Phosphorus Translocation by Red Deer on a Subalpine Grassland in the Central European Alps

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ABSTRACT
We examined the role of red deer (Cervus elaphus L.) in translocating phosphorus (P) from their preferred grazing sites (short-grass vegetation on subalpine grasslands) to their wider home range in a subalpine grassland ecosystem in the Central European Alps. Phosphorus was used because it is the limiting nutrient in these grasslands. When we compared P removal of aboveground biomass due to grazing with P input due to the deposit of feces on a grid of 268 cells (20 m × 20 m) covering the entire grassland, we detected distinct spatial patterns: the proportion of heavily grazed short-grass vegetation increased with increasing soil-P pool, suggesting that red deer preferably grazed on grid cells with a higher soil-P pool. Biomass consumption related to increased proportion of short-grass vegetation, and therefore P removal, increased with increasing soil-P pool. However, within the two vegetation types (short-grass and tall-grass), consumption was independent from soil-P pool. In addition, P input rates from defecation increased with increasing soil-P pool, resulting in a constant mean net P loss of 0.083 kg ha⁻¹ y⁻¹ (0.03%–0.07% of soil-P pool) independent of both soil-P pool and vegetation type. Thus, there was no P translocation between grid cells with different soil-P pools or between short-grass and tall-grass vegetation. Based on these results, it is likely that the net rate of P loss is too small to explain the observed changes in vegetation composition from tall-herb/meadow communities to short-grass and from tall-grass to short-grass on the grassland since 1917. Instead, we suggest that the grazing patterns of red deer directly induced succession from tall-herb/meadow communities to short-grass vegetation. Yet, it is also possible that long-term net soil-P losses indirectly drive plant succession from short-grass to tall-grass vegetation, because nutrient depletion could reduce grazing pressure in short-grass vegetation and enable the characteristic tall-grass species Carex sempervirens Vill. to establish.

Key words: Cervus elaphus; elimination pattern; grazing pattern; phosphorus removal/input; succession; Swiss National Park.

INTRODUCTION
There are many potential effects of large herbivores on vegetation. Apart from increasing or decreasing primary production and changing species compo-
sition, species richness, and the physical structure of the vegetation itself (Collins and others 1998; Gough and Grace 1998; Knapp and others 1999; Virtanen and others 2002), large mammalian grazers may also accelerate nutrient turnover (Detling 1988; McNaughton and others 1997; Frank and Evans 1997; Knapp and others 1999). Spatial patterns of nutrients can be altered by grazers such as sheep, horses, and rabbits, which feed over a wide area, but defecate in a small area (Edwards and Hollis 1982; Willo and others 2000). Such feeding behavior results in a gradual impoverishment of the wider grazing range but a continued enrichment of small areas within it. Bokdam (2001) found that the excreta of cattle was deposited at resting places that covered only 2.5% of their grazing range in a Dutch heathland, and that 75% of the heathland was still excreta-free after 10 years of grazing. In the European Alps, the traditional system of dairy farming may also promote such patterns. Nutrients accumulate around huts and stables, where cattle rest and are milked (Spatz 1980). In contrast, other large herbivores feed in small and nutrient-rich areas, but defecate in much larger areas (Putman 1986). Various studies have shown that female red deer prefer nutrient-rich grasslands for grazing (Charles and others 1977; Clutton-Brock and others 1987; Gordon 1989), and that nutrients are transferred from these small grazing sites into the wider home range (Schoenecker and others 2002).

We believe that such a change in nutrient transfer took place on subalpine grasslands in the Swiss National Park (SNP). Agricultural management ceased with the foundation of the park in 1914, when domestic livestock (cattle and sheep), which had grazed on the subalpine grasslands for several centuries, were removed from the park area. Braun-Blanquet and others (1931) reported that tall-herb/meadow communities dominated the vegetation around the abandoned stables and on former cattle resting places where high input of cattle excreta had enriched the soil nutrient concentrations. Where cattle predominantly grazed but did not rest, tall-grass pastures dominated by the evergreen sedge Carex sempervirens Vill. developed. Soon after the park’s establishment, locally extinct red deer remigrated into the area (Haller 2002), and the nutrient-enriched tall-herb/meadow communities of abandoned subalpine grasslands became preferred nocturnal grazing sites for hinds (Stüssi 1970).

