Is livestock grazing a key factor for changing vegetation patterns in lime rich coastal dunes in the Netherlands?

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Received: 26 February 2018 / Revised: 27 January 2020 / Accepted: 4 February 2020 / Published online: 19 February 2020
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Abstract
In 1990, livestock grazing was introduced in Meijendel, a 1800 ha lime-rich coastal dune area, at a density of 0.06–0.07 LLU ha⁻¹ year⁻¹ (1:12–18 ha) to counteract encroachment of tall grasses and shrubland on dune grassland and increase the bare sand area. Monitoring was based on four digital orthophotos (1975–1990–2001-2009) with a high spatial resolution (pixel size 25 × 25 cm). The changes were tested using Generalized Estimating Equations. Habitat changes occurred, but contradicting our hypothesis, there was no significant impact from the grazing on bare sand, grassland or shrubland within 11 and 19 years post livestock introduction. (1) After several decennia of decreasing bare sand, there was a significant increase between 2001 and 2009, irrespective of livestock presence. (2) The changes in grasslands and shrublands are independent of the livestock, but dependent on distance to the coast. (3) Bare sand and shrub cover determine the space left for the dune grasslands. It appears other factors than livestock grazing must have induced the changes. Changes in climate conditions and nitrogen load might have stimulated bare sand. An interaction with the end of Marram planting in 1990 cannot be concluded from available data. The disease-led reduction of rabbit grazing from the mid-1950s led to an expansion of the dominant shrub Hippophae rhamnoides. However, Hippophae shrubland typically regresses to grasslands on its collapse after 25–40 years. Tree species like Crataegus, Betula and Quercus will gradually dominate the landscape for far longer. Active removal of these indigenous species is necessary to prevent future loss of dune grasslands.

Keywords Coastal dunes · Blowout · Aerial photographs · Livestock grazing · Oryctolagus cuniculus · Hippophae rhamnoides

Introduction
Lime-rich coastal sand dunes in The Netherlands are species rich and harbour 66% of the Dutch flora (De Vries et al. 1994). Under the Natura 2000 legislation (Council of the European Communities 1992) almost all habitats of the coastal sand dunes are protected. These habitats are a mosaic of bare sand, dune grasslands, shrubland and forests. ‘Grey dunes’, i.e. fixed coastal dunes with herbaceous vegetation, are designated as a priority habitat type (H2130) by Natura 2000. Grey dunes not only contain a great number of vascular plants, but they are also rich in bryophytes, mosses and lichens (Schaminée et al. 1995, 1996, 1998; Stortelder et al. 1999).

During the last five decades, the vegetation in Dutch coastal dunes has changed considerably. Patches with bare sand and open grasslands have changed into areas mainly dominated by tall grasses and sedges, shrubs and forest that has gradually led to a rather monotonous vegetation and a considerable loss of local biodiversity (Ketner-Oostra and Sýkora 2004; Kooijman et al. 1998; Veer and Kooijman 1997). These processes have been associated with several phenomena: (1) the outbreak of
Myxomatosis and Viral Haemorrhagic Disease (RVHD) in populations of rabbits (Oryctolagus cuniculus), (2) routine planting of Marram grass (Ammophila arenaria) and shrubs and trees to stabilize dunes, (3) planting of shrub and tree species for mainly aesthetic reasons and wood production, (4) a high input of airborne nitrogen, (5) changes in land use, including abandonment of agricultural practices like livestock grazing and crop cultivation, (6) the impact of changes in climate conditions, (7) an ongoing anthropogenization of the landscape (e.g. recreation), accompanying a rapid spread of introduced non-native species, and (8) autonomous succession (Provoost et al. 2011; Remke et al. 2009; Noordijk 2007; Jones et al. 2004; Eerens and Van Dam 2001; Drees and Olff 2001; Kooijman et al. 1998; Ten Harkel 1998; Ten Harkel and Van der Meulen 1996; Schaminée et al. 1995, 1996, 1998; Stortelder et al. 1999; Bakker et al. 1974; Boerboom and Westhoff 1974; Ranwell 1960). As large herbivores were thought to counteract the process of stabilisation and monotonisation (Van Dijk 1992; Kooijman and De Haan 1995; Provoost et al. 2011) and the fact that large herbivores strongly prefer graminoid species (Lamoot et al. 2005), extensive grazing by domestic livestock has been introduced in many nature reserves in The Netherlands and elsewhere.

