
389 390	In the SL scenario, dead wood generated by windthrow was salvaged, and 60% of advanced seedlings were destroyed (Table 1). This percentage was the most suitable for
390	advanced seedlings were destroyed (Table 1). This percentage was the most suitable for
390 391	advanced seedlings were destroyed (Table 1). This percentage was the most suitable for empirical data in salvaged sites, and Ohsato <i>et al.</i> (1996) also reported that the percentage of
390 391 392	advanced seedlings were destroyed (Table 1). This percentage was the most suitable for empirical data in salvaged sites, and Ohsato <i>et al.</i> (1996) also reported that the percentage of forest area disturbed by salvage logging was approximately 60%.
390 391 392 393	advanced seedlings were destroyed (Table 1). This percentage was the most suitable for empirical data in salvaged sites, and Ohsato <i>et al.</i> (1996) also reported that the percentage of forest area disturbed by salvage logging was approximately 60%. (c) Salvage logging and scarification (SLSC)
390 391 392 393 394	advanced seedlings were destroyed (Table 1). This percentage was the most suitable for empirical data in salvaged sites, and Ohsato <i>et al.</i> (1996) also reported that the percentage of forest area disturbed by salvage logging was approximately 60%. (c) Salvage logging and scarification (SLSC) In the SLSC scenario, dead wood generated by windthrow was salvaged, and the

Post-windthrow management scenario	Dead wood generated by the windthrow	Advanced seedlings	Dwarf bamboo (Sasa)
Windthrow (WT)	Left intact	20% destroyed	Undestroyed
Salvage logging (SL)	100% salvaged	60% destroyed	Undestroyed
Salvage logging and scarification (SLSC)	100% salvaged	100% destroyed	99% destroyed

400 2.6 Calibration

399

To adjust the performance of the model to better fit the empirical data, some parameters 401 were calibrated by comparing the simulation results and the empirical data. Previous studies 402also calibrated parameters used to calculate the biomass growth, litterfall, decomposition rate 403of soil organic carbon, etc. (Lucash et al. 2019; Haga et al. 2019; Haga et al. 2020). In addition 404 to these parameters, parameters related to the calculation of the LAI, net primary production 405 (NPP), tree-grass competition, and decomposition rate of dead wood were also calibrated in 406 this study. The SPOTPY package (Houska et al. 2015) in Python ver. 3.10 was used to perform 407408 the calibrations.

411	The mortality shape, maxANPP, maxBiomass, monthly wood mortality, and KLAI
412	were used as tuning parameters, and the aboveground biomass growth of trees and the litterfall,
413	LAI, and NPP at the sites were calibrated for each species. For the aboveground biomass growth,
414	the volume table of living trees according to tree species and area (Hokkaido 2010) was used
415	to provide empirical data. In addition, the aboveground biomass growth curves were extended
416	to age 140 by nonlinear regression using the least squares method because data for some species
417	were insufficient in terms of stand age. Regarding litterfall, data from deciduous broadleaved
418	forests (Tomakomai, Uryu, Ashoro in Hokkaido, northern Japan) and evergreen coniferous
419	forests (Otanomousdaira in Nagano, central Japan) collected in the monitoring site 1000 project
420	(Ishihara et al. 2011) were used as empirical data. Data from Muraoka et al. (2010) and Sumida
421	et al. (2018) were used as empirical data for the LAIs of deciduous broadleaved forests and
422	evergreen coniferous forests, respectively. Data from Ishii (2019) were used to provide
423	empirical data on the NPP in cool-temperate forests.
424	We selected the best parameter set based on the following conditions: (1) the simulated

We selected the best parameter set based on the following conditions: (1) the simulated LAI and NPP at the sites were within the range of the empirical values and (2) the root mean squared errors (RMSEs) of the aboveground biomass and litterfall between ages 130 and 140 were minimal (Fig. S2.1, S2.2).