The vegetation development since the foundation of the park is well documented by time-series data on vegetation structure and composition. These data were collected every 5–10 years on more than 150 permanent plots established on these subalpine grasslands as early as 1917 (Achermann and others 2000; Grämiger and Krüsi 2000; Wildi and Schütz 2000). Between the park’s establishment and 1960, tall-herb/meadow communities were completely replaced by short-grass pastures. This process was accompanied by significant changes in vegetation composition (Achermann and others 2000): tall-growing herb and grass species (for example, Aco-

\[ \text{nitum compactum} \text{ Rchb.}, \ \text{Chenopodium bonus-henricus} \text{ L.}, \ \text{Deschampsia caespitosa} \text{ (L.) B.}, \ \text{Trisetum flavescens} \text{ (L.) B.} \] were replaced by small-growing grasses such as Festuca rubra L. and Briza media L. Later in the century (1970/1980), short-grass areas in proximity to tall-grass pastures (former cattle-grazing areas) were invaded by Carex sempervirens. Both the changes from tall-herb/meadow communities to short-grass pastures and the development of short-grass to tall-grass pastures may have been driven by red deer nutrient translocations.

Focusing on phosphorus (P), we hypothesized that on subalpine grasslands in the SNP: (a) red deer hinds prefer to graze on P-rich sites, (b) preferential grazing will deplete these sites, and (c) P is translocated from preferred nighttime grazing sites (short-grass vegetation on subalpine grasslands) to rarely grazed tall-grass vegetation or to daytime ranges in the surrounding forests or alpine grasslands.

**Study Site**

The study was conducted in a subalpine grassland ecosystem (Alp Stabelchod) within the SNP. The park was founded in 1914 and is located in the southeastern part of Switzerland (46°40′N, 10°15′E). It occupies an area of approximately 170 km² with 85 km² covered by vegetation (subalpine/alpine grasslands and forests). The elevation ranges between 1,400 and 3,174 m.

Alp Stabelchod (10.7 ha) is located at an elevation of 1,950 m and has an uniform slope of 6° in a southerly direction. The parent material consists of mainly dolomite sediments. The average annual temperature is 0.2°C ± 0.76 (mean ± SD) and the mean precipitation is 925 mm ± 162 (recorded at the park’s weather station: Buffalora 1,977 m). The growing season is from early June to the end of September. The two vegetation types found on Alp Stabelchod today are easily recognizable. As a result of intensive grazing, the vegetation height of the short-grass type is approximately 2 cm; by contrast, the tall-grass type, which is dominated by Carex sempervirens tussocks, exceeds 20 cm in
height. Mountain pine (*Pinus montana* Miller) is the predominant stand-forming species in the surrounding forests (Risch and others 2003).

**Materials and Methods**

Plant and mineral soil sampling was conducted on a systematic grid of 268 cells (20 m × 20 m), which encompassed the entire grassland area of Alp Stabelchod (10.7 ha). We focused on P cycling, because P offers the following advantages over other nutrients: (a) P is the limiting nutrient in subalpine and alpine grassland ecosystems (Dietl 1994); (b) P is mainly excreted with dung and not with urine (Wu and others 2000), which makes it much easier to quantify nutrient return in excreta (compared for example, to nitrogen); and (c) soil-P is immobile and therefore leaching losses are low (Hilal and others 1973).

**Vegetation**

The proportion (%) of both short-grass and tall-grass vegetation was visually estimated in each grid cell (20 m × 20 m) in the summer of 1998. Short-grass was defined as vegetation that was grazed to approximately 2 cm vegetation height. Vegetation composition was sampled in July and August 1998 on a subplot (1 m × 1 m) located at the center of each grid cell using the method of Braun-Blanquet (1964). Names of plant species followed Hess and others (1984).