In late 1990, year-round grazing by livestock was introduced in the coastal dunes of Meijendel near The Hague, The Netherlands (GPS 52°7’N; 4°20’E). The livestock density was 0.06–0.07 LLU/ha−1·year−1 (1:12–18 ha). Generally speaking, Wallis de Vries et al. (1998) consider this to be a relatively low level of impact. However, the level of LLU had been adjusted to the low level of palatable biomass in coastal sand dunes (e.g. Ebrahimi 2007) and expected winter survival of the animals. In some years, some animals of the herds barely survived winter, which meant a relatively high impact on this nutrient-poor environment. By trampling and grazing, the vegetation was expected to be damaged to a large degree, opening up the shrubland and forest to the benefit of species rich grasslands (see De Bonte et al. 1999). The hypothesis of this study is that the introduction of livestock will lead to regressive succession, as expressed in an increase of bare sand, a decrease in the area of shrubland, and a subsequent increase of dune grassland. This paper focusses on the changes in area cover of these three classes (bare sand, grassland and shrubland), by comparing three livestock-grazed areas and three ungrazed controls within the coastal dunes of Meijendel in The Netherlands.

**Material and methods**

**Aerial images**

Along the Dutch coast, sequential mapping of aerial photographs has shown drastic changes in vegetation cover (e.g. Van Dorp et al. 1985; Kruijssen et al. 1992; Van Til and Mourik 1999; Janssen 2004), but in complex systems like coastal sand dunes, manual and especially manual sequential mappings have their limitations (Janssen 2004; Assendorp 2010). Drawing lines between vegetation types in discrete landscapes with crisp (sharp) boundaries is relatively easy. But drawing lines in a landscape with fuzzy spatial transitions between vegetation types, such as coastal dunes where every metre can be different to the next, the interpretation of vegetation types is highly dependent on personal interpretation. Therefore, other remote techniques are required and used for vegetation mapping and land surveys (e.g. Campbell 2006; Groom et al. 2006; Potter 2013). Assendorp (2010) demonstrated that geometrically and radiometrically corrected orthophotos with a high spatial resolution largely overcomes the difficulties of manual mapping.

Four sets of high-resolution false colour infrared digital orthophotos of Meijendel were available with clear time intervals: 1975, 1990, 2001 and 2009. All four were taken in the same period of the year (end June/beginning July). The years 1975 and 1990 are ±330 2 μ scanned analogue images (34 × 34 cm), which have been geo-referenced and radiometrically corrected to one orthophoto. The years 2001 and 2009 were
direct digital recordings and also radiometrically corrected. For the technical details of orthophoto production see Assendorp (2010). The feature space data were processed in ArcGis 10.3. The accuracy of the classification of scanned...
analogous false colour images of 1975 and 1990 and of the
digital photos of 2001 and 2009 were 80–95% as demonstrat-
ed by Assendorp (2010). This percentage is generally accept-
ed as appropriate for a vegetation structure map (Foody 2002;
Assendorp 2010). All four sets of orthophoto images were
resampled to a pixel size of 25 × 25 cm. At this resolution,
bare sand and shrubland/trees are defined (crisp) objects and
consist of multiple pixels in the image. Grasslands differ with-
in a 25 × 25 cm pixel and have a reflection at the subpixel level
(fuzzy). Bare sand has a specific reflection and is easily sep-
In the six
vegetation cover and no grey humic sand on the surface.
Shrubland/trees, defined as 100% coverage of woody plants,
are also easily distinguished from the rest in a traditional su-
terred within one class.

**Selection of research areas**

As the focus of this research is on groundwater inde-
dependent dunes, dune slacks and water bodies were left
out and in the case of infiltration ponds given an extra
margin of 10 m around. Similarly, all human-disturbed
open areas were left out. These include (i) areas with
extraction wells for artificial recharge for drinking water
production, (ii) intentionally denuded areas to stimulate
blowouts, (iii) the foreshore and Fore Dunes as grazing
is prohibited, (iv) parts of the Parabolic dunes which
were intentionally planted with shrubs and trees, and
(v) the larger dune valleys as these valleys are an
unnatural feature, and have no ungrazed counterpart. Up to
1990, all over Meijendel except for the Fore Dunes
compulsory plantings of Marram grass was equally
applied.

