



Forest recovery in set-aside windthrow is facilitated by fast growth of advance regeneration

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Abstract

• **Key message** The disturbance of a research plot by a windstorm allowed us to study the role of the seedling bank in the regeneration processes. The released advance regeneration dominated among the saplings; taller individuals retained their position until the end of the study. Pioneer species occurred sporadically. Seven years after the disturbance, the windthrow was covered by a dense thicket of young trees.

• **Context** The dominant role played by advance regeneration in natural regeneration processes after intense wind disturbances is still a matter of dispute.

• **Aims** We took advantage of a windstorm in one of our research plots to study the role of the seedling bank released by the disturbance in the regeneration processes.

• **Methods** We collected data in 70 plots, recording the survivorship of seedlings, annual height growth, and signs of browsing. The height ranking was analyzed with Kendall's concordance coefficient, and the height growth rates were compared using Dunn's test.

• **Results** The density of seedlings increased from 6.7/m² in 2008 to 8.1/m² in 2010 and then decreased to 1.2/m² in 2015. The density of saplings increased continuously from 0.14 to 1.9/m². The highest size differentiation occurred in sycamore maple; the individuals which were taller before the windstorm retained their position until the year 2015. The only species that was recruited mainly from germinants was European hornbeam.

• **Conclusion** The advance regeneration released by the windstorm played a major role in the regeneration process, while pioneer species occurred only sporadically. Seven years after the disturbance, the windthrow was already covered by a dense thicket of young trees.

Keywords Advance regeneration · Forest dynamics · Natural disturbance · Regeneration processes · Seedling recruitment · Temperate forests

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Contribution of the co-authors JS and ZM designed the study, ZM and EM collected the field data, AG and AT conducted data analyses, and JS and AG wrote the text.

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1 Introduction

Natural disturbances have been widely recognized as a driving force in forest ecosystem dynamics (Frelich 2002; Johnson and Miyanishi 2007). While small disturbances usually release the advance regeneration of late-successional species (Woods 2004), large and intense disturbances that can kill most of the canopy trees in the whole stand allow for colonization of the disturbed area by pioneer species (Bobic 2007; Jaloviar et al. 2017). However, the mortality of canopy trees after a large and intense disturbance can be much lower than 100% and the role of pioneer trees can be limited by competition with the remnants of the old stand (Nagel et al. 2006).

Hurricanes, downbursts, and tornadoes are examples of large and intense disturbances (Turner et al. 1997, Greenberg and Collins 2015). Mortality rates of canopy trees after wind disturbance rarely exceed 50% (Wolf et al. 2004; Woods 2004). Partly damaged live trees belong to important legacies (Franklin et al. 2007) of wind disturbances. Therefore, the conditions for the development of a young generation of trees (both advance regeneration and new recruitment) vary according to the local density of mature trees that have survived the disturbance. Windthrows and breakage of stems brought about by severe winds do not necessarily lead to tree mortality (Peterson 2000; Canham et al. 2001). Some species are able to survive and rebuild damaged crowns or to develop secondary shoots from windthrown stems lying on the ground (Peterken 1996). A substantial role in the development of the new generation of trees could be played by vegetative reproduction of these species (Koop 1987).

The rates of regeneration of forest communities following intense wind disturbances vary among different forest types. In tropical forests, regeneration processes can last less than 10 years (Brokaw 1985; Marra et al. 2014), while in boreal forests, the regeneration processes span several decades (Rich et al. 2007). Rates of forest regeneration in temperate forests are difficult to determine, as in most cases large-scale wind disturbances in these forests are followed by salvage-logging and tree planting (Lindenmayer et al. 2012), and the opportunities to study natural processes are very few (Palik and Kastendick 2009; Buma and Wessman 2012; Vodde et al. 2015). According to the results of studies conducted in mixed temperate forest, regeneration processes can be very variable in terms of speed (Gill et al. 2017): in poor habitats regeneration can be relatively slow (Dobrowolska 2015), while under good site conditions regeneration processes can be very fast (Bartels et al. 2016).

In the case of wind disturbance, an important role could be played by the seedling bank, present under the canopy prior to the disturbance (Nagel et al. 2006; Franklin et al. 2007). Seedling banks usually consist of shade-tolerant species and thus can contribute to maintaining a big share of shade-tolerant species in post-disturbance stand development

(Nagel et al. 2007; Palik and Kastendick 2009). They can play a significant role after wind disturbances, as little damage is done by wind to the young trees growing on the forest floor. Moreover, the spatial heterogeneity of light intensity following wind disturbance provided by slowly dying crowns of broken or uprooted trees allows for acclimation to the changing light conditions even for very shade-tolerant species (Čater and Diaci 2017).

One of the important problems associated with natural regeneration after disturbances is the disturbance interactions (Veblen et al. 1994; Buma and Wessman 2011). In coniferous forests, some blown down areas experience more severe wildfires (Kulakowski and Veblen 2007; Buma and Wessman 2011) or insect outbreaks (Stadelmann et al. 2013). In mixed deciduous temperate forests, wildfires and insect outbreaks are rare, but an important interaction can occur between wind disturbance and herbivory by ungulate browsers. Gaps created by windstorms are strongly preferred by foraging ungulates (Bobic 2007), but accelerated growth rates of young trees in wind-disturbed areas (Kupferschmid et al. 2013) could allow the highly palatable species to regenerate successfully after windstorms (Keren et al. 2017). In our paper, we studied regeneration processes after a windstorm that blew down a patch of forest in the Roztoczanski National Park, SE Poland. One of the permanent research plots, which had been measured 1 year before, was located in the middle of the area affected by the windstorm. After disturbance of the entire site, the windthrow was set-aside for studying natural regeneration. The fact that field data were collected immediately before the disturbance allowed us to study the regeneration process in detail; we could follow the fate of young trees released by the disturbance along with the establishment of new cohorts of seedlings.