**Soil-Phosphorus Pool**

Five mineral soil cores (1.5-cm diameter) were taken to a depth of 20 cm at the edges and the center of each grid-cell subplot (1 m × 1 m) immediately after completing the floristic survey in the summer of 1998. The shallow soils prevented deeper soil sampling. The soil cores from each subplot were combined, dried to constant weight at 60°C, passed through a 2-mm sieve, and analyzed for organic P concentration with the Tecator Flow Injection Analyser System 5012 for organic P.

**Phosphorous Input by Feces**

The number of fecal pellet groups was counted in each grid cell (20 m × 20 m) in July 1997. Additionally, we cleared all feces from 46 systematically selected grid cells (every sixth grid cell) in early May 1998. New pellets were then collected from these cells monthly from late May until the end of September and dried to constant weight at 60°C. We compared the 1998 input of dung in the 46 grid cells with the corresponding numbers of fecal pellet groups counted in July 1997. Yearly input of dung into each of the 268 grid cells was estimated by using the resulting linear regression equation:

\[
y = 274.9x + 142.06
\]

where *y* is dung dry weight (g), and *x* is number of fecal pellet groups counted in July 1997; (*n* = 46, *R*² = 0.62, *P* < 0.001).

We determined the average feces P concentration on 28 randomly selected feces samples. Samples were fine-ground and analyzed with the Tecator Flow Injection Analyser System 5012 for organic P.

**Phosphorus Removal by Grazing**

Based on the vegetation survey conducted in 1998, we stratified Alp Stabelchod into short- and tall-grass grid cells. We found 22 pairs of short- and tall-grass grid cells on a soil-P concentration gradient from 144 to 275 mg P kg⁻¹ soil, which met the following criteria: (a) the difference in soil-P concentration between cell pairs did not exceed 3 mg P kg⁻¹, and (b) the difference in short-grass proportion exceeded 50%. Five additional grid cells with concentrations between 93 and 135 mg P kg⁻¹ were selected in the tall-grass vegetation only, because the short-grass community did not contain grid cells with concentrations lower than 144 mg P kg⁻¹.

Before red deer returned from their winter ranges located outside the park (immediately after snowmelt), we installed two grazing-proof wire baskets measuring 28 × 48 × 20 cm with a mesh size of 1.5 cm in the center of each selected grid cell (44 baskets in short-grass and 54 baskets in tall-grass) in early June 2001. As control plots, two similar-sized areas were additionally established in each grid cell on unprotected vegetation. In mid-September, plants were clipped to a height of 2 cm aboveground on all plots and oven-dried to constant weight at 60°C. The differences in biomass between the protected plots and unprotected controls corresponded to the amount of dry biomass (in g) consumed by red deer annually. To avoid
underestimation of both plant production and biomass consumption (see, for example, McNaughton and others 1996), we used dry weights from baskets with a single clipping in September for our calculations because the productivity of monthly clipped vegetation was lower (936 versus 945 kg ha\(^{-1}\)).

The P concentration in leaf tissue (Leaf-P) of grazed vegetation was determined by establishing additional baskets in each of the 49 grid cells in early June. Plants were clipped several times to a height of 2 cm above ground until mid-September, mimicking the grazing behavior of red deer. Plant biomass was collected separately for each plot in a paper bag and oven-dried to constant weight at 60°C. Samples were dry-ashed in a muffle furnace at 450°C for 6 h, leached with 2N HNO\(_3\), and filtered. Analyses were conducted via inductively coupled plasma (ICP) for total P within the plant material (Weetman and Wells 1990). Because leaf-P was significantly related to soil-P in short-grass but not in tall-grass vegetation (see Results), we used the following equation to estimate leaf-P and to calculate P removal as a function of soil-P for the short grass:

\[
y = 0.0018x + 0.5943
\]

where \(y\) is leaf P (g kg\(^{-1}\)), and \(x\) is soil-P (g kg\(^{-1}\)) \((n = 22, R^2 = 0.32, P = 0.006)\).