From West to the Central and the East, the test grazing
areas are represented by the Helmduijen (43 ha), Kijfhoek/
Bierlap (65 ha) and Pallantsduin (43 ha), whereas Prinsenduin
(43 ha), Lange Strooken (65 ha) and Vinkenduin (43 ha) re-
present the ungrazed control areas (Fig. 1); in total 302 ha. The
study areas are positioned within the areas of the (ii) Parabolic
Dunes and the (iv) Inner Dunes (Van der Meulen et al. 1985).

In late 1990, year-round grazing with Galloway cattle and
Norwegian Fjord horses was started, just after the 1990
orthophotos were taken. It is assumed that the impact of rabbit
grazing on the three vegetation classes is equal. The rabbit
populations fluctuate over the years because of variations in
the impact of epidemic diseases (Myxomatosis from around
1954 and RVHD in 1989, just before the introduction of the
livestock). Rabbits are counted, but the transects cross over
the areas of this research.

### Statistical analysis and confounding factors

With late 1990 as the starting point of grazing, the changes of
the crisp vegetation classes in hectares between 1975 and
1990 as well as between the test starting date of 1990 and
2001 and equally between the test starting date of 1990 and
2009 have each been calculated per research area in order to
view time-dependent impact of grazing on the outcome, while
acknowledging the changes in impact by starting levels. The
interval between 1975 and 1990 is considered as a period
without large herbivore grazing. By subtracting the outcome
per plot at the various times from status at the start, data are
transferred into change values: the change in amount of hect-
ares covered by the various vegetation types per time interval
(before and after start of grazing) per period. Two main sets of
data were obtained: one with grazing and the other without
(control). Subsequently the values were tested as to what ex-
tent grazing affected the relative distribution of the three types
of vegetation using Generalized Estimating Equations (GEE)
analysis with dummy variables (Kleinbaum et al. 1998). As
potentially confounding factors, the starting values (at \( t = 0 \))
and the relative locations in the dune area (1 = West; 2 =
Central; 3 = East; Fig. 1) were taken into account as variables.
The regression equation (eq. 1) used was:

\[
y = (a_1 x_1) + (a_2 x_2) + (a_3 x_3) + a_4
\]

in which:

- \( y \quad \text{change in hectares per time interval} \)
- \( x_1 \quad \text{dummy variable, taking the values 0 for not grazed control area and 1 for grazed area} \)
- \( x_2 \quad \text{time interval} \)
- \( x_3 \quad (x_1) \times (x_2) \)
- \( a_1 - 4 \quad \text{coefficients} \)

In a second stepwise regression analysis confusing was
checked by the introduction of location: West, East and
Central, all coded by a discrete character, and starting value
at 1975 to detect potential impact by this factor on the out-
come. In the latter analysis, variation in the change of hectares
was associated with variation of the above factors, together
with dummy coding (0: ungrazed versus 1: grazed).

Fitting of the equation model was evaluated by Wald Chi-
square test. GEE analyses were done using STATA, version 12
(StataCorp, College Station, Texas, US). Graphic representation
was performed via GraphPad Prism, version 6 (LaJolla, CA,
USA). Throughout the study, using two-sided testing, \( P \) values
of 0.05 or lower were considered to be of statistical signi-

Apart from the statistical analysis on the changes, the
weighted average of the three crisp classes of the three grazed
and three ungrazed controls is calculated and fitted to figures
in MS Office Excel 2010.
Results

Impact by livestock

For the period of fifteen years before (1975–1990), and respectively eleven (1990–2001) and nineteen (1990–2009) years after the start of livestock grazing, the changes in relation to the start of livestock grazing (1990) are presented (Fig. 2a–c) as well as the weighted average of the three areas of grazed versus ungrazed (Fig. 3a, b). The changes in hectares of bare sand, grassland and shrubland between grazed and ungrazed areas are - in contrast with the hypothesis - not different (Fig. 2a–c; Table 1). A similar conclusion was reached for the mean proportion of the three structural types (bare sand, grassland and shrub; Fig. 3a, b). A decline of bare sand up to 2001 and an increase of shrubland up to 1990, as direct outcomes of progressive succession, is clearly visible in both the grazed and ungrazed control areas.

The results for each crisp class is described in different sections. The classes of bare sand and shrubland are described first, because bare sand and shrub development seem to determine the space left for the dune grasslands.