Advance regeneration plays an important role in the development of the new stands following large and intense disturbances (Veblen 1986, Nagel et al. 2006, Čater and Diaci 2017). This is especially true for windstorms, as wind does little harm to the seedlings and saplings (Franklin et al. 2007). That gives an advantage to shade-tolerant species, which usually form the advance regeneration and are in a better position compared to the seedlings recruited after the disturbance (Diaci 2017); in extreme cases, it can even contribute to the long-term dominance of late-successional species (Jaloviar et al. 2017).

In our study, we formulated five specific hypotheses concerning regeneration processes:

1. Mortality rates of canopy trees after wind disturbance differ strongly among species, and some badly damaged trees of deciduous species survive and are able to regenerate vegetatively.
2. Advance regeneration of shade-tolerant species suffers little damage during the windstorm and is most abundant in the regeneration process after disturbance.

3. Sycamore maple and hornbeam will increase in relative abundance among the regenerating trees because they thrive under high light intensity.
4. In mixed deciduous temperate forests, seedlings of pioneer tree species will establish only during the 3 years after the windstorm; after 4 years the density of saplings is so high that it will prevent further recruitment of shade-intolerant species.
5. Fast growing seedlings and saplings of palatable species are able to reach safe heights despite intense browsing by ungulates in the area of the windthrow.

2 Methods

2.1 Study area

The Roztocze National Park (RNP; coordinates 50° 31'–50°40' N, 22° 53'–23° 07' E) is situated in the central part of the Roztocze Highlands in the South-East part of Poland. The landscape is characterized by long chains of Late Cretaceous limestone hills reaching 390 m a.s.l. surrounded by thick layers of postglacial deposits, sand, or loess. The most common soil types are podzols and cambisols. The climate of the Roztocze region is characterized by warm and usually dry summers (mean monthly temperature in July is 17.2 °C) as well as cold and snowy winters (mean monthly temperature in January is –4.3 °C, long-term average number of frost days is 38.1 per year and the long-term average yearly duration of the snow cover is 71 days). Mean yearly temperature is about 7.3 °C. The yearly amplitude of the mean temperatures often exceeds 22 °C. The growing season lasts usually 210 days. The mean annual precipitation in the Roztocze region ranges from 650 to 750 mm (Kaszewski 2008).

The most common forest types are mixed European beech-silver fir-hornbeam stands (25% of the Park area), Scots pine stands with a small admixture of deciduous trees (19%) and almost pure silver fir stands (8.5%). Out of 40 tree species native to Poland, 32 occur naturally in the RNP. However, 90% of the forest stands are dominated by three species: European beech *Fagus sylvatica* L., Scots pine *Pinus sylvestris* L., and silver fir *Abies alba* Mill. (Izdebski et al. 1992). The current area of the RNP is 8483 ha, 95% of which is covered by forests. The majority of the forests have retained their natural character and they have been recognized as remnants of the European primary forests (Sabatini et al. 2018). The main types of natural disturbance in RNP are windstorms (winter storms and downbursts associated with summer storms) and accumulation of wet snow in the early autumn, before the leaves of deciduous trees are shed. No large wildfires or insect outbreaks have been recorded since the establishment of the park (Maciejewski and Szwagrzyk 2011).

2.2 Field data collection

The original research plot, with a size of 0.5 ha (rectangle 50 × 100 m), was established in 1973 to study changes in species composition and the structure of secondary forest stands. All trees exceeding DBH of 7 cm were measured (DBH and height) and numbered within 10 × 10 m quadrats. A set of circular subplots of 0.5 m² each was established with regular spacing (Fig. 1) to study the natural regeneration of trees and the composition of the herbaceous vegetation. The re-measurements of the plot were completed in 1997 and in 2008, by the end of July. The time between consecutive measurements (1997, 2008) was too long to provide reliable information about the demographic processes in seedlings and saplings.

In August 2008, the area was hit by a windstorm, resulting in a blowdown with a size of about 2.5 ha. The entire research plot was within the limits of the area disturbed by the wind.

In the year 2009, the circular plots were re-measured again, and an additional 36 circular plots for monitoring tree regeneration and forest floor vegetation were added (Fig. 1). Seedlings and saplings growing in the small circular plots were numbered and labeled to follow their fate. In the years 2010, 2011, and 2015 the measurements of the heights of the seedlings and saplings were repeated in all 70 circular sample plots. Signs of ungulate browsing were recorded for all seedlings and saplings in every year throughout the study. The recording of browsing was conducted twice during the growing season (in spring and in summer), and for each individually tagged seedling or sapling, the location of the browsing (top of stem, side branches) was described.