We then multiplied the proportion of short-grass in each grid cell with the mean P removal in the biomass of the short-grass stratum and added the proportion of tall-grass multiplied by the mean P removal of the tall-grass stratum to determine P removal per grid cell and year. Annual net P loss (P removal minus P input) was also calculated for each grid cell.

**Data Analysis**

Before analysis, all data on dry biomass and short-grass cover (%) were transformed using natural log and arcsin square root transformation, respectively, because they did not fulfill the normality and homogeneity criteria (Sokal and Rohlf 1995). Data on leaf tissue nutrient concentrations were not transformed, because they already met these criteria. We used linear regression analyses to test the relationships between the independent variable soil-P pool and the dependent variables short-grass proportion, dry biomass consumed, leaf-P, P removal, P input and net P loss (removal – input). The effect of the soil-P pool on the dependent variables dry biomass consumed, leaf P, and P removal was tested separately for both strata on the scale of individual baskets using one-way analysis of variance (ANOVA). One-way ANOVAs were also used to test whether short-grass proportion, dry biomass consumed, P removal, P input, and net P loss (removal – input) per grid cell (grassland scale) depended on the soil-P pool. We used two-way ANOVA to compare (a) biomass consumption in short-grass versus tall-grass vegetation with short and tall-grass as fixed factors and (b) P removal with P input per grid cell over both the soil-P pool gradient and the short-grass cover gradient with P removal and P input as fixed factors. We calculated mean P removal/input for grid cells with the same short-grass cover before analysis (n reduced from 268 to 40). We compared mean annual net P loss with the successional development of the vegetation to estimate whether net P loss is important in driving succession. We derived the succession stage of grid cells by comparing the vegetation composition of each grid cell with the vegetation composition of the long-term data from 59 permanent plots, as described in detail by Wildi and Schütz (2000). We described the relationship between soil-P pool and vegetation succession using a linear regression model.

**RESULTS**

**Spatial Patterns on Alp Stabelchod**

The organic P concentration in the mineral soil (soil-P) ranged from 0.072 to 0.322 g P kg\(^{-1}\) in the 268 grid cells. Based on an average bulk density of 0.547 g cm\(^{-3}\) (short-grass = 0.599 g cm\(^{-3}\); tall-grass = 0.515 g cm\(^{-3}\); \(P = 0.33\)), we estimated that soil-P pools ranged from 79 to 352 kg P ha\(^{-1}\) (Figure 1A). Soil-P pools generally were highest in the eastern part of Alp Stabelchod north and south of the cottage, whereas lowest values were found in the southwestern part adjacent to the forest (Figure 1A). The proportion of short-grass vegetation was highly correlated to soil-P pools \((n = 40, R^2 = 0.63, P < 0.001)\), (Figure 1C), with the highest short-grass cover (more than 95% per grid cell) in the eastern part (average soil-P pools of 251 kg P ha\(^{-1}\)) and no short-grass cover in the western part of Alp Stabelchod (average soil-P pools of 154 kg P ha\(^{-1}\)), (Figure 1B).