428

429 **2.6.2** Aboveground biomass and the LAI of Sasa and tree–grass competition

430	We calibrated the parameters related to Sasa for the following three processes: (1)
431	simulating the aboveground biomass of Sasa between 27 and 32 years after scarification to
432	match the empirical data; (2) simulating the LAI of Sasa between 27 and 32 years after
433	scarification to match the empirical data; and (3) simulating tree species composition 32 years
434	after scarification to match the empirical data. The RMSE was used to evaluate the objective
435	functions of (1) and (2), and the Bray-Curtis index of dissimilarity was used to evaluate the
436	objective function of (3). Finally, the parameter set in which the sum of the three standardized
437	objective functions was minimized was selected. The MaxANPP, maxBiomass, monthly wood
438	mortality, KLAI, BTOLAI, and grass threshold multiplier were used as the tuning parameters.
490	

- 439
- 440 **2.6.3** *Dead wood decomposition rate*

The dead wood decomposition rate was calibrated by tuning the time until dead wood 441 442reached decay class 3 using the "Wood decay rate" parameter. The actual time until dead wood reached decay class 3 was estimated by the following processes. First, the decomposition rate 443constant (k) in the climate of the simulated landscape was estimated using the equation in Russel 444 et al. (2015). Second, the retention ratio of dead wood (R) was calculated by the following 445equation: R = -exp(kt) (t: the elapsed time since the trees died). The retention ratio of the mass 446 of dead wood in decay class 3 was calculated by using the wood density of dead wood in each 447decay class provided in Ugawa et al. (2012). Finally, the actual time until dead wood reached 448 decay class 3 was estimated from k and R. 449

451 **2.6.4** Soil organic carbon

452	We selected the parameter set in which the change in soil organic carbon in the first
453	five years was the lowest because no data on changes in soil organic carbon over time were
454	available. In LANDIS-II, there are four pools of soil organic carbon. We carefully avoided the
455	decomposition rates of SOM2 and SOM3 exceeding those of the surface and SOM1, which are
456	more active pools (Lucash et al. 2019).

457

458 **2.7** Validation

The results of the simulation of post-windthrow management scenarios were validated 459460 by comparison with empirical data. We used data of permanent plots for measuring forest recovery after the windthrow caused by Typhoon Thad in 1981 and following various post-461462windthrow managements carried out in the UTHF. The data were collected in 1982, just after the windthrow, and in 2014, and included species and diameter at breast height (DBH) and 463 464 height measurements. We determined the age of each individual according to the DBH, calculated aboveground biomass by the DBH and height, aggregated the aboveground biomass 465of each individual into cohorts of 10-year age intervals, and constructed the initial communities 466 based on the empirical data from 1982, simulated forest succession for 32 years in a single grid, 467 and compared the simulated results with the empirical data from 2014. Additionally, to 468 quantitatively evaluate the accuracy of validation, we used relative errors and the Brey-Curtis 469

470 index of dissimilarity for the aboveground biomass and species composition, respectively.471

