PERSISTENCE OF SPATIAL VARIANCE AND SPATIAL PATTERN IN THE ABUNDANCE OF A SUBMERGED PLANT

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Abstract. Organisms usually benefit from heterogeneous conditions, but, by doing so, may reduce the degree of heterogeneity. The question therefore arises how heterogeneity is maintained. We investigated within-year spatiotemporal patterns in a monospecific stand of a submerged plant (fennel pondweed, Potamogeton pectinatus), with the novelty that we distinguished between different forms of heterogeneity: spatial variance (the frequency distribution of densities) and spatial pattern (the spatial distribution of densities). We repeatedly measured plant biomass that was affected by swan predation, winter mortality, and summer regrowth. Spatial variance was enhanced mostly by swan foraging, despite the fact that swans appear to exploit patches to the same threshold level. Spatial pattern, which had vanished after swan foraging, reestablished due to spatial pattern in winter mortality and was further enhanced by plant regrowth. We found that variance and pattern each have their own temporal dynamics and are maintained by different biological processes. We therefore advocate that it is pivotal to distinguish between variance and pattern in the study of spatial heterogeneity.

Key words: Cygnus columbianus bewickii; density dependence; herbivory; heterogeneity; macrophyte; optimal foraging behavior; plant-animal interactions; Potamogeton pectinatus; spatial scale; vegetation patterns.

INTRODUCTION

In recent years it has become increasingly clear that for many problems we cannot ignore the fact that the world is heterogeneous in space (Sparrow 1999). Spatial heterogeneity is by no means “unnecessary complexity” (Pickett and Cadenasso 1995); the general notion is that organisms benefit under heterogeneous conditions (Sparrow 1999), both on the individual and on the population level (Fortin et al. 2005, Fryxell et al. 2005, Klaassen et al. 2006a, b). Spatial heterogeneity in food abundance (i.e., patchiness), for example, provides a forager with an opportunity to bias foraging effort toward areas that are, on average, richer in resources (Valone and Brown 1989, Klaassen et al. 2006b). Small-scale patchiness allows a forager that strategically regulates time in patches as well as movement between patches to achieve a higher energy gain rate (Klaassen et al. 2006b), whereas patchiness at the landscape scale allows mobile foragers to compensate for local and temporal variability in resource abundance, facilitating long-term population persistence (Fryxell et al. 2005). Accordingly, for conservation of animal populations it is vital to understand how heterogeneity is maintained in the ecosystem. This problem is particularly interesting in study systems where heterogeneity does not simply reflect heterogeneity in habitat characteristics (so where the template is homogeneous; Bascompte and Sole 1998), thus where heterogeneity is the result of some form of self-organization (Rietkerk et al. 2002).

Spatial heterogeneity consists of spatial variance, i.e., the frequency distribution of densities, and spatial pattern, i.e., the spatial distribution of densities (Li and Reynolds 1995, Wiens 2000). Variance and pattern are linked by the fact that pattern cannot exist without variance, but on the contrary there can be variance without pattern (i.e., a random spatial configuration; Tenhumberg et al. 2001). Obviously, variance and pattern can only persist if supporting effects are stronger than counteracting effects (Hutchings 1997, Adler et al. 2001).
Here we study spatiotemporal patterns in the abundance of a submerged plant, fennel pondweed *Potamogeton pectinatus* (L.). This is a pseudo-annual macrophyte that survives the winter via belowground tubers (Van Wijk 1988, Santamaría and Rodríguez-Girone 2002). In autumn (September) the distribution of tubers is strongly heterogeneous (Klaassen et al. 2006), despite the fact that pondweed occurs in a near-monospecific stand and on a template that is relatively homogeneous regarding abiotic factors (see Appendix A). The marked spatial variance and spatial pattern in tuber abundance allows the migratory Bewick's Swan *Cygnus columbianus bewickii* (Yarrell), that relies on submerged plants at migratory stopovers, to achieve a higher energy gain rate (Klaassen et al. 2006). An enhanced energy intake rate at stopover sites is thought to be pivotal for the swans to successfully complete their migratory journeys (Hedenström and Alerstam 1998). Our study system is the last refueling station for the swans, before they arrive at the wintering grounds (Nolet et al. 2006b; Fig. 1). The dependency of the swans on spatial heterogeneity in their food distribution is especially interesting because foraging itself has an enormous impact on the abundance of plants (Adler et al. 2001, Klaassen et al. 2006a, Nolet et al. 2006b).

We are interested in how spatial variance and spatial pattern in the abundance of tubers develop throughout the year, and whether three important biological processes in the pondweed's life cycle act the same on variance and on pattern. These processes are (1) “plant growth” (the production of tubers in summer), (2) “swan foraging” (the feeding on tubers by swans in autumn), and (3) “winter mortality” (the decomposition of tubers in winter) (Jonzén et al. 2002, Nolet 2004; Fig. 2). We monitored the natural development of variance and pattern at a small spatial scale, equal to the foraging scale or grain of swans (Wiens 1989, Van Eerden et al. 1997, Klaassen et al. 2006b). We illustrate here how spatial variance and spatial pattern in pondweed tuber abundance are affected by the different biological processes, and thus how heterogeneity is maintained in this study system.