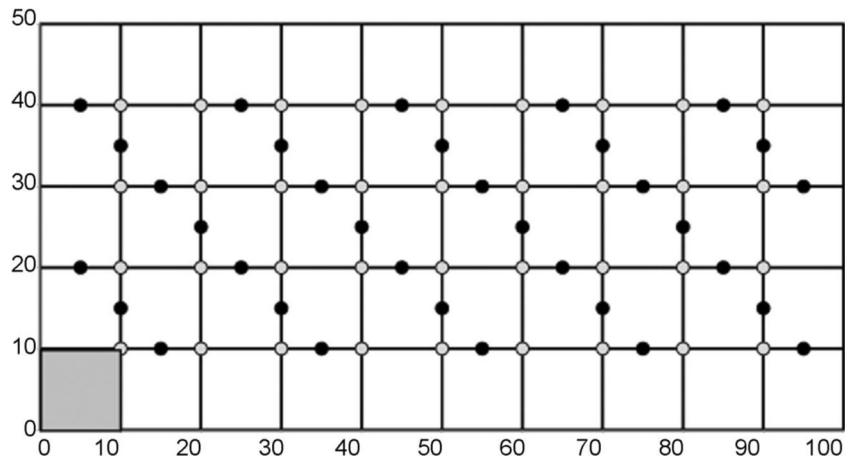
New individuals established in the circular plots were also recorded and measured. The vitality of the mature trees in the wind-disturbed area was checked visually every year at the end of the growing season, and the time of death of each canopy tree was determined.

2.3 Methods of data analysis

First, we analyzed the distribution of the heights of the trees belonging to three major species (sycamore maple *Acer pseudoplatanus* L., European hornbeam *Carpinus betulus* L., and European beech) as well as their growth rates in consecutive periods. We checked the normality of these distributions by statistical tests: visualizing Q-Q (quantile-quantile) plot, Shapiro-Wilk test, and Bartlett's test. The distributions were characterized by high skewness and data heteroscedasticity. However, because the sample sizes for the groups were large enough (Rutherford 2001), we employed Welch's ANOVA followed by the Games-Howell post hoc test along with the non-parametric Kruskal-Wallis test followed by Dunn's test.

To determine the differences in the height increments among the three species over the four studied time periods,

Fig. 1 Scheme of study area; mature trees were measured and mapped in entire 50 × 100 m plot, while circular sample plots (area—0.5 m²) were used for measuring forest regeneration. Filled (34) circles—plots established in 1973, hollow (36) circles—plots established in 2009; one 10 × 10 m quadrat represents basic unit, in which canopy trees were measured and numbered prior to windstorm



we calculated the intra-species Gini coefficients as measures of their distribution inequalities. After arranging the data values in order from the lowest value to the highest value, we employed the following formula (Dixon et al. 1987):

$$GST = \ln 2 \sum x_i = \ln(2i - n - 1)x_i$$

where:

- GST Gini coefficient value for species S in time period T,
- n the number of individuals for species S in time period T,
- x_i height increment value for i th individual,
- \bar{x} mean value for all individuals of species S in time period T.

To determine whether the young trees maintain their initial height advantage, we compared the ranks of individual trees among the consecutive measurements. We tested the consistency of height rankings during the study period using Kendall’s coefficient of concordance (KCC) (Legendre 2005). In our analysis, we took into account only the trees for which we had height measurements in all years, i.e., in 2009, 2010, 2011, and 2015. We calculated the KCC for each pair of height rankings from the above-mentioned years along with chi-squared significance tests to identify which of them were concordant with one another and what was the strength of this association.

To compare the intensity of the ungulate browsing among tree species, we used the data showing how many times the individual trees had been browsed during the period of our study. We analyzed the number of browsing episodes along with the species identity of young trees. As the Cameron and Trivedi test (Cameron and Trivedi 1990) suggested that data may be underdispersed, we used a generalized Poisson regression (Hilbe 2014) to estimate the parameter values in the model where the response variable was the number of browsing events per individual and its species identity was an explanatory variable. The indicator variable was the expected difference in the log count between particular species and the

reference group of sycamore maple. Hence, the incidence rate for particular species was the exponent of the coefficient value multiplied by the incidence rate for the reference.

Data availability The datasets generated during and/or analyzed during the current study are available from the corresponding author on reasonable request.

3 Results

3.1 Tree mortality after windstorm

Prior to the windstorm, the stand density was 534 trees/ha and the basal area (BA) was almost 40 m²/ha. According to tree numbers, Scots pine was less than 42%, hornbeam almost 30% and beech 24%. By BA, Scots pine was the dominant species, with almost 47% of BA, beech was slightly over 31%, and hornbeam had less than 16% (Table 1).

The total mortality in the study plot amounted to 113 trees, i.e., 40.45% of all trees prior to the windstorm. Mortality calculated on the basis of BA was even higher: 48.94%. The only species that was completely eliminated from the stand was wild cherry *Prunus avium* (L.) L. The highest mortality by tree number was recorded in Scots pine (59%). High mortality (48%) occurred also in European beech and in sessile oak *Quercus*

Table 1 Stand characteristics in 0.5 ha study plot prior to windstorm

	Number of trees	Basal area (m ²)
European beech	64	6.23
Wild cherry	2	0.34
Sessile oak	4	0.33
Scots pine	111	9.26
European hornbeam	79	3.16
Sycamore maple	7	0.55
Total	267	19.86

Table 2 Tree mortality after windstorm

	Number of dead trees in year					Cumulative tree mortality (%) after 7 years
	2009	2010	2011	2012	2015	
European beech	1	14	16			48.4
Wild cherry		2				100.0
Sessile oak	1		1			50.0
Scots pine	50	2	9	1	4	59.5
European hornbeam	2		4			7.6
Sycamore maple			1			14.3
Total	54	18	31	1	4	40.4

petraea (Matt.) Liebl. (50%). The mortality rate calculated by BA was higher in European beech (61%) than in Scots pine (55%).