2.8 *Simulation*

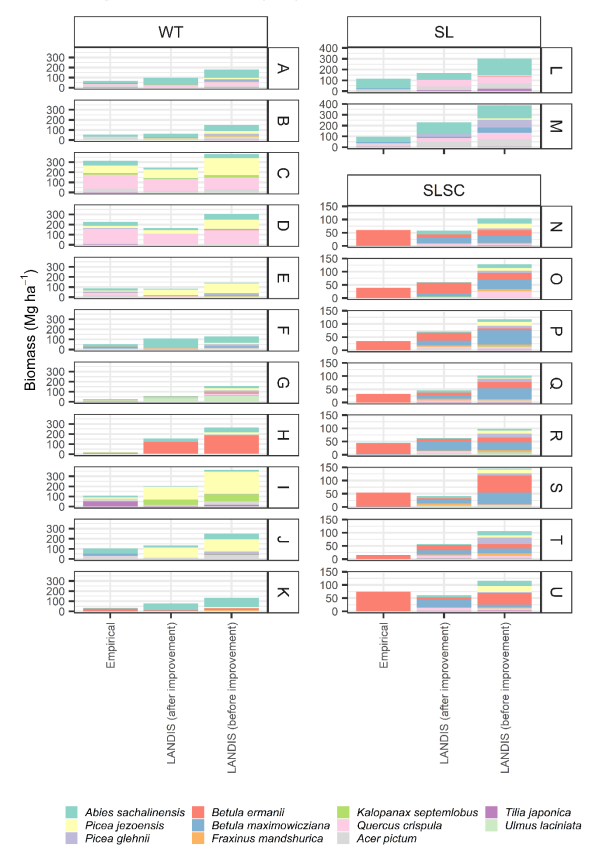
473	The simulation was replicated 10 times for each scenario to capture the stochasticity
474	related to seed dispersal, cohort establishment, and the selection of windthrow sites. The
475	duration of the simulation was 115 years. The results were classified into the following two
476	categories: stands where windthrow occurred only in year 15 (1,947 sites) and stands where
477	windthrow occurred in both year 15 and year 65 (486 sites). The 10 replicates of the mean
478	aboveground biomass for each category and post-windthrow management scenario were
479	ensemble-averaged. It is noted that the aboveground biomass of windthrow sites was
480	averaged in each category.

3. Results

3.1 Validation of and improvements in LANDIS-II

485	The relative errors of the total aboveground biomass from empirical data versus
486	LANDIS (after improvement) were in the range from -0.27 to 5.7. The Brey-Curtis index of
487	dissimilarity in species composition between empirical data and LANDIS (after
488	improvement) were in the range from 0.17 to 0.87. Although there were slight differences in
489	species composition and aboveground biomass in some plots, the tree species composition
490	and aboveground biomass in each plot coincided more with the empirical data after the
491	incorporation of the regeneration process on downed logs than before, regardless of which
492	post-windthrow management practice was implemented (Fig. 3, 4). Before the incorporation
493	of this regeneration process, dead wood-dependent species such as <i>P. jezoensis</i> and <i>P. glehnii</i>
494	were simulated to be able to regenerate under SL and SLSC, which are scenarios that leave
495	behind little well-decayed dead wood. However, their simulated regeneration decreased after
496	the incorporation of regeneration on downed logs into the model; accordingly, the simulated
497	aboveground biomass was also closer to the empirical values. Additionally, the relative error
498	and Brey-Curtis index were substantially closer to 0 and relatively lower than those of
499	empirical data versus LANDIS (before improvement) (0.22 to 10.3 and 0.20 to 0.92,
500	respectively). Furthermore, the RMSE of aboveground biomass growth for each species in the
501	calibration ranged from 714 to 3,158 g m ⁻² (Fig. S2.1).

502	Although the differences in tree species composition and aboveground biomass
503	between the simulated and empirical data were small after the improvement of the model, the
504	simulated aboveground biomass was still slightly overestimated, and the simulated species
505	composition was slightly different from that in the empirical data in some WT and SL plots
506	(Fig. 3, 4). Under SLSC, the difference in aboveground biomass between the simulated and
507	empirical data was minimal, and Betula spp. was dominant in both the simulated and
508	empirical data (Fig. 3, 4). In particular, the differences in species composition were relatively
509	large in plots E, G, H, I, and J under WT and in plot L under SL, and the differences in
510	aboveground biomass were relatively large in plots F, G, H, I, and K under WT and in plot M
511	under SL.



Trees aboveground biomass recovery 32 years after windthrow.

513 Fig. 3 The results of validations of aboveground biomass and species composition 32 years

514	after windthrow. Empirical: empirical aboveground biomass data for each species 32 years
515	after windthrow and post windthrow management; LANDIS (after improvement): the results
516	for the aboveground biomass of each species 32 years after windthrow and post windthrow
517	management simulated by LANDIS-II NECN succession in which the process of regeneration
518	on downed logs was implemented; LANDIS (before improvement): the results for the
519	aboveground biomass of each species 32 years after windthrow and post windthrow
520	management simulated by the original version of LANDIS-II with NECN succession. WT: the
521	scenario in which dead wood generated by windthrow is left intact; SL: the scenario in which
522	dead wood generated by windthrow is salvaged; SLSC: the scenario in which dead wood
523	generated by windthrow is salvaged, followed by scarification. Alphabets on the right side of
524	each figure indicate the IDs of the permanent plots.