**Methods**

**Study site**

Data were collected in the Lauwersmeer, a shallow freshwater lake in The Netherlands (53°22′ N, 06°13′ E; Fig. 1). In parts <0.7 m deep (750 ha) fennel pondweed grows from April to August. Tubers are mainly formed in August (Santamaria 2002, Santamaria and Rodríguez-Gironés 2002). In October the lake is visited by 1500–7000 Bewick’s Swans (peak numbers; Beekman et al. 1991).

**Sampling**

Tubers were sampled by taking 40 cm deep cores (diameter 10 cm) and sieving the sediment through a 3-mm sieve. Tubers were dried at 70°C for 72 h and weighed individually to the nearest 0.001 g. Tuber abundance is expressed as dry mass biomass density (grams per square meter).

**Spatial variance**

Tuber biomass densities were recorded in 33 squares (1×1 m, 12 cores/m²), arranged in a regular 3×11 array (20 m between squares) in September (2001, 2002, 2003, 2004), November (2001, 2003, 2004), and March (2002,
In 2002 the majority of pondweed tubers did not sprout within the study area, possibly as a result of an exceptionally warm period in February 2002 (R. H. G. Klaassen and B. A. Nolet, personal observation), resulting in extremely low tuber densities in September 2002. Because no swans visited the area in October, we did not sample in November 2002. A distance of 20 m between squares ensures that biomass densities are independent (for none of the sampling occasions could a positive correlation be detected between the densities in squares that were 20 m apart \( R^2 = 0.00 - 0.07, P = 0.08 - 0.96, n = 10 \); see also Nolet and Mooij [2002]). For every sampling episode we calculated the average and the coefficient of variation (CV) of biomass density.

**Spatial pattern**

Tuber biomass densities were recorded in each 1 m \( \times \) 1 m square (12 cores/m\(^2\)) within a 10 m \( \times \) 10 m plot in September (2001, 2003), November (2001, 2003), and March (2002, 2003, 2004). For abovementioned reasons no sampling was conducted in September and November 2002. For different lag distances (i.e., distances between squares), the magnitude of spatial autocorrelation was quantified by Moran’s \( I \) index (Moran 1950, Sawada 1999). In order to summarize the degree of spatial pattern for a certain sampling occasion we summed the absolute Moran’s \( I \) indices for all lag distances (1–9 m; Adler et al. 2001). Spatial relationships were visualized by autocorrelograms, i.e., a plot of the Moran’s \( I \) index against lag distance (Adler et al. 2001).

**Abiotic factors**

All study plots were located on the northeastern shore of the same creek (400 m \( \times \) 1200 m; Fig. 1). Throughout this area the water depth (standardized against the regulated water table) varied from 26.5 to 46.5 cm. The granulometric composition of the sediment was described along a single axis (using principal components analysis; see Nolet et al. 2001), which correlated most strongly with the silt fraction (percentage soil particles < 63 \( \mu \)m; Nolet et al. 2001). Within the study area the silt fraction varied from 1.5 to 4.8 % (95% confidence limit), which equals only 6% of the variation in silt fraction observed in the whole creek. Additional information about the (low) variation in abiotic factors throughout the study area is presented in Appendix A.

**RESULTS**

Tuber biomass densities were highest just after the growing season, as a result of a strong increase due to plant growth (Fig. 3a). Tuber biomass production in the summer was constant despite different initial (March) densities (see Fig. 4), indicating that per capita pondweed growth is depressed at higher densities (Fig. 4a). Swan foraging and subsequently winter mortality strongly decreased tuber biomass density (Fig. 3a). The exploitation by swans is a function of initial (September) density (Fig. 4b). From the exploitation pattern we can infer that swans exploit patches down to a threshold density of 8.5–13.0 g/m\(^2\) (negative of regression intercept...
of $y$-axis) and that 88–96% (regression slope) of the tuber biomass is accessible for the swans (Nolet et al. 2006a). Winter loss is similarly a function of initial (November) density in which more biomass is lost from a higher initial biomass (Fig. 4c). Tuber biomass densities are measured with an error of $\sim 20\%$ (as determined in the field; Nolet and Mooij 2002), which may cause spurious correlations between initial density and density changes. However, we determined by simulation that the slopes of such spurious correlations are smaller than the 95% confidence interval of the observed slopes, except for the correlation between tuber
density in March and the amount of biomass produced in summer (plant growth; see Appendix B). Hence, the reported correlations are no artifacts.