**Phosphorus Removal, Phosphorus Input, and Phosphorus Balance**

Our clipping experiment (individual baskets) showed that biomass consumption by red deer was significantly higher in the short-grass area compared to the tall-grass vegetation \((n = 22,\)
In short-grass vegetation, red deer consumed 945 kg dry biomass per hectare per year, or 85% of the annually produced 1,110 kg ha\(^{-1}\) total biomass. In contrast, only 438 kg ha\(^{-1}\) \(\times\) y\(^{-1}\), or 17% of the annually produced 2,537 kg ha\(^{-1}\) total biomass, was consumed in tall-grass vegetation. Consumption did not depend on soil-P pool in either vegetation type (short-grass: \(n = 44, R^2 = 0.06, P = 0.27\), tall-grass: \(n = 54, R^2 = 0.04, P = 0.31\), (Figure 2A), but leaf-P concentration increased with increasing soil-P pool in short-grass vegetation (\(n = 22, R^2 = 0.32, P = 0.006\), (Figure 2B), with values ranging from 0.88 g kg\(^{-1}\) (soil-P pool = 158 kg ha\(^{-1}\)) to 1.13 g kg\(^{-1}\) dry biomass (soil-P pool = 299 kg ha\(^{-1}\)). In contrast, we did not detect a relationship between leaf-P and soil-P pools in tall-grass vegetation (mean P concentration = 1.04 g kg\(^{-1}\) dry biomass, \(n = 27, R^2 = 0.12, P = 0.08\) (Figure 2B). Despite the strong correlation between short-grass leaf-P and soil-P pool, P removal (basket scale) was not correlated with soil-P pool in either vegetation type (mean P removal short-grass: 0.95 kg P ha\(^{-1}\) \(\times\) y\(^{-1}\), \(n = 44, R^2 = 0.07, P = 0.08\); tall-grass: 0.49 kg P ha\(^{-1}\) \(\times\) y\(^{-1}\), \(n = 54, R^2 = 0.06, P = 0.07\) (Figure 2C). At the grassland scale, high P removal rates were detected for the eastern part of the Alp (Figure 3A) due to a higher proportion of short-grass vegetation (\(n = 40, R^2 = 0.94, P < 0.001\), (Figure 3C). These grid cells had proportionally higher biomass consumption, which in turn was correlated to the soil-P pool (\(n = 268, R^2 = 0.47, P < 0.001\)) (Figure. 3D). Overall, P removal rates ranged from 0.08 to 0.95 kg P ha\(^{-1}\) \(\times\) y\(^{-1}\) in the grassland.

The quantity of dung was highly variable among the grid cells (0.14–16 kg dry weight). Based on an average fecal P concentration of 3.92 g P kg\(^{-1}\), we calculated P input rates ranging from 0.014 to 1.576 kg P ha\(^{-1}\) \(\times\) y\(^{-1}\) (Figure 3B). The spatial pattern of feces-P additions was similar to that of P removal through grazing (Figure 3B) and was positively related to short-grass cover (\(n = 40, R^2 = 0.57, P < 0.001\), (Figure 3C) and soil-P pool (\(n = 268, R^2 = 0.0.25, P < 0.001\), (Figure 3D).

Rates of P removal and P input increased with increasing short-grass cover and soil-P pool (Figure 3C, D). Removal rates ranged from 0.305 kg P ha\(^{-1}\) \(\times\) y\(^{-1}\) (soil-P pool = 79 kg ha\(^{-1}\)) to 0.988 kg P ha\(^{-1}\) \(\times\) y\(^{-1}\) (soil-P pool = 352 kg ha\(^{-1}\)), whereas input rates were between 0.207 and 0.916 kg P ha\(^{-1}\) \(\times\) y\(^{-1}\). We found a highly significant positive correlation between P removal through grazing
and P input by feces ($n = 268$, $R^2 = 0.39$, $P < 0.001$); for example, high P removal due to grazing in the eastern part of Alp Stabelchod was balanced by high fecal P inputs (Figure 3A, B).

Overall, P removal rates were significantly higher than P-input rates ($P < 0.001$), averaging a net P loss of 0.083 kg P ha$^{-1}$ y$^{-1}$ independent of soil-P pool ($n = 268$, $R^2 = 0.0007$, $P = 0.65$) and short-grass cover ($n = 40$, $R^2 = 0.0002$, $P = 0.92$) (Figure 4A, B). On the single grid cell scale, we found that the soil-P balance varied between losses of 0.74 kg P ha$^{-1}$ y$^{-1}$ and gains of 0.66 kg P ha$^{-1}$ y$^{-1}$. Grid cells crossed by hiking trails had higher P losses (216 g P ha$^{-1}$ y$^{-1}$) due to smaller P additions.
from feces (Figure 4C, D), compared with the average losses of 50 g P ha\(^{-1}\) y\(^{-1}\) calculated for undisturbed grid cells.