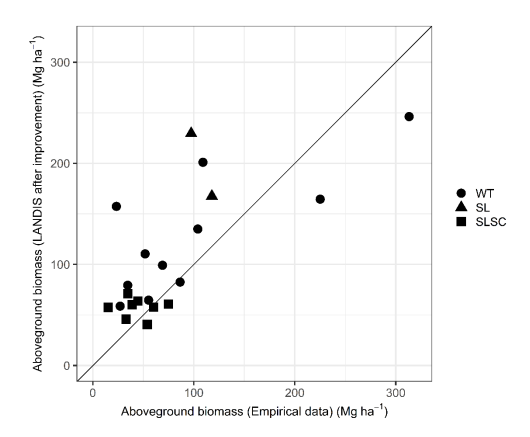
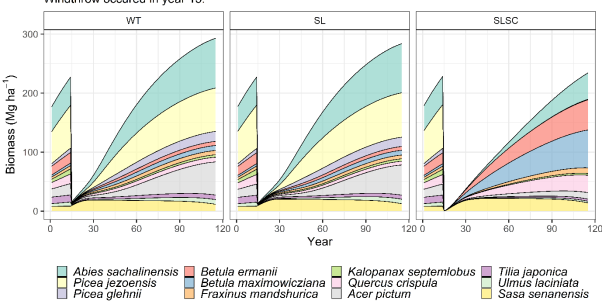


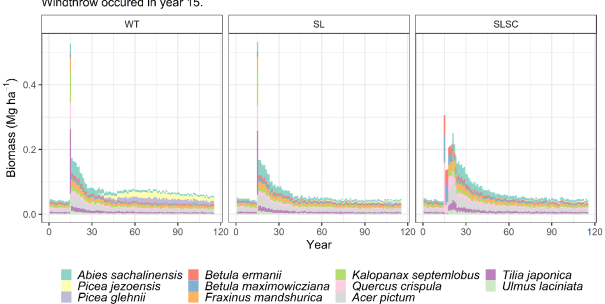
Fig. 4 Scatter plot of the tree aboveground biomass 32 years after the windthrow and
following management from empirical data and LANDIS-II simulation results after
improvement. WT: the scenario in which dead wood generated by windthrow was left intact;
SL: the scenario in which dead wood generated by windthrow was salvaged; and SLSC: the
scenario in which dead wood generated by windthrow was salvaged, followed by
scarification.

3.2 *Recovery after a single windthrow and subsequent management*

534	In stands where windthrow occurred only in year 15, the recovery of aboveground
535	biomass was slower under SLSC than under WT and SL (Fig.5). In the long term, the
536	dominant species under SLSC were B. ermanii and B. maximowicziana, which are quite
537	different from the dominant species under WT and SL (Fig.5). The species composition and
538	aboveground biomass under SL were little different from those under WT (Fig.5). Under
539	SLSC, the establishment of early-successional species such as <i>B. ermanii</i> , <i>B.</i>
540	maximowicziana, and Q. crispula increased soon after windthrow (Fig.6). Under WT and SL,
541	the establishment of mid-successional species such as T. japonica and A. pictum increased
542	soon after windthrow (Fig.6). In addition, the establishment of <i>P. jezoensis</i> and <i>P. glehnii</i>
543	increased more than 20 years after windthrow under WT; however, such an increase was not
544	detected under SL or SLSC (Fig.6).
545	



Changes in species composition at windthrow area Windthrow occured in year 15.

