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Shifting Rangeland Mineral Resource Limitations: Ecological Responses to Atmospheric Nitrogen Deposition

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Shifting rangeland mineral resource limitations : ecological responses to atmospheric Nitrogen deposition

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Key points : Deposition ofm fixed nitrogen (N) from the atmosphere into natural and managed systems is increasing worldwide, primarily as a result of fossil fuel combustion and agricultural fertilization practices . While historically most rangelands have been subjected to relatively low N deposition rates , \tilde{N} deposition is projected to increase in rangelands . This unintentional widescale fertilization has the potential to dramatically shift mineral resource limitations in rangeland ecosystems . In contrast to forested ecosystem response to N deposition , one of the most evident rangeland responses to N deposition is a shift in community composition , with declines in biodiversity and potentially shifts in the dominant functional groups . Evidence to date suggests that many other ecological processes , such as interactions with insect herbivores and carbon (C) and N cycling may in turn be affected by this shift in species composition . Increasing predictive capability of rangeland responses to N deposition is crucial for developing appropriate management strategies .

Key words : nitrogen deposition , biodiversity , nutrient limitations , global change

Introduction

Deposition of anthropogenically‐fixed nitrogen from the atmosphere onto plant and soil surfaces is profoundly changing N availability and ecological stoichiometry in ecosystems worldwide . Since N is typically a major limiting resource in temperate terrestrial systems , this change in nutrient availability may have major consequences for plants and soils in both natural and managed rangelands . Historically N deposition research has focused on forests and , to a lesser extent , mesic grasslands in Europe and North America due to the high rates and long‐term , chronic deposition in these areas . However , rates of deposition are currently increasing and the areas of impact expanding , with estimates of large increases in deposition over the next 100 years in rangeland areas including Asia , Africa , and South America(Lamarque et al . , 2005) . Understanding , and potentially mitigating , N deposition influences on rangelands requires development of predictive models of N enrichment on rangeland plant , community , and ecosystem properties and processes .

Anthropogenic activities have more than doubled the annual inputs of fixed N into the biosphere , and these inputs are expected to continue to increase in the coming decades (Vitousek et al., 1997; Lamarque et al., 2005). Some of this fixed N becomes volatilized from agricultural activities (primarily as NH_x) or emitted as a result of fossil fuel combustion (primarily as NO_y), and is later deposited from the atmosphere onto terrestrial systems (Lovett , 1994) . Deposited N can then become incorporated into the biota through root or microbial uptake of N deposited into the soil surface and via foliar uptake of gaseous and particulate N compounds (Lovett , 1994) . This deposited N may affect ecosystem properties , including fundamentally shifting the absolute and relative availability of nutrients (Vitousek et al . , 1997) .

Aber et al .(1998) developed a conceptual model overviewing the major influences of N deposition temperate forests . In this model , ecosystems at stage 0 are N limited , but chronic N deposition shifts systems to become progressively less N limited . By stage 3 systems are N saturated . Following N saturation , net primary production (NPP) and N mineralization decline , and N leaching losses are high . Aber et al . (1998) further suggest that land use history determines the initial degree of N limitation at the site , and that the forest stand type (deciduous or evergreen) dictates the rate at which sites progress through the stages and become N saturated . Predicting , and potentially mitigating , rangeland responses to N deposition requires a conceptual model similar to that developed by Aber et al . for forested systems . Considerable uncertainty remains in how rangeland response to N deposition will differ from the forest ecosystem responses described by the Aber et al . model . Here I present an overview on some of the potential effects of N deposition on rangeland plants , communities , and ecosystems , with particular emphasis on key ways in which rangeland responses may differ from forested ecosystems .