Changes in bare sand

Between 1975 and 1990 and 1990–2001, there is a steady decrease in the area of bare sand. In the period 2001–2009, an increase in bare sand is visible, though not significant (Fig. 2a; Table 1). Before the introduction of livestock, between 1975 and 1990 aerial photographs show that bare sand mainly evolves into grasslands, but no significant differences could be observed between the grazed and ungrazed areas because in the pre-1990 time frame they were both ungrazed. From 1990 onwards, we expected clear differences due to grazing. Surprisingly, there is no difference between the grazed and the ungrazed areas, and -surprisingly- the change in the ungrazed areas is more pronounced, but statistically not different.

Between 2001 and 2009, there is a clear increase of bare sand both in the grazed and ungrazed sites with no significant difference ($p > 0.05$; Table 1). Time ($p < 0.04$) and location ($p < 0.01$) explains the differences in the changes in area of bare sand, irrespective of grazing. There is an overall reduction in the bare sand areas between 1975 and 2009, while the reduction in the central dune area is more pronounced than in the west and the east dune areas. Again, there is no significant effect because of livestock grazing.

Fig. 2 Changes in relation to the year 1990 of the three classes (bare sand, shrubland, grassland) area cover in hectares; mean ± 1 x standard deviation of three grazed and three not grazed areas. (a) Changes in relation to the year 1990 in bare sand area cover in hectares. (b) Changes in relation to the year 1990 in shrubland area cover in hectares. (c) Changes in relation to the year 1990 in grassland area cover in hectares.
Within the studied time interval of 34 years, the coverage of bare sand varied between 2.7% and 21.5% (Fig. 3a, b; online appendix Table 1a, b). The grazed and ungrazed areas show similar trends in bare sand cover. In 1975, the weighted average proportion of bare sand was 14.5% in the pre-1990 grazed area and 21.5% in the control. In 1990, it had decreased to 8.9% and 8.4% in the post 1990 grazed versus the control area. The lowest percentage was reached in 2001 with 5.8% and 2.7% after which it increased to 8.3% and 6.2%, respectively.

### Changes in shrubland

Between 1975 and 1990 prior to the start of the grazing, shrubland increased by an average of three hectares in both grazed and ungrazed sites (Fig. 2b; Table 1). This is according to the expectations, because all sites were ungrazed before 1990. Between 1990 and 2001, the average change was a decrease of about 4 ha, again in both grazed and ungrazed areas. In between 2001 and 2009 however, there is almost no change in the extent of shrubland. Again, location ($p < 0.01$) is the parameter explaining the changes in shrubland between the grazed and the ungrazed areas. In the west part of the dunes, the decrease in shrub area is significantly ($p < 0.01$) higher than in the central area with the central being significantly ($p < 0.01$; Table 1) higher than in the east. A clear west to east effect in the reduction in shrubs is demonstrated. In the ungrazed situation of Lange Strooken and Vinkenduin the decline starts in 2001; in the other four areas the decrease started already in 1990. Aerial photographs show an overall regressive succession as when Hippophae shrubs start to die, the shrubs open up and gradually shift to grassland.

In all years and areas, the proportion of shrub varied between 30 to 42% of the area (Fig. 3a, b; online appendix Table 1a, b). In all six areas, the dominant shrub is *Hippophae rhamnoides*. The main changes in coverage are mainly caused by the growth and decline of this species. Between 1975 and 1990, the proportions of shrub increased from 35.2 to 40.7% (grazed) and from 37 to 41.9% (ungrazed). Aerial photographs show an expansion of closed thickets of Hippophae shrubs. From 1990 to 2001, this tendency is followed by a decline from 40.7 to 30.5% (grazed) and from 41.9 to 37.8% (ungrazed). From 2001 to 2009, in the grazed area, shrubland increases slightly from 30.5 to 32.0%, whereas in the ungrazed areas the shrubland further decreases from 37.8 to 32.2%.

### Changes in grasslands

In the ungrazed areas, the average change between 1975 and 1990 amounts to 5 ha (bare sand changes mainly into grasslands); in the grazed area, almost no changes are observed (Fig. 2c; Table 1). From 1990 to 2001, there is an increase...
of about 5 ha in both the grazed and ungrazed areas. Between 2001 and 2009, the extent of grasslands stabilize but there is a difference by location. The further east, there is a reduced change (p < 0.04) to grassland.