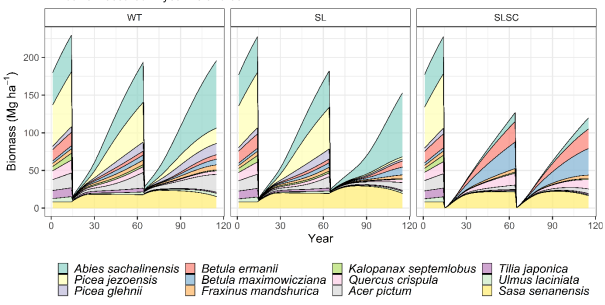


Cohorts biomass established in each year at windthrow area Windthrow occured in year 15.

Fig.6 The species composition of cohorts established in each year in stands where windthrow 553occurred only in year 15. WT: the scenario in which dead wood generated by windthrow is 554left intact; SL: the scenario in which dead wood generated by windthrow is salvaged; SLSC: 555the scenario in which dead wood generated by windthrow is salvaged, followed by 556scarification.

3.3 *Recovery after the two windthrows and subsequent management*

560	In the stands where windthrow occurred in years 15 and 65, the recovery of
561	aboveground biomass after the second windthrow (year 65) was slower under SL than under
562	WT, and dead wood-dependent species hardly regenerated after the second windthrow under
563	SL (Fig.7). On the other hand, there were no differences in species composition or aboveground
564	biomass recovery after the first and second windthrow under SLSC (Fig.7). Regardless of the
565	post-windthrow management scenario, there was also no difference between the species
566	composition of the cohorts established after the first windthrow and that of the cohorts
567	established after the second windthrow (Fig.8).



Changes in species composition at windthrow area Windthrow occured in year 15 and 65.

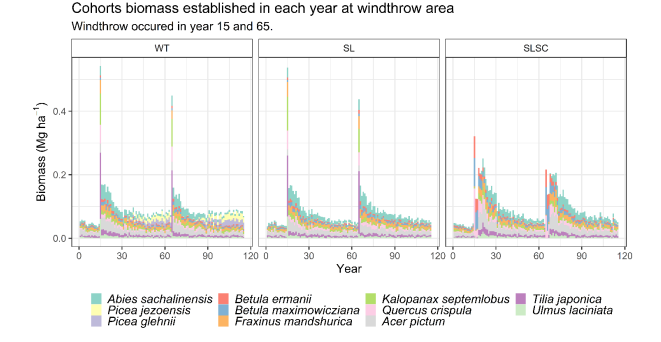


Fig.8 The species composition of cohorts established in each year in stands where windthrows
occurred in years 15 and 65. WT: the scenario in which dead wood generated by windthrow is
left intact; SL: the scenario in which dead wood generated by windthrow is salvaged; SLSC:
the scenario in which dead wood generated by windthrow is salvaged, followed by
scarification.

583 **4. Discussion**

584 4.1 Improvements in LANDIS-II

585	The trends in forest recovery associated with post-windthrow management were
586	represented more precisely by LANDIS-II that incorporated the regeneration process on
587	downed logs in addition to tree-grass competition (Scheller et al. 2021) than by the original
588	LANDIS-II model (Fig. 3). Before the improvements, the simulations showed dead wood-
589	dependent species such as <i>P. jezoensis</i> and <i>P. glehnii</i> regenerating even under SL and SLSC,
590	which are scenarios with less well-decayed dead wood. We successfully represented the
591	establishment process of dead wood-dependent species by limiting the establishment
592	probabilities of dead wood-dependent species on the basis of the number of downed logs in
593	decay classes 3, 4, and 5. This change resulted in a more precise representation of the species
594	composition and aboveground biomass 32 years after windthrow and the subsequent
595	management. Although multiple empirical studies have demonstrated the importance of dead
596	wood for tree regeneration (Harmon and Franklin 1989; Takahashi et al. 2000; Nakagawa et
597	al. 2001; Weaver et al. 2009), this process has not yet been implemented in forest landscape
598	models. Forest landscape models have often been used to evaluate the effects of harvesting or
599	various natural disturbances, and these disturbances likely also affect dead wood dynamics in
600	forest ecosystems. Therefore, forest landscape models that incorporate the regeneration
601	process on downed logs will contribute to improving the accuracy of predictions of forest
602	dynamics in future research.