The mortality was distributed unevenly among the tree species in consecutive years after the windstorm. Most of the Scots pines that were broken or uprooted by the windstorm died in the first year, while in the case of European beech the mortality was concentrated in the second and third years after the windstorm (Table 2). In European beech, almost 30% of the uprooted trees survived until 2015. The extreme case was European hornbeam where out of 43 uprooted trees only five had died by the year 2015, and out of nine broken hornbeams, eight trees survived and rebuilt their crowns, so total mortality in that species amounted to less than 8%.

3.2 Changes in densities of seedlings and saplings

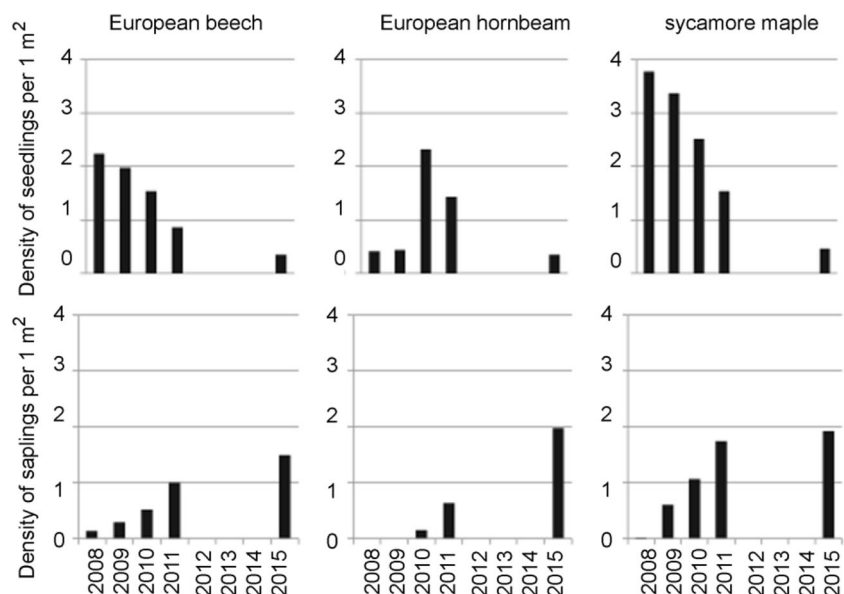
The density of seedlings prior to the windstorm amounted to 6.7 individuals per square meter (ind./m²), and the most abundant species among the seedlings was sycamore maple (56%)

followed by European beech (33%). At the same time, the density of the saplings was much lower (0.14 ind./m²), and the most abundant species among the saplings was European beech (89%), sycamore maple had 9%, and the other species occurred only as single individuals (Fig. 2).

The total densities of seedlings of all species increased slightly in the first 2 years following the windstorm and reached 8.1 ind./m² in 2011; then they decreased to 4.4 ind./m² in 2011 and 1.2 ind./m² in 2015 (Fig. 2). That decline was a result of two processes: height growth resulting in reaching the sapling stage (height larger than 0.5 m) and mortality increasing with time as the sapling layer grew denser. European beech was abundant among seedlings since the beginning of this study (33% of all seedlings in 2008), and it slightly increased its share among seedlings by 2015. Seedlings of Scots pine arrived in 2010 and were still present in 2011; however, by the year 2013, they had died.

The density of the saplings increased throughout the period under study: it reached 0.3 ind./m² in the year 2009 and increased steadily to 1.9 ind./m² in 2015. Already in 2009,

Fig. 2 Densities of seedlings and saplings in study area in years 2008–2015



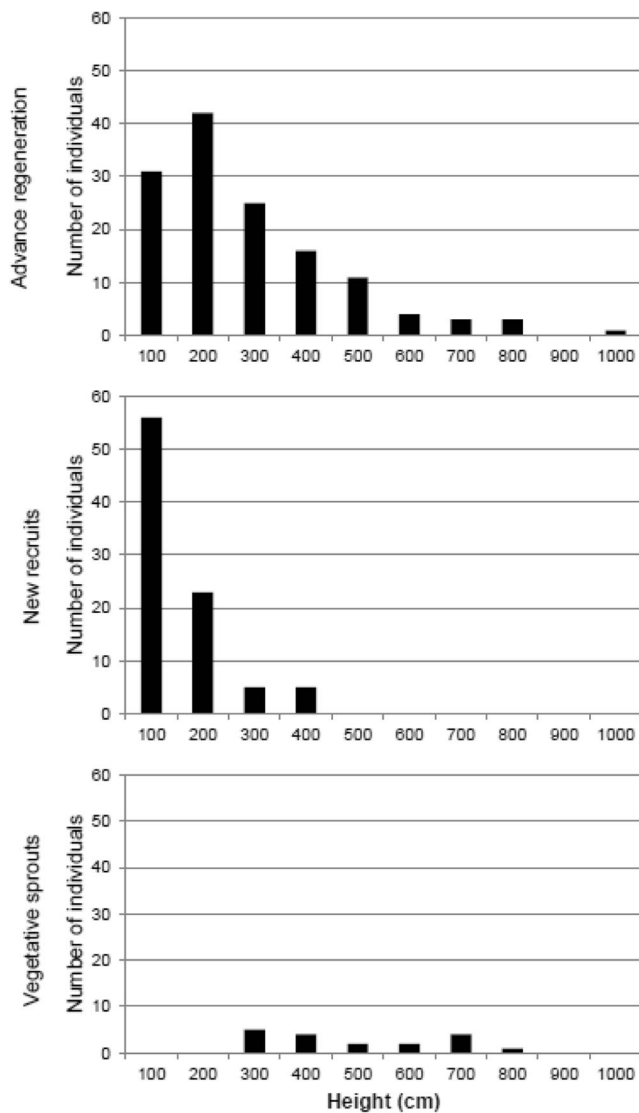


Fig. 3 Comparison of height distributions of advance regeneration, new recruits and vegetative sprouts 7 years after windstorm