The loss of grasslands by the increasing share of bare sand in the period 2001–2009 is explained by a larger decrease in shrubland. Grazing pressure and time variables do not explain the model variables.

In the mosaics of bare sand and shrubland, the dune grassland in all the years of observation and in all areas forms the largest portion of the three crisp classes, illustrating the relative openness of the coastal dune landscape. The dune grassland varies between 41 and 64% of the area (Fig. 3a, b; online appendix Table 1a, b). Between 1975 and 1990, the extent of grassland in the grazed area remains the same, while in the ungrazed areas the grassland proportion increased from 41.5 to 49.7%, but the changes are different. Between 1990 and 2001, both grazed and ungrazed areas of grassland show an increase of 13.3 and 9.8%, respectively. In the 2001–2009 period, the grassland slightly decreased with 4% in the grazed area. In the ungrazed situation, the grassland increased slightly by 2.1%.

In overall conclusion, the extent of grasslands is determined by the space left over from the changes in the development of both bare sand and shrubland.

**Discussion**

We analysed the effect of the introduction of livestock grazing on changes in the presence and changes in cover extent of the crisp classes of bare sand, grassland and shrubland in the coastal dunes of Meijendel. We hypothesized that due to grazing and trampling caused by the livestock, and the rejuvenating effects of blowouts, the areas of bare sand, and grassland would increase, while simultaneously shrub encroachment at least would be stopped, and even reduced.

Regressive succession occurred, but contrary to our hypotheses, no significant impact of the introduced livestock could be demonstrated in the observed changes of the extent of bare sand, grassland and shrubland with the grazing intensity applied. In the ungrazed areas, similar changes occurred as in the grazed. The question arises which factors could explain this unexpected outcome.

**Bare sand**

Which factors can explain the decrease in bare sand up to 2001 and its subsequent increase in all six areas irrespective of livestock grazing?

Bare sand in coastal dunes is normally present as blowouts and in paths. The decrease of bare sand from 1975 until 1990 can be linked to the compulsory planting of Marram grass all over Meijendel as in other dune areas (e.g. Van Dorp et al. 1985; Van Til 1999). Annually, in Meijendel up to 1.2 million Marram tufts were planted and in many places this was repeated, stimulating a shift towards stabilisation (Van der Meulen and Jungerius 1989; Mensing 2002). In 2001, bare sand mainly consisted of small spots and long paths, as well as in the grazed Helmduiten (Fig. 4a) as in the ungrazed Prinsenduin (Fig. 5a).

The paths originate from monitoring volunteers, rangers and (illegal) visitors, all taking the same routes over and over again. Though expected, introducing livestock in 1990 did not result in an extra increase of bare sand in the grazed areas. Grazed and ungrazed areas have a similar decrease (Fig. 2a) and shift (Fig. 4a, 5a). Between 2001 and 2009, there is a substantial and comparable increase of bare sand in the grazed Helmduiten (Fig. 4b) as well as in the ungrazed Prinsenduin (Fig. 5b). In 2009, the paths are still there, but the larger part of the new bare sand is formed by blowouts with areas much broader than paths. These new blowouts are mainly found on slopes with a south-west orientation with climate extremes in spring and summer. Former Marram grass planters confirm that the areas, which were regularly involved in Marram planting, opened up again as part of a natural process irrespective of livestock grazing. Whether the Marram planting hampered the 1990–2001 development of blowouts is impossible to conclude from our data.

Since 1990, the airborne nitrogen load in the dunes has almost halved (Noordijk 2007) and this is likely to have consequences on the increase of bare sand. Primary succession on bare sand starts with algal crusts (Van den Ancker et al. 1985; Pluis and De Winder 1989). Cyanobacteria are the initial colonizers mostly members of the genera Microcoleus, Oscillatoria and Tychonema, in more stable conditions, to be followed by the dominating green algae Klebsormidium flaccidum. These organisms all depend on airborne nitrogen (Pluis and De Winder 1989) and a substantial reduction must have had its implications on the colonization, growth and cohesion of these algal crusts.

A temperature jump in the average mean from 9.2 °C to 10.5 °C, starting in 1988 (KNMI 2013), and just before the introduction of livestock, is likely to have caused more droughts in the coastal sand dunes stimulating blowouts, irrespective of livestock grazing. Several spring droughts and dry winter winds from the East in the period 2001–2009 are also likely to have stimulated sand blowouts (Aggenbach et al. 2018). Model calculations based on changing climate conditions confirm this theory (Witte et al. 2008, 2012).