603	In this study, we tried our best to validate forest development after various post-
604	windthrow management scenarios; however, its duration (32 years) was somewhat short
605	relative to the duration of main simulation analysis (100 years), and the empirical data used
606	for validation were data related to a single windthrow event due to lack of additional
607	empirical data. Establishing permanent plots and continuous monitoring surveys in forests
608	managed in a variety of ways would be critical for more precise validation. Evaluating biases
609	among results from multiple simulation models would also be useful.
610	
611	4.2 Long-term effects of post-windthrow management on species composition and
612	aboveground biomass
613	The decrease in dead wood and the destruction of advanced seedlings due to salvage
613 614	The decrease in dead wood and the destruction of advanced seedlings due to salvage logging had little effect on the species composition or aboveground biomass recovery in the
614	logging had little effect on the species composition or aboveground biomass recovery in the
614 615	logging had little effect on the species composition or aboveground biomass recovery in the stands. However, the complete destruction of advanced seedlings due to scarification resulted
614 615 616	logging had little effect on the species composition or aboveground biomass recovery in the stands. However, the complete destruction of advanced seedlings due to scarification resulted in a delay in the recovery of aboveground biomass and a conversion to birch-dominated forests,
614615616617	logging had little effect on the species composition or aboveground biomass recovery in the stands. However, the complete destruction of advanced seedlings due to scarification resulted in a delay in the recovery of aboveground biomass and a conversion to birch-dominated forests, and these effects lasted for 100 years. In stands where windthrow occurred once, there was no
 614 615 616 617 618 	logging had little effect on the species composition or aboveground biomass recovery in the stands. However, the complete destruction of advanced seedlings due to scarification resulted in a delay in the recovery of aboveground biomass and a conversion to birch-dominated forests, and these effects lasted for 100 years. In stands where windthrow occurred once, there was no difference in species composition or the recovery of aboveground biomass between WT and SL
 614 615 616 617 618 619 	logging had little effect on the species composition or aboveground biomass recovery in the stands. However, the complete destruction of advanced seedlings due to scarification resulted in a delay in the recovery of aboveground biomass and a conversion to birch-dominated forests, and these effects lasted for 100 years. In stands where windthrow occurred once, there was no difference in species composition or the recovery of aboveground biomass between WT and SL (Fig.5). Many of the surviving advanced seedlings had already exceeded the height of the <i>Sasa</i>

623	the advanced seedlings had little effect on long-term forest recovery after windthrow and
624	salvage logging. In contrast to these results, many empirical studies have reported delays in
625	forest recovery and increases in early-successional species due to intensive salvage logging
626	(Donato et al. 2006; Ilisson et al. 2007; Morimoto et al., 2011; Fischer and Fischer 2012). More
627	intensive salvage logging than that we assumed in this study, i.e., the destruction of more than
628	half of the advanced seedlings, could delay forest recovery more than our simulation predicted
629	and result in a forest more similar to that predicted under SLSC. Although the effects of salvage
630	logging on the species composition and aboveground biomass recovery were limited, the
631	establishment of <i>P. jezoensis</i> and <i>P. glehnii</i> was less common under SL than under WT (Fig.6).
632	These two species can establish only on well-decayed downed logs or stumps (Takahashi et al.
633	2000; Nakagawa et al. 2001), and the decrease in dead wood due to salvage logging resulted in
634	a decrease in the establishment of these species.
635	Short-term empirical studies have reported that Betula spp., which are major pioneer
636	species in boreal and hemiboreal zones, dominate forests that have undergone salvage logging
637	followed by scarification (Yoshida et al. 2005; Prévost et al. 2010; Aoyama et al. 2011; Suzuki
638	2020). Our results showed that the effects of scarification remained even 100 years after

scarification was performed (Fig.5). In our simulation, seedlings of *B. ermanii* and *B. maximowicziana* became established several years after *Sasa* was destroyed by scarification;
these seedlings immediately grew up and dominated the canopy after scarification. However,
the growth of a dense stand of *Betula* was diminished, probably due to competition with rapidly