sycamore maple was the most abundant species among the saplings (69%); in the next year, its share remained high and then gradually declined to 34% in 2015. In the same time, the share of European hornbeam among the saplings increased from zero in 2009 to 35% in 2015. European beech was dominant among the saplings prior to the disturbance (89% of saplings in 2008). In the years 2009–2015, it maintained an almost constant share among the saplings (Fig. 2).

3.3 Growth rates of young trees

The three main groups of young trees were the advance regeneration—released by the disturbance, the new recruits—established after disturbance and vegetative sprouts of the trees damaged by the wind. The advance regeneration was the most numerous group, attaining the greatest heights. Newly recruited young trees were much lower than the

advance regeneration. The vegetative sprouts were less numerous, but they had the largest mean height of these three groups (Fig. 3).

The analysis was focused on the three most common species: European beech, hornbeam, and sycamore maple. The average heights of the young generation of trees in the wind-disturbed area ranged in the year 2015 from 190.6 cm for European hornbeam to 230.5 cm for sycamore maple. However, the height distributions were strongly right-skewed, with many low saplings and a few tall ones (Fig. 4). The tallest young tree recorded in our plots was sycamore maple with the height of 940 cm; the tallest European hornbeam reached 755 cm and the tallest European beech 600 cm. The tallest specimens of European hornbeam were vegetative shoots produced by uprooted mature trees, and their average height was 458.6 cm. The young trees of generative origin for hornbeam were on average 115.2 cm tall.

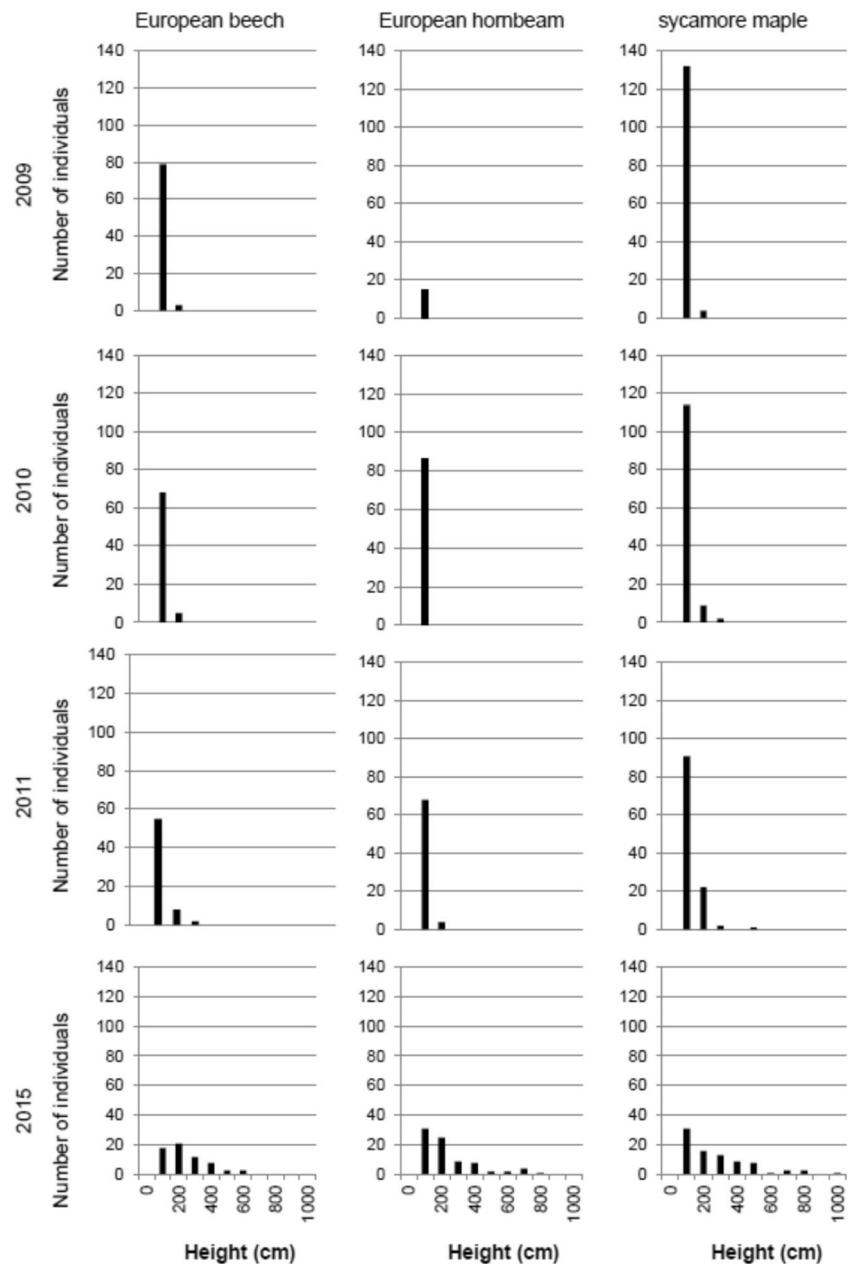
Height increments for the entire period under study (2009–2015) were calculated only for the young trees present throughout the entire period under study, and that group amounted to 232 individuals. The height increments were strongly right-skewed (Fig. 5), with average values ranging from 165.5 cm in European beech through to 189.2 cm in European hornbeam and to 198.7 cm in sycamore maple. Height increments calculated for young European hornbeams of generative origin were much smaller (113.4 cm). The largest height increment (824 cm) was recorded in sycamore maple.