Plant mineral nutrition

At the individual level , Aber et al .(1998) suggest that N deposition leads to increases in foliar N concentration ; subsequent increases in processes such as photosynthetic rates and grow th are expected until negative effects of declining Ca :Al and Mg :N ratios dominate . Studies mesic rangeland plant responses to simulated and natural N deposition typically indicate strong responses to N for the first stages of N deposition , with generally positive responses to N additions in terms of leaf or shoot N , photosynthesis, and growth (e.g., Power et al., 1995; Kirkham, 2001; Throop, 2005). However, with the exception of the extremely long-running Park Grass experiment (Crawley et al., 2005), the short-term nature of most rangeland N fertilization studies makes results relevant only to the early stages of the Aber et al . model , while systems are still N limited . Long‐term , chronic N addition experiments are needed to understand shifts in plant mineral nutritional trajectories following system N saturation .

How might these patterns differ for drier rangelands in which water , rather than N , limits plant growth ? Studies to date suggest that dry rangeland plant responses to N are highly species-specific, with some species showing positive or neutral response to N additions, and others showing co-limitation by water and N (e g., Lajtha and Whitford, 1989; Drenovsky and Richards , 2004 ; James et al . , 2005) . Across a precipitation gradient , there was conflicting evidence for shifts from N to water limitation with decreasing precipitation , with responses dependent upon the indices considered (Hooper and Johnson , 1999) . The authors conclude that co-limitation affects plants at ecophysiological, community, and ecosystem levels, and that these influences are in turn affected by biogeochemical feedbacks in resource availability . Thus , a mechanistic understanding of individual plant-level responses to N deposition across varying precipitation regimes may require broader scale assessments of how N deposition-induced changes in larger scale (e.g. community and ecosystem) processes in turn affect mineral resource availability .

Community composition

In contrast to the ecosystem-focused Aber et al . model, some of the strongest impacts of N deposition in rangelands appear to be shifts in the composition of plant communities and interactions among organisms . Nitrogen deposition has been identified as the third main threat to biodiversity worldwide (behind changes in land use and climate) ; and grasslands , savannas , and Mediterranean systems may be among the most susceptible to N deposition as these systems are often limited , at least in part , by N (Sala et al . , 2000) . An analysis of functional group responses to N deposition across biomes suggests that rare species , perennials , N fixers , and native species may be most susceptible to local extinction following N enrichment (Suding et al . , 2005) . In rangelands , empirical evidence for decreased plant diversity in response to N deposition is particularly clear for European mesic grasslands and heathlands . A strong negative correlation between deposition load and species richness and cover has been documented in Great Britain (Stevens et al., 2006), and it has been estimated that a cumulative load of 714 kg N ha would cause species richness to decline by 25% (Stevens et al . , 2004) . Substantial declines in diversity may occur quite rapidly (within three years) with high rates of N additions , but would likely occur only after 10 or more years of chronic , low‐level additions characteristic of N deposition (Dise and Stevens , 2005) . Proposed driving mechanisms for species losses include shifts in plant competitive abilities with altered resource availability , with N deposition in California grasslands causing competitive exclusion of native forbs by invasive non-native grasses (Weiss, 1999) and losses of forb diversity (Zavaleta et al., 2003). In contrast, coastal sage scrub subject to N deposition transitions from dominance of shrubs to dominance of grasses (Egerton-Warburton et al . , 2001 ; Wood et al . , 2006) . In Minnesota , species richness declined rapidly with N fertilization , and led to a shift from C₄ warm-season grasses (with typically high N use efficiencies) to C₃ cool-season grasses (with typically low N use efficiencies) (Wedin and Tilman, 1996). Though there are many system-specific differences in particular groups that are lost, there is a clear common pattern of declining rangeland diversity with N inputs . This is in strong contrast to forested ecosystem responses where shifts in dominant canopy species are not commonly reported . However , given the lifespan of trees , changes in community composition of forested ecosystems would be expected to occur over much longer temporal periods . Given strong controls of community composition on many ecological processes, deposition-induced changes in plant community diversity will likely be a key variable for understanding and predicting rangeland responses to N deposition .