643	recovering Sasa. As a result, the recovery of aboveground biomass under SLSC was slower
644	than that under WT and SL, and birch-dominated forests developed. Fischer et al. (2002)
645	applied an individual-based stand dynamics model and predicted that birches would dominate
646	immediately after windthrow; however, their populations gradually decreased at more than 30
647	years after windthrow. This may be due to the relatively short longevity of the Betula spp.
648	focused on in Fischer et al. (2002), Betula pendula Roth and Betula pubescens Ehrh., whose
649	longevity is 40-100 years (Jónsson 2004). On the other hand, <i>B. ermanii</i> and <i>B. maximowicziana</i> ,
650	which were focused on in this study, are known as long-lived pioneers and can live for more
651	than 300 years (Watanabe 1994). Therefore, our predictions showed the birch-dominated forest
652	remaining for 100 years after scarification. In the UTHF, B. maximowicziana has also been
653	dominant for over 100 years at sites that burned in the 1910s (Suzuki 2020). This result suggests
654	that the progress of forest succession can slow after long-lived pioneer species become
655	dominant (Chazdon 2008; Donate et al. 2012). Seed dispersal is also one of the major factors
656	influencing forest dynamics. Although variations in seed dispersal related to the wind direction,
657	topography and rich versus poor seed production are not represented, fundamental processes of
658	seed dispersal such as the effective and maximum seed dispersal distances and mature age are
659	represented in LANDIS-II. Therefore, there could be few effects to our discussion.

4.3 Long-term effects of the interaction between post-windthrow management and frequent
windthrows on species composition and aboveground biomass

663	The decrease in dead wood due to salvage logging caused a delay in the recovery of
664	aboveground biomass and a change in species composition after windthrow occurred again.
665	The decrease in newly established cohorts after the first windthrow, especially cohorts of P.
666	jezoensis and P. glehnii, due to salvage logging delayed forest recovery after the second
667	windthrow occurred 50 years after the first windthrow; these conditions resulted in a decrease
668	in aboveground biomass, especially that of <i>P. jezoensis</i> and <i>P. glehnii</i> . Birch-dominated forests
669	developed after the second scarification just as they did after the first scarification; this occurred
670	because stand development restarted from the "bare land" produced by scarification, which
671	thoroughly destroyed advanced seedlings and removed Sasa. The repeated windthrows and
672	salvaging (SL) at 50-year intervals slowed the recovery of aboveground biomass and greatly
673	reduced the biomass of <i>P. jezoensis</i> and <i>P. glehnii</i> after the second windthrow compared with
674	those in WT stands (Fig.7). This could be attributed to the lower number of advanced seedlings,
675	especially of <i>P. jezoensis</i> and <i>P. glehnii</i> , under SL than under WT when the second windthrow
676	occurred. Under WT, the species composition and aboveground biomass recovery after the
677	second windthrow were the same as those after the first windthrow because there was plenty of
678	well-decayed dead wood after the first windthrow, and dead wood-dependent species were able
679	to become established (Fig.8). However, fewer seedlings established between the first and
680	second windthrows (from years 16 to 64) under SL than under WT (Fig. 8); moreover, the
681	advanced seedlings were destroyed due to salvage logging after the second windthrow. Thus,
682	there were much fewer surviving advanced seedlings after the second windthrow under SL than