The results from comparing the mean height increments among the tree species using Welch's ANOVA followed by the Games-Howell post hoc test were consistent with the results of a non-parametric Kruskal-Wallis test followed by Dunn's test (Table 3). The European beech had significantly smaller height increments than sycamore maple and European hornbeam in the beginning of the study, but for the entire period, the differences among those three species were insignificant (Table 3). European beech was the least variable in height growth, while in European hornbeam and sycamore maple, apart from individuals with large height growth rates, there were also many slow growing individuals (Fig. 5). This was also supported by the results of calculating the Gini coefficient, where the highest values were ascertained for sycamore maple and the lowest values for beech (Table 4).

3.4 Size hierarchy among young trees

On average, the European beech had the lowest Gini coefficient, and sycamore maple had the largest one. The values of the Gini coefficient fluctuated among years, and the only exception was hornbeam, in which the initial value of the coefficient was relatively low and rapidly increased in the second year of the measurements (Table 4).

Fig. 4 Height distributions of young trees in wind-disturbed area in years 2008–2015



On the basis of Kendall's concordance coefficient (Table 5), we rejected the null hypothesis that any two height rankings were uncorrelated at the 0.05 significance level. In general, trees that had been taller in the beginning of the study tended to be also taller in the year 2015, and the tendency among smaller trees was consistent in a similar way (Table 5).

3.5 Sum of height of young trees per area unit

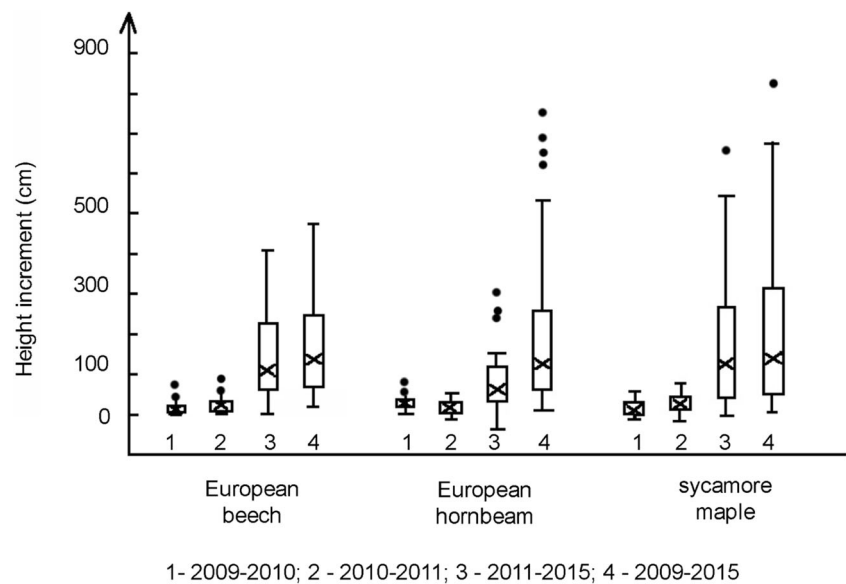
The sum of heights of the young trees increased from 2 m/m² in 2009 to almost 14 m/m² in 2015. The most spectacular increase was recorded in the case of European hornbeam: the sum of the height of seedlings and saplings for that species

in 2009 was 10 times lower than the sum of young individuals of European beech, and in the year 2015, the sum of the height of young European hornbeam was substantially higher than that of European beech (Fig. 6). The species with the greatest sums of height was sycamore maple.

3.6 Natural regeneration and deer browsing

Browsing of seedlings and saplings by ungulate herbivores was common throughout the study period; out of 127 young individuals of European beech, European hornbeam, and sycamore maple present in the sample plots from 2009 to 2015, 102 were browsed at least once. Only 25 individuals present

Fig. 5 Comparison of height increments of young individuals: 1—in the first 2 years after disturbance, 2—in the third and fourth years after disturbance, 3—from the fourth to seventh years after disturbance, and 4—over period of 7 years after disturbance. x—median, filled circle—outliers



from 2009 to 2015 were not browsed at all. The most browsed species was sycamore maple and the least browsed was beech (Table 6).

The results of the generalized Poisson regression indicated that European beech was about 0.37 times less likely to be browsed than sycamore maple. The difference between sycamore maple and hornbeam (0.43 times less likely to be browsed) was insignificant (Table 7). A similar model run without sycamore maple suggested that European hornbeam was more frequently browsed than European beech (coefficient ~ -0.843) and that the difference was statistically significant (p value ~ 0.02).