On the other hand, changes in climate conditions also led to more intense summer rains and a longer growth period that stimulates an increased and more palatable biomass production over a longer period during a year. Civerolo et al. (2008) found an interaction between a higher nitrogen input and climate change because of higher summer rainfall, making it difficult to pinpoint the changes in our data to any one factor.

This research shows that the location (west, central, east) does not explain the shifts in area of bare sand over time,
though the wind intensity is presumed to have a greater effect in the western part of the dunes (Arens et al. 2013). Depending on the presence of organic matter, dune sand is more sensitive to erosion by wind or by water. Yellow lime-rich sand, mainly in the western part of the dune, is without cohesion and is more easily transported away by wind. Further east, grey

![Fig. 4](image1.png)  
**Fig. 4** Major part of the grazed Helmduijen area showing the $25 \times 25$ cm pixels where the land cover changed into bare sand. (a) The changes into bare sand from 1990 to 2001 (red). They are predominantly linear structures in reality observed as paths. (b) The changes from 1990 to 2001 (red) combined with the changes from 2001 to 2009 (pink). The substantial increase in the 2001–2009 period is clearly visible, predominantly surface structures in the field observed as blowouts.

![Fig. 5](image2.png)  
**Fig. 5** Major part of the ungrazed Prinsenduin area showing the $25 \times 25$ cm pixels where the land cover changed into bare sand. (a) The changes into bare sand from 1990 to 2001 (red). They are predominantly linear structures in reality observed as paths. (b) The changes from 1990 to 2001 (red) combined with the changes from 2001 to 2009 (pink). The substantial increase in the 2001–2009 period is clearly visible, predominantly surface structures in the field observed as blowouts.
lime-poor top soils contains more organic matter. This sand resists wind erosion, but because of its water repellency, the top layer can be easily washed downslope and this happens especially with heavy rainfall after a period of dry weather. Once the top layer is eroded, wind erosion picks up the underlying yellow sand leading to blowouts (Jungerius and Van der Meulen 1988). Between 2001 and 2009, it is believed that both processes lead to a substantial and comparable increase of bare sand.

Apart from changing nitrogen loads and changing climate factors, very slow increasing numbers of rabbits since the outbreak of RVHD in 1989 might lead to more collapsing warrens in all areas resulting in more points where wind erosion processes could take hold.

**Shrubland**

Rabbit grazing is likely to have a great impact on the shifts in shrubland. Rabbits not only graze the grasslands, have warrens, which can collapse that opens up the land to wind erosion, but especially favour nutrient rich seedlings of shrub and tree species (Jungerius and Van der Meulen 1988; Drees and Olff 2001; Van Tongeren 2006). The increase of shrubland can be explained by sudden and steep decreases in rabbit densities due to two rabbit diseases, which became epidemic in Meijendel. These are Myxomatosis from around 1954 and RVHD from around 1989 (Drees and Olff 2001; Scheffer 2012). Myxomatosis is considered by site managers to be responsible for shrubland increase between 1975 and 1990 and for its unexpected decrease between 1990 and 2001 (Fig. 3a, b). *Hippophae rhamnoides* can grow in nutrient poor and sandy pioneer conditions due to bacterial nodules on its roots (Oremus 1982). In the absence of rabbit grazing in the years directly after 1954, many seedlings or young sprouts of shrub species, including Hippophae, survived. The young plants invaded the surrounding area through clonal spreading, forming male and female monocultures of Hippophae shrubs. The life span of this shrub is about 25–40 years (Zoon 1995). The degeneration process is initiated by nematodes causing malfunctioning of the root nodules (Oremus 1982; Maas et al. 1983; Zoon 1986, 1995), finally resulting in dieback of the plants (Westhoff and Van Oosten 1991). The massive expansion from 1954 would therefore logically lead to a massive decrease starting around 1985 (Fig. 2b; 3a, b). Hippophae shrubs typically return to grasslands (regressive succession: this study; see also Provoost et al. 2011; Cornelissen et al. 2014), but the fine scale analysis of 25 × 25 cm field size pixels reveals the degenerating process at an earlier stage and better than manual mapping. On the other hand, new blowouts can generate new opportunities for Hippophae especially in a low density of rabbits due to both diseases being still epidemic. Provoost et al. (2011) report that livestock grazing in *Calamagrostis epigejos* dominated grasslands also led to a renewal of Hippophae shrubs, but in the Meijendel situation, the decreases outweigh (local) increases. This degenerating process has also been noticed in other dune areas (Aggenbach et al. 2017). The process is the same in all six areas, but the changes in the extent of shrubland are related to the location and the area seems to be smallest in the eastern areas. *Hippophae rhamnoides* as a pioneer species, has a preference for lime rich sands, which are devoid of nematodes feeding on its root nodules. The chances of finding thesefavoured conditions diminish from west to east, leading to reduced Hippophae populations.