683	under WT. This difference explains why the recovery of aboveground biomass after the second
684	windthrow was slower under SL than under WT. Furthermore, the seedling community without
685	dead wood-dependent species (due to salvage logging after the first windthrow) grew into
686	mature trees after the second windthrow; as a result, dead wood-dependent species hardly
687	regenerated under SL after the second windthrow. Although previous studies have suggested
688	that salvage logging may affect the long-term regeneration dynamics of dead wood-dependent
689	species (Suzuki et al. 2019; Morimoto et al. 2019a; Hotta et al. 2020), these effects have not
690	yet been directly observed due to the difficulty of performing long-term monitoring in forest
691	ecosystems. Previous simulation studies have not evaluated these effects of salvage logging
692	because the regeneration process on downed logs had not yet been incorporated into forest
693	landscape models (Dobor et al. 2019). However, we successfully quantitatively evaluated the
694	effects of salvage logging on the regeneration dynamics of dead wood-dependent species by
695	using a process-based simulation model that includes the regeneration process on downed logs.
696	

697 5. Implications for forest management

When windthrow occurred only once, salvage logging hardly affected the species 698 composition and aboveground biomass of the forest within 100 years after the windthrow. 699 However, the number of advanced seedlings decreased, and the species composition of the 700 seedlings changed due to the decrease in dead wood caused by salvage logging. Therefore, 701 when windthrow occurred again within a short interval, salvage logging decreased the 702aboveground biomass of the forest and changed its species composition after the second 703 windthrow. To conserve the species composition and aboveground biomass of forest 704 ecosystems under climate change, which is predicted to increase windthrow frequency (Usbeck 705et al. 2010; Donat et al. 2011; Gregow et al. 2017; Laapas et al. 2019), it is preferable to leave 706 707 the dead wood generated by windthrow and not perform scarification after windthrow. Because this paper describes a case study in hemiboreal forests in northern Japan, forest landscapes in 708 709 different climatic zones and dominated by different tree and grass species might show different results. The promotion of similar studies in various regions worldwide would be useful to obtain 710711a more robust conclusion. Additionally, this study involved assumptions regarding scenario settings (e.g., windthrow regimes and the destruction rate of advanced seedlings). Although 712these assumptions were based on real situations and findings of previous studies as much as 713possible, examining the effects of these assumptions could also be an important topic of future 714studies. Besides, leaving the dead wood after windthrow could result in insect outbreaks (Mezei 715et al. 2017). However, salvage logging to prevent insect outbreaks would decrease the diversity 716

717	of forest ecosystems, which can result in even more serious insect outbreaks (Hughes and
718	Drever 2001). In addition, the simulation results of Dobor et al. (2019) suggest that over 95%
719	of the dead wood generated by windthrow in the landscape must be salvaged to prevent insect
720	outbreaks; however, this is nearly impossible, because pieces of dead wood at inaccessible sites
721	are left in the forest. Therefore, salvage logging could not effectively prevent insect outbreaks,
722	changes forest tree species composition and decreases the aboveground biomass of forests in
723	the long term.

724 Acknowledgments

725	We would like to thank the University of Tokyo Hokkaido Forest for allowing us to
726	use various data, including tree measurement data from the permanent plots in wind-disturbed
727	areas (reference to validations for WT and SL), restoration experimental areas (for SLSC),
728	and environmental data (soil map) for the target area. Additionally, we would like to thank Dr.
729	Robert M. Scheller and the Hokkaido University Ecosystem management laboratory for their
730	helpful discussions about our study. Comments from anonymous reviewers greatly improved
731	the manuscript.
732	
733	Funding
734	Funding for this study was supported by a KAKENHI grant from the Japan Society
735	for the Promotion of Science (Grant Number JP17H01516, 18J20266, and 21J21458); the
736	Environment Research and Technology Development Fund (JPMEERF16S11508) of the
737	Environmental Restoration and Conservation Agency of Japan; and the Ministry of Education,
738	Culture, Sports, Science and Technology Japan TOUGOU Grant Number
739	JPMXD0717935498.
740	
741	Appendix A. Supplementary materials
742	
743	Appendix B. Source code availability

744	The source code for our improved version of LANDIS-II NECN succession is
745	available at https://github.com/hagachi/Extension-NECN-Succession/tree/feature-
746	initdecayrate.
747	
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