4 Discussion

This study fortuitously captured data immediately before wind disturbance, and then we were able to follow the dynamics of the regeneration processes from there on a yearly basis, until the young trees in the study area had formed a dense thicket. Despite the small size of the study area and lack of repetitions,

this has provided a unique and clear picture of the regeneration processes.

4.1 Mortality rates and patterns in canopy trees

The tree mortality after a windstorm (40%) was similar to that in many other studies on wind disturbance (Peterson 2000; Woods 2004; Szwagrzyk et al. 2017). European hornbeam seems to be most adapted to regenerating in a vegetative way (Faliński and Pawlaczyk 1993), and sycamore maple also regenerated well, especially trees that had been uprooted by the wind. In European beech, the ability to rebuild the crown was very low, and that agrees well with the results of earlier work (Fichtner et al. 2013). The process of mortality following the windstorm was prolonged; only in the case of Scots pine did all broken and uprooted trees die within 1 year after the disturbance.

The tree species present in the study area responded to the disturbance in different ways. European beech was characterized by high mortality and abundant recruitment. Two species—European hornbeam and sycamore maple—displayed low mortality and high recruitment. Scots pine was a loser

Table 3 Results of Dunn’s test (presented in form of table filled with p values computed for each pairwise comparison) for height increments of young individuals of various tree species (* $p < 0.05$; ** $p < 0.01$, *** $p < 0.001$)

	2009–2010		2010–2011		2011–2015		2009–2015	
	European beech	Sycamore maple	European beech	Sycamore maple	European beech	Sycamore maple	European beech	Sycamore maple
European hornbeam	0.0000***	0.0000***	0.1079	0.4224	0.0468*	0.1854	0.1987	0.3833
Sycamore maple	0.074		0.1305		0.1970		0.2812	

Table 4 Gini coefficients for height increments of young individuals of three species over various time periods

Gini coefficient				
	2009–2010	2010–2011	2011–2015	2009–2015
European beech	0.4347693	0.3493414	0.4084994	<i>0.4052743</i>
European hornbeam	0.2784479	0.4836758	0.4567751	<i>0.4701092</i>
Sycamore maple	0.5999638	0.457208	0.5092737	<i>0.4974311</i>

Italic numbers are statistically significant values

as it suffered high mortality and had no effective recruitment. According to the classification of Batista and Platt (2003), the trees present in the study area could be classified as resilient (European beech), usurpers (European hornbeam and sycamore maple), and susceptible (Scots pine). No tree species displayed the resistant syndrome, which means low mortality and low recruitment.

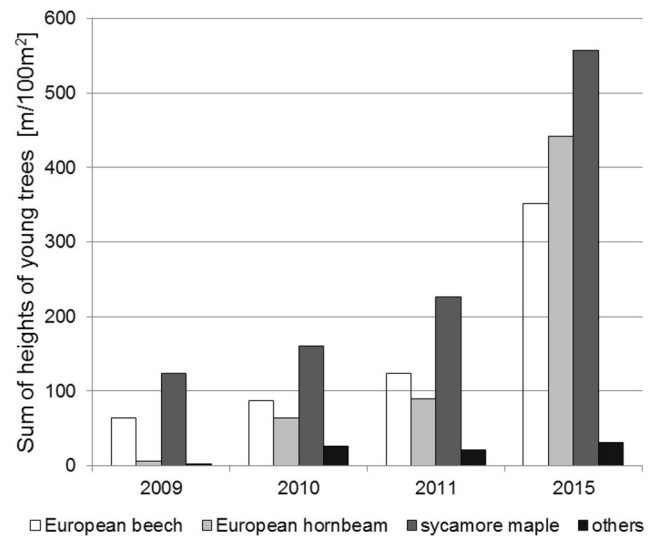
4.2 Role of advance regeneration

Before the windstorm, the understory of the analyzed stand was covered with abundant seedlings, mostly of sycamore maple; however, the growth of this species was suppressed, so among the saplings, the most abundant species was European beech. The strategy of increasing the likelihood of long-term survival by reducing height growth rates has been described by Woods (2008) for sugar maple *Acer saccharum* Marshall. Our observations suggest that sycamore maple responds to light conditions in a similar way. On the other hand, some individuals of European beech growing in a similar light environment were able to attain the sapling stage, and this agrees well with the well-known shade tolerance in beech, which is higher than in any other deciduous tree in the Central European temperate forests (Ellenberg 1988).

The released advance regeneration formed the largest group among the young generation of trees, and along with vegetative sprouts, it dominated among the tallest saplings. That leaves little room for the new recruits; however, they were still abundant, but their chances in competition for light

Table 5 KCC values for each pair of height rankings during investigated time interval

Height rankings	Kendall's W
2009, 2010	0.68
2009, 2011	0.614
2009, 2015	0.534
2010, 2011	0.779
2010, 2015	0.556
2011, 2015	0.616

**Fig. 6** Sum of heights of young trees [m/100 m²]

with their taller neighbors are low (Čáter and Diaci 2017). The dominance of advance regeneration is not surprising, as similar situations had been already described (Nagel et al. 2006; Vodde et al. 2015).

4.3 Advantages of fast growth rates

The growth of the advance regeneration was surprisingly fast. The maximum height growth rates in sycamore maple reached 163, 195, and 190 cm in the 3 years after the windstorm. In European hornbeam, the growth rates of new stems of vegetative origin were even higher. In the year 2015, 7 years after the disturbance, the sum of the heights of the young trees had reached 1380 m/100 m²; this value is almost twice as large as the sum of the heights of trees in 30-year thickets of European beech growing under good habitat conditions (Szymkiewicz 1971).