The shrub *Crataegus monogyna* and tree species *Betula pendula*, *B. pubescens* and *Quercus robur* also demonstrate similar rabbit-related expansions in Meijendel. In the mid 1970’s, tree ring research in Meijendel was done on *Crataegus* within a sea-to-inland transect (three kilometres west-east). About 80–85% of all *Crataegus* specimens originated from the first few years after the outbreak of myxomatosis in 1954 (Salman and Van der Meijden 1985). Due to the life span of *Crataegus* (200 years or more), this species might eventually dominate the dunes in future decades before a massive collapse in 150 years time. A relatively sudden decline of *Betula pubescens* in the dune valley Bierlap (30 ha) in Meijendel was considered to be related to a lowering of the water table due to the extraction of groundwater. This was not the case (Van der Meulen and Wanders 1985). Sequential vegetation mapping showed the real cause of its expansion and decline (Maasdam 1988). Aerial photographs of 1938 showed that Betula invaded most of the valley from west to east after the abandonment by the farmers around 1890. Later years (1962, 1975, 1980) showed that the degeneration of Betula forest also went from west to east through the valley. Most of the Betula trees had disappeared by 1985 (Maasdam 1988). Based on the nutrient poor soil of the Bierlap valley, these Betula trees had a life span of about 70–90 years (see Fitter and Peat 1994; Beck et al. 2016). Prunus spinosa in the riverine sand dunes of Junner Koeland (The Netherlands) shows the same life span process of increase after myxomatosis and its latter collapse (pers. comm. J. Bokdam; M. Gleichmann).

**Dune grasslands**

In traditional progressive succession schemes, grasslands are the outcome of stabilized sand overgrown by vascular plants and mosses after a period of algal crusts. This process takes several decennia (Mensing 2002; Arens et al. 2007, 2009). These grasslands are gradually invaded by shrub species, which expand, and with progressive succession end up as dune forest, reducing the grassland area. In Meijendel, the shrub encroachment is related to the sudden collapse of the rabbit population due to diseases, starting with the 1954 myxomatosis, offering sudden opportunities for seedlings and
young sprouts of Hippophae rhamnoides and other shrub and tree species. Over a period of 35 years, Hippophae shrub increased at the expense of the grassland area. Due to aging, the shrub collapses and grasslands re-emerge (regressive succession). Because of the now acidified soil, the species composition will differ in the grassland compared to before the Hippophae expansion.

On the other hand, because of the increase of bare sand since 2001, the extent of the grasslands decrease in all research areas irrespective of livestock grazing. This means that it is plausible that -at least in the Meijendel situation- the changes in the extent of space for the grasslands is dictated by the increase and decrease of bare sand and of shrubland; grasslands fill in any left-over space. The regressive succession from Hippophae shrubland into grassland has also been noticed in other areas (Aggenbach et al. 2017) and is likely to be a regular phenomenon along the lime-rich coastal sand dunes. But this also means that apart from bare sand taking a part of the total grassland share, the cutting of indigenous long living shrubs and trees (like Crataegus and Quercus) is unavoidable to maintain a large amount of the EC priority (Council of the European Communities 1992) grasslands habitat.

Acknowledgements First of all we thank Dunea, the dune drinking water company that manages the Meijendel dunes, for giving the opportunity for this study and financing it. The initiative and support of Georgette Lettz, at the start of the research head of the department of Customers & Nature of Dunea, is gratefully acknowledged. Many thanks to my fellow PhD Nils van Rooijen discussing the results and helping out on the manuscript. We thank the reviewers for their useful comments on the first draft of the manuscript. Christopher Briggs is gratefully acknowledged for improving the English for the revised manuscript.

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