The relative variance in tuber abundance was highest just after the winter. The growth of plants strongly reduced spatial variance (Fig. 3b), as might be expected for a process that is slowed down at higher densities. On the contrary, swan foraging in particular and, to a lesser extent, winter mortality increased the level of variance. (The spatial variance within the 10 × 10 m plot, which was used to quantify spatial pattern, changed in similar ways [Fig. 3b].) Both in March and September a clear spatial pattern exists: high densities neighbor with high densities and low densities neighbor with low densities (Fig. 5a–c). This is also reflected in autocorrelograms where the degree of autocorrelation is higher for the first lag distances (Fig. 5d–f). The Moran’s $I$ index differed significantly from 0 for lag distances 1–2 m in March 2002, 1–5 m in March 2003, 1–6 m in March 2004, 1–3 m in October 2001, and 1–6 m in October 2003. In November, just after the feeding activities of the swans, the spatial pattern has vanished. For none of the lag distances did Moran’s $I$ index differ significantly from 0 for both November 2001 and November 2003. We have thus observed that plant growth decreased the degree of variance, whereas it increased the degree of pattern. Winter mortality enhanced both variance and pattern, demonstrating that effects on variance and pattern are not always inversely related. Similar temporal patterns in spatial variance and spatial pattern emerged when tuber abundance was expressed in numbers rather than biomass densities (results not
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Spatial variance and spatial pattern each have their own temporal dynamics, with different processes sometimes having different effects on variance than on pattern. For example, swan grazing strongly increased small-scale spatial variance, whereas it strongly reduced the small-scale spatial pattern. In this study system spatial variance seems to be maintained mainly by the feeding activities of the swans, whereas spatial pattern seems to be maintained by winter mortality (and to a lesser extent plant growth).

The direction of the effect of a process on the degree of variance and pattern is not straightforward but depends on specific circumstances. We might, for example, expect that spatial variance decreases if a forager depletes patches down to a fixed threshold level (as swans apparently do, cf. Klaassen et al. 2006a, Nolet et al. 2006a, b). In contrast, we observed that swan foraging strongly increased the level of variance, which is explained by the swans giving-up foraging (and leaving the pondweed bed) before all patches are exploited (Nolet et al. 2006c). Our studies were conducted at a migratory stopover site where the Bewick’s Swans maximize their daily energy intake during their relatively short visit (Nolet and Klaassen 2005). At wintering sites swans probably do not maximize their intake rate but satisfy their daily energy requirements, and thus deplete patches down to a much lower threshold (Van Eerden et al. 1997). The effect on the dynamics of spatial variance may however depend on the swans’ foraging mode, i.e., whether they use a flexible or fixed patch leaving threshold (Nolet et al. 2006c). A general rule for the effects of a process on variance and pattern remains to be established (Adler et al. 2001), if any such generalization can be made.

In contrast to spatial variance, spatial pattern had disappeared after the foraging activities of the swans, but had reestablished after the winter. Parasitic infections are thought to be an important cause for winter mortality. A limited outreach of infections could actually produce an aggregated spatial pattern, as has, for example, been shown in an insect host–parasitoid system (Maron and Harrison 1997). The exact mechanism of how spatial pattern can reoccur during the winter period remains to be established.

For research on the movement behavior of Bewick’s Swans we have also measured tuber abundance at the same study site and in the same years in eight 10 × 10 m plots (24 cores/plot; for details see Klaassen et al. 2006a). A lower level of spatial variance was observed at this scale (for example, variance in September was 0.13 and 0.06 for small and large plots, respectively), corroborating a negative scaling relationship between the scale of measurement and the observed level of spatial variance (Wiens 1989, Levin 1992). More interesting, spatial variance had decreased over the winter for these large scale plots (data not shown), which is opposite to the change for small-scale plots. There is thus some indication that the direction of an effect is scale dependent. Consequently, it is pivotal to conduct a study at the right spatial scale. It is a challenge to explain the persistence of spatial variance and spatial pattern at multiple spatial scales.

The term “spatial heterogeneity” has been used for both spatial variance and spatial pattern, causing confusion about what heterogeneity precisely implies (Wiens 2000, Adler et al. 2001). Furthermore many studies are not explicit in whether they study variance or pattern, despite the fact that these are very different forms of spatial heterogeneity. In this study we have shown that spatial variance and spatial pattern are different and independent aspects of a heterogeneous environment that have their own temporal dynamics. Whether a single process is really responsible for maintaining variance or pattern can only be assessed by exclusion studies. In any case, we advocate that it is important to make a distinction between variance and pattern to be able to ultimately understand spatial heterogeneity in a system.

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**APPENDIX A**

A figure showing that study plots are relatively homogeneous in sediment characteristics (Ecological Archives E089-170-A1).

**APPENDIX B**

Estimating the slope for spurious correlations between biomass changes and initial biomass density (Ecological Archives E089-170-A2).