In contrast to mesic systems , N deposition into dry rangelands may have fewer consequences for community composition if plants are limited by soil moisture rather than N . However , relatively little is known about mineral nutrient‐induced shifts in species composition in arid and semi‐arid rangelands . In the northern Chihuahuan Desert , fertilization studies suggest that N deposition may cause increased grass cover and decreased legume abundance , as well as lead to a shift in the dominant grass species(Baez et al., 2007). In the same desert, N treatments shifted species composition, leading to a loss of C_4 summer annuals (Gutierrez and Whitford , 1987) . Species diversity in dry rangelands may be maintained in part by resource pulses (Chesson et al . , 2004) , suggesting that temporal relationships between precipitation and N deposition patterns could also affect community patterns . Additional N fertilization studies in dry rangeland systems are needed to develop an understanding and generalize about how , if at all , N deposition and precipitation patterns interact to affect plant communities .

In addition to driving changes in plant community composition due to alteration of mineral resource availability , evidence to date suggests that N deposition may affect relationships between plants and secondary stresses such as insect herbivory . These shifts may in turn influence community and ecosystem processes (Throop and Lerdau , 2004) . In mesic grasslands , simulated N deposition affects the survival or performance of insect herbivores (e.g., Power et al., 1998; Throop, 2005); these changes may affect herbivore populations (e.g., Haddad et al., 2000) and may ultimately affect plant community composition (Power et al ., 1998 ; Carroll et al ., 2003) . In the case of the dominant heathland shrub *Calluna vulgaris* in Britain, N depositioninduced increases in susceptibility to secondary stresses such as insect herbivory and winter frost injury appears to be a factor in community change from shrubland to grassland(Power et al . , 1998 ; Carroll et al . , 2003) . Similarly , simulated N deposition increased the fungal pathogen load for C⁴ grasses in a Minnesota grassland , apparently as a result of increased foliar N concentration (Mitchell et al., 2003). Conversely, N deposition-induced shifts in plant communities may also affect herbivore and pathogen diversity . In Sweden , declines in butterfly species richness appear to be due , at least in part , to changes in plant community composition driven by N deposition . Butterfly species most likely to be lost are those reliant on nutrient‐poor conditions , with greatest local extinctions occurring for those species in areas with greatest encroachment of woody plants into grasslands (Ockinger et al . , 2006) . Similarly , checkerspot butterfly populations in California have declined in areas where N

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deposition facilitates invasion by non-native grasses and competitive exclusion of butterfly host plants. This cascade of events is prevented with moderate grazing as cattle graze preferentially on grasses (Weiss , 1999) .

Ecosystem processes

Aber et al . (1998) focus their model on the ecosystem level , positing that net primary production (NPP) increases with initial N deposition , but begins to decline as the system becomes N saturated (stage 2) . Simulated N deposition typically increases NPP in mesic rangelands(Dukes et al . , 2005 ; Bassin et al . , 2007 ; Chung et al . , 2007) , but it is unclear whether sustained N deposition will eventually cause declines in NPP similar to those exhibited by forests . These declines occur as a result of deposition-induced losses of base cations , mobilization of aluminum , and subsequent nutrient imbalances and aluminum toxicity (Aber et al . , 1998) . Initial increases in NPP in response to N deposition may not occur in drier rangelands where water and N co‐limit production , although eventual declines in NPP could still occur if nutrient imbalances or aluminum toxicity occur . However, the response of NPP to N deposition in rangelands may be much more complex than in forested systems if depositioninduced shifts in community composition substantially affect NPP . Studies with manipulated plant community diversity patterns have found that simulated N deposition leads to greater stimulation of NPP in high diversity communities than low diversity communities (Reich et al . , 2001 ; Chung et al . , 2007) . Thus , there may be complex interactions between diversity and NPP whereby N deposition initially causes increases in NPP , but that these increases are offset by declines in plant diversity and by a subsequent dampening of the positive influence of N on NPP .