**Net Phosphorus-Loss and Succession**

The vegetation within the 268 grid cells on Alp Stabelchod represents succession stages from short-grass vegetation dominated by *Festuca rubra* L. (early stages) to tall-grass vegetation with a predominance of *Carex sempervires* Vill. (late stages). We found a negative correlation between succession stage and soil-P pool (\(n = 268, R^2 = 0.33, P < 0.001\)). Because we observed a constant P loss independent of soil-P pool, we described the correlation in a linear model (Figure 5). The model predicts a decrease in soil-P pool from 250 kg P ha\(^{-1}\) in the earliest succession stages to 112 kg P ha\(^{-1}\) in the latest stages. If we assume (a) an average net P loss rate of 0.083 kg P ha\(^{-1}\) y\(^{-1}\) (= increase from 0.03% yearly soil-P pool loss in earliest stage to 0.07% loss in latest stage) and (b) that there are no other P sinks or P sources existing, it would take 1,660 years for the soil-P pool found in the most P-rich parts of Alp Stabelchod today to be depleted to the current levels observed in the P-poor parts near the forest edge.

![Figure 4. Relationship between a net phosphorus (P) loss and soil-P pool, b net P loss and cover of short-grass vegetation. Spatial patterns of c net P loss (removal > input) and d net P gain (input > removal) on the subalpine grassland of Alp Stabelchod.](image)

![Figure 5. Pattern of the soil-phosphorus (P) pool in relation to the succession stage of the vegetation in 268 grid cells on Alp Stabelchod. A linear model was fitted. Low number of succession stage = early succession stage; high number = late succession stage.](image)
**DISCUSSION**

**Biomass Consumption**

The subalpine grassland ecosystem of Alp Stabelchod is characterized by distinct spatial patterns. Soils in the eastern part of the grassland around former stables are P-rich and gradually become impoverished toward the western part and the forest edges. Vegetation and soil patterns were found to be highly correlated: increasing proportion of heavily grazed short-grass vegetation and decreasing proportion of rarely grazed tall-grass vegetation were positively related to increasing soil-P. Consequently, the total amount of biomass consumed by red deer was higher in P-rich than P-poor areas. Also, other studies have found that nutrient-rich grasslands dominated by *Festuca* are the preferred grazing sites for red deer hinds (Charles and others 1977; Clutton-Brock and others 1987; Gordon 1989). Hinds select plant material that is low in fiber and high in nutrients (Moss and others 1981; Staines and others 1982; Welch and Scott 1995) to satisfy their high energy requirements during pregnancy and lactation (Georgii 1980). Because (a) red deer hinds prefer to graze on nutrient-rich sites, and (b) continuous grazing is known to maintain a high nutrient concentration in plants throughout the growing season (Cargill and Jefferies 1984; Iason and others 1986; Gauthier and others 1995; Fox and others 1998), female deer tend to feed on areas that have already been grazed earlier in the season (Clark and others 1995; Märki and others 2000). Our results agree with these findings.

**Phosphorus Balance**

We found that P-removal rates were higher from P-rich than from P-poor areas, because biomass consumption by red deer increased with increasing soil-P pool. However, P-input rates due to the deposit of feces were also higher where biomass consumption was higher. Neff (1968) and Charles and others (1977) reported that the feeding patterns of large herbivores, such as red deer, may be fairly well represented by feces patterns, thus supporting our results. In contrast to our hypothesis, net P loss on Alp Stabelchod was not restricted to P-enriched areas, which are preferably grazed by red deer, but was independent of grazing intensity, soil-P pool, and P concentration in leaf tissue.

Because we found higher P removal than P input rates on Alp Stabelchod (average annual net P loss, 0.083 kg P ha\(^{-1}\)), it seems that the soil-P enrichment caused by cattle is slowly being reversed by red deer. Under today’s grazing regime, we estimated that it would take 1,660 years for the soil-P pool in the most P-rich parts of Alp Stabelchod to be depleted to the levels observed in the P-poor parts near the forest edge. Overall, our yearly net P losses of 0.03–0.07% of the soil-P pool were comparable to results reported for vegetation communities grazed by elk in the Rocky Mountain National Park (Schoenecker and others 2002), where substantial reductions of nitrogen pools were observed in willow and aspen communities, while pine forests became enriched. For meadow or grassland/shrub communities, however, almost no changes in nitrogen pools were found—that is, losses were less than 2% over 50 years.