Individuals that were taller than their neighbors at the time of the windstorm maintained their dominant position throughout our study. Suppressed individuals present under the forest canopy as well as seedlings established after the disturbance played a minor role in the regeneration process. Similar findings were reported by Uriarte et al. (2004) from subtropical forests of the Caribbean affected by recurring hurricanes.

The extremely high growth rates displayed by sycamore maple and European hornbeam in the area affected by the windstorm indicated that these species are able to take advantage of the resources (light, nutrients) that were made available by the mortality following the disturbance. A similar situation was described for sugar maple by Leak (1987). Thus, European hornbeam and sycamore maple seemed to benefit from the wind disturbance. Both species were classified as “competitors” within the framework of the Grime’s classification of life history strategies (Brzeziński and Kienast 1994). However, in the context of the findings from recent studies

Table 6 Frequency of browsing [%] for three species in years 2009–2015

Numbers of years when individual was browsed	European beech	European hornbeam	Sycamore maple
0	45	11	3
1	39	22	23
2	14	44	32
3	2	22	30
4	0	0	12

(Maciejewski and Szwagrzyk 2011; Szwagrzyk et al. 2012), their life history strategies should be considered as a combination of “C” (competitor) and “R” (ruderal) strategies.

4.4 Marginal role of pioneer species

In the first year after the windstorm, the crowns that were still alive of broken or uprooted trees covered much of the blow-down area, thereby creating partly shady conditions on the forest floor; similar conditions have been reported from other wind-disturbed areas left without any salvage-logging (Peterson 2000). Seedlings that were already present prior to the disturbance took advantage of the release (Veblen 1986; Nagel et al. 2006) and grew at high rates; in addition, herbaceous vegetation developed rapidly in the blowdown area. Like in other studies (Peterson 2000; Vodde et al. 2015), the only places suitable for the establishment of pioneer tree species were in the pits and mounds created by the windfallen trees. The single species that germinated in large numbers after the windstorm was European hornbeam, and this was probably due to abundant seed rain (Jankowska-Błaszczuk and Grubb 1997). In addition, wild cherry germinated after the disturbance, but was then eliminated by browsing pressure. Typical pioneer species, like silver birch *Betula pendula* Roth., goat willow *Salix caprea* L., and European aspen *Populus tremula* L. (Ellenberg 1988), played minor roles. In the first year after the windstorm 97 seedlings of Scots pine were found in 70 circular sample plots, but the number of Scots pine germinants declined dramatically. Most of the pine seedlings died in only a few years; only one of two specimens growing on elevated root plates attained the sapling stage. Similar results have been reported in several studies (McClure and Lee 1993, Cook 2000), and their authors suggested that a mineral seedbed is a factor controlling the

Table 7 Results of fitting generalized Poisson regression model to deer browsing count data

Species	Coefficient	<i>p</i> value for significance test
European hornbeam	−0.2447	0.1303
European beech	−0.9932	0.0055

regeneration of shade-intolerant, opportunistic species, and such microsites are very limited in wind-disturbed areas not subjected to salvage-logging.

Shrubs, like European elder *Sambucus nigra* L., red-berried elder *Sambucus racemosa* L., common dogwood *Cornus sanguinea* L., or dog rose *Rosa canina* L., took advantage of the disturbance and established on the elevated root plates and on windthrow mounds. The study by Herrera and García (2009) suggested that birds, using elevated places as perches, can disperse the seeds to wind-disturbed areas.

4.5 Role of deer browsing in regeneration process

Prior to the windstorm, 40% of the sycamore maple and 21% of the European beech saplings had been browsed by deer. These figures are comparable with the levels of browsing pressure recorded in other national parks in southern Poland (Miścicki and Żurek 2015; Bodziarczyk et al. 2017). After the windstorm, the percentage of browsed saplings increased to 61%; however, the high growth rates of the young trees in most cases compensated for the losses resulting from browsing and allowed the saplings to escape browsing of terminal shoots by growing tall enough in one or two growing seasons. An additional factor was the rapid arrival of numerous saplings of European hornbeam, and that species is strongly preferred by deer (Faliński and Pawlaczyk 1993), but according to recent findings, it is also the most resilient species growing in the temperate deciduous forest of Central Europe (Kuijper et al. 2010). Apparently, in the few years after the windstorm, the browsing pressure by ungulates was not able to slow down the regeneration processes of the trees. The successful growth of the seedlings and saplings could be partly attributed to the rapid development of herbs and shrubs in the windthrow area, which provided an alternative food source for the ungulates. However, the ungulate herbivory probably contributed significantly to the elimination of the pioneer species: willows, aspens, and wild cherries.

5 Final remarks

In the 7 years since the windstorm, the analyzed stand was a patchwork of groups of mature trees that survived the disturbance and dense young thickets dominated by released advance regeneration. European beech, which used to be one of the dominant species in the tree stand prior to the windstorm, lost many trees in the canopy, but retained its position due to the ample advance regeneration released by the disturbance. European hornbeam survived the windstorm very well and regenerated in the disturbed area in generative and vegetative ways. The species that was much more abundant in the regeneration than in the mature stand prior to the windstorm was sycamore maple.

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Compliance with ethical standards

Conflict of interest The authors declare that they have no conflicts of interests.

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