Mounting concerns over rising concentrations of atmospheric carbon dioxide , along with the possibility of economic incentives for carbon (C) sequestration , underscore the importance of understanding the extent to which N deposition influences system C sequestration capacity . If N deposition affects NPP , these changes could translate into altered C storage , although N deposition has less potential to strongly affect C sequestration in rangelands than forested systems because of minimal capacity in rangelands for C storage in woody tissue (Townsend et al., 1996). Once again, empirical evidence suggests that N depositioninduced changes in community composition may be an important driver of rangeland C sequestration response to N deposition . Soil organic C storage increased in a Minnesota grassland under simulated N deposition , but only under diversity treatments where the plant species present exhibited elevated foliar lignin concentration in response to N deposition(Dijkstra et al . , 2004) . Nitrogen deposition could potentially lead to large changes in rangeland C sequestration if deposition leads to increased woody plant biomass . However , although positive correlations between N deposition rates and woody plant expansion have been reported (Kochy and Wilson , 2001 , 2005) , conclusive drivers of woody plant expansion in rangelands remain elusive (Archer et al . , 1995) .

Ecosystem processes mediated by microbial activity may be strongly affected by N deposition , with Aber et al . (1998) proposing that N mineralization mirrors NPP with declines following N saturation , while nitrification and N leaching increase following N saturation . Indeed , simulated N deposition can increase N mineralization rates in rangelands (Morecroft et al . , 1994 ; West et al . , 2006) . However , as with foliar N and NPP , it is not clear whether chronic N deposition‐induced shifts in mineral resource availability will cause eventual declines in N mineralization in rangelands . Also in congruence with the Aber et al . model , simulated N deposition caused increased nitrification and leaching in a dry coastal dune grassland (ten Harkel et al . , 1998) and increased nitrification in a Mediterranean grassland (Barnard et al . , 2006) . These patterns may be the result of altered resource availability to microbes driving changes in metabolic activities and biomass , although evidence to date suggests that simulated N deposition in rangelands can also strongly affect microbial community composition and function(Bradley et al . , 2006 ; Chung et al . , 2007) . However , changes in these microbially‐mediated processes may also be indirectly affected by changes in plant species richness , with the positive response of microbial biomass to N additions positively related to species richness in a Minnesota grassland (Chung et al . , 2007) . Leaching of N from a Minnesota grassland was also enhanced with N additions , but once again these were affected by plant species richness , with greater losses from monoculture plots as compared to high diversity plots (Dijkstra et al . , 2007) . Predicting rangeland ecosystem responses to N deposition will thus require understanding not only how N enrichment directly affects microbially‐mediated processes , but also the nature of N deposition impacts on plant community composition, and how these changes in turn indirectly mediated ecosystem-level processes

Conclusions

Projected increases in the deposition of anthropogenically-fixed N into rangelands will drastically shift mineral resource availability in these systems . The majority of N deposition research to date on the impacts of atmospheric N deposition has taken place in forested ecosystems . As such , much uncertainty remains regarding rangeland response to chronic , low‐level N additions . This is particularly true in drier rangelands where water is typically a major limiting or co-limiting resource . Developing predictive models for rangeland response to N deposition will require long-term experimental work spanning from the physiological to the ecosystem level . One clearly apparent contrast from the conceptual model developed by Aber et al . (1998) for N deposition influences in forested ecosystems is that decreases in plant community diversity are a major consequence of N deposition in rangelands . Thus , N deposition in rangelands may have both direct consequences from altered mineral resource availability , as well as indirect consequences mediated by shifts in plant community composition . The sustainability , forage quality and quantity , and aesthetic value of rangelands may all be affected by N deposition . Indeed , a recent economic

analysis suggested that the net financial benefits for decreasing N deposition loads may be particularly high in grassland ecosystems (Wamelink et al . , 2007) . These benefits can only be clearly calculated , however , with a comprehensive understanding of rangeland responses to N deposition . Rangeland researchers must face the challenge of developing predictive models of N deposition impacts on rangeland plant , community , and ecosystem properties and processes . Characterizing these responses and making generalizations across sites that differ in dominant vegetation and precipitation regime is crucial for understanding , and potentially mitigating , the negative consequences of N deposition on rangelands .

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