Although the estimated 1,660 years for soil-P depletion seems to represent a very slow process it is likely that the processes discussed are even slower, because we might have overestimated the annual net P loss in grid cells crossed by hiking trails. Park rangers often remove pellets during trail maintenance, which would result in a higher annual net P loss compared to undisturbed cells. Calculations based on the annual net P loss of undisturbed grid cells only indicated that it would take 2,770 years to remove the accumulated P from cattle grazing. Using this scenario, the net P loss would vary between 1% and 2.2% of the soil-P pool per 50 years.

We are aware that these are estimates and that the calculated rate of P-depletion could have been influenced by under- or overestimations of the variables measured in our study. For example, the method used for estimating aboveground productivity and biomass consumption in grazing ecosystems can produce considerable biases (for a detailed discussion, see McNaughton and others 1996), which in our case could lead to an under- or overestimation of net P losses. Additionally, our study encompassed several years of measurements, during which grazing patterns or the population size of red deer could have changed. However, based on annual records, we are confident that red deer population size stayed quite constant between 1997 and 2001 (Haller 2002). We therefore are convinced that our approach is accurate enough to gain a good understanding of functions and processes in this grassland ecosystem. However, a future experimental approach, in which grazing would be separated from P-translocation effects by establishing a combined grazing pressure and fecal pellet redistribution gradient, would be helpful to test our interpretations.
Effects of Net Phosphorus Loss on Succession

Interactions between soil fertility and vegetation development can be complex. An accumulation of P and an increase in P mineralization are generally found during primary succession (Frizano and others 2002), although these trends are much more variable during secondary succession. Increases (Johnson and others 2001) as well as decreases in inorganic P (Abadin and others 2002) have been reported.

Our estimated average P loss (0.83 g P ha\(^{-1}\) y\(^{-1}\)) is so small that it is not likely that P depletion has been the driving force behind the floristic changes observed in the heavily grazed parts of Alp Stabelchod during the past 60 years (Schütz and others 2003). Within a few decades, tall-herb/meadow communities lost dominance in the most P-rich parts of the grassland and were replaced by a Festuca rubra L. dominated short-grass pasture. Parallel to these changes in community structure, we found marked increases in the number of plant species in the P-rich parts, whereas no change in species richness was found in the P-poor, rarely grazed tall-grass communities (Schütz and others 2003). Additionally, plants with physiological and morphological adaptations to grazing became more abundant since abandonment of the grassland: (a) small-growing species (for example Carex capillaris L., C. verna Chaix, Prunella vulgaris L., Trichophorum pumilum (Vahl.) Schinz et Thellung, Viola rupestris F.W. Schmidt), (b) species with morphological (Carlina acaulis L., Cirsium acaule (L.) Scop.) or chemical protection (Ranunculus acer L.), and (c) annuals such as Euphrasia montana Jordan and Gentiana nivalis L. (Schütz and others 2003). Based on these indications, we therefore suggest that disturbance by grazing rather than P translocation by red deer was the major driver behind the observed vegetation development. Grazing by red deer reduced the dominance of the competitive tall-growing species found in tall-herb/meadow communities (Grime 1979; Palo and Robbins 1991; Olofsson 2001; Wohlgemuth and others 2002), enabling a larger array of life strategies to be competitive and species richness to increase.

We therefore conclude that vegetation changes from tall-herb/meadow communities to short-grass pastures were caused by grazing disturbance rather than by P translocation. Phosphorus translocation, however, will likely have a long-term effect on vegetation development because depletion will eventually lead to a reduction in grazing pressure. The reduced grazing pressure will in turn enable the establishment of tall-grass vegetation and finally forest stands. Our results indicate that cyclic succession within the grassland is unlikely to occur, because the P-rich short-grass vegetation is not depleted in favor of P-poor tall-grass vegetation. Thus, shifts in the grazing preference of red deer from short- to tall-grass vegetation are not likely.

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