Trophic Interactions,
Community Organization,
and the Kluane Ecosystem

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The conservation of biodiversity depends on functioning natural ecosystems and communities. There are various scientific reasons for conserving natural ecosystems. One is that we need to retain representative portions of the natural world as baseline controls for human impacts elsewhere. Another is that we need to preserve species that may later be of benefit to humans. To conserve natural systems, however, we have to understand how disturbances affect natural systems and whether those effects will cause irreversible changes. Alternatively, in human-dominated systems, we need to understand how changes will affect our own livelihood through the stability of the system.

Communities are highly complex, consisting of a large number of species that interact with other species. Studies of communities ask two main questions. First, how are they structured? That is, how many levels, such as plants, herbivores, predators, and top predators, are there? Second, how does the ecosystem work? For example, will the system be affected if we take out one of the species or several of the species? Alternatively, are some species more important to the viability of the community? These questions are relevant to the idea that systems have a large number of species that perform the same function and that there is a large amount of redundancy (Walker 1991). This idea implies that one can afford to lose species without radically altering the integrity and functioning of the community. Redundancy in the function of species in the community is a necessary safety net for that community, and, if we lose that redundancy (i.e., reduce the diversity of species), then we run the risk of a collapse in the system from some small external perturbation.

These questions are also relevant to systems that have already been subjected to changes such as eutrophication of lakes, fragmentation of forests, acid rain pollution of lakes and forests, introduction of exotic species, and creation of protected areas as islands within human ecosystems. Removal of the lower trophic levels, the plants, will always change a community and result in habitat loss. Nearly all human endeavor affects lower levels and, hence, alters habitats. However, when herbivores or predators are removed from a system the effects are less predictable and depend on which species are affected. Does the removal of species at higher trophic levels change the nature of the whole community? We know from studies of some ecosystems such as the marine intertidal and coastal systems of the Pacific Coast of North America that, when top predators (in this case sea otters) are removed from the system, the system changes radically from one type of community to another and stays that way until the top predators return (Estes et al. 1989, Wootton 1994a). Therefore, in this ecosystem the removal of a predator radically alters the community. Not all communities, however, may react in this way when a predator is removed.

Because communities are complex and contain large numbers of species, we cannot study every component and put it together like a jigsaw puzzle. Indeed, it is likely that the sum of the component parts (the dynamics of individual populations) will not tell us how the combination of species will interact with each other. We must therefore study the whole community as a functioning unit and try to find general rules for how communities work. It is unlikely that simple generalities will be appropriate for such complex systems, and we recognize that in reality communities will have complex structure and dynamics. However, we must start our research with a simple idea and then expand on that as we learn more from experimentation. In this chapter we discuss trophic-level theory and lay the theoretical groundwork for the experiments described in the rest of this book.

3.1 Trophic-level Theory

3.1.1 Community Structure

Community structure has been studied by comparing food webs (i.e., the connections between different species in natural history studies of communities) across the world. Over the past decades several hundred descriptive studies of food webs have been exhaustively analyzed (Cohen 1978, Pimm 1982, Briand and Cohen 1987, Paine 1988, Menge and Farrell 1989, Schoener 1989, Hall and Raffaelli 1991, Polis 1991, Schoenly et al. 1991). Despite the incomplete data, a number of general patterns in food webs can be seen (Lawton 1989). Ecologists have suggested rules for how many species one might expect, the types of interactions between species, and the number of trophic levels in different parts of the world.

The functioning of these communities—their dynamics—has also been studied by comparative methods and by perturbation experiments. These studies have focused on how one trophic level affects another or on the nature of the interactions between species in the community. Interactions between species can be either direct or indirect. Direct interactions are those in which there is a physical relationship between species, such as a predator eating its prey or direct interference between competitors. Indirect interactions along a food chain are those that are one step or more removed. Thus, predation is a direct interaction, but the effect of the prey population on the vegetation as a consequence of that predation one level above is an indirect interaction. Other indirect interactions involve exploitation competition, in which one predator affects the food supply of another predator, apparent competition, in which two prey species share the same predator, and indirect mutualism, in which a predator affects its prey, which in turn affects the competition between another prey and therefore its own predator (Wootton 1994b). Indirect effects can also arise when one species affects the interaction between two other species, sometimes by a change in behavior (Schmitt 1998). Questions asked at this level of generality involve how far perturbations travel as indirect linkages along the food chain. For example, if the main prey for a predator is removed, does the predator switch to a secondary prey, and how does this affect the secondary prey and its own competitors? Alternatively, if a predator is removed, how does this alter the competition between various prey species?

3.1.2 Control of Trophic-level Biomass

Bottom-up hypotheses assume that systems are regulated by nutrient flow from below (White 1978, 1984) because plants are essential to the levels above. Comparative studies have suggested that low soil nutrients and low productivity of plants result in fewer trophic levels, that higher productivity of plants results in more trophic levels, and that a gradient of increasing primary productivity (and more levels) can be traced from high to low lati-
Ecosystem Dynamics of the Boreal Forest

Trophic Interactions and Community Organization

TROPHIC INTERACTIONS

ECOSYSTEM DYNAMICS OF THE BOREAL FOREST

TROPHIC INTERACTIONS AND COMMUNITY ORGANIZATION

3.1.3 Case Studies in Mammalian Communities

Long-term studies have thrown some light on trophic-level dynamics. For example, 40 years of monitoring the large-mammal communities in the Serengeti, Tanzania, revealed the complexities of trophic-level interactions (Sinclair 1995, Mduma et al. 1999). Herbivorous mammals can be divided into migrants and residents. Migrant species, such as the wildebeest (Connochaetes taurinus), are regulated through their food supply of grasses (monocots). Although top predators (lions, hyenas) feed on wildebeest, they are not limited by this food source, but rather are limited by the supply of resident herbivores. In turn, top predators limit the density of resident herbivores as well as lower level predators such as cheetah (Acinonyx jubatus) and wild dogs (Lycaon pictus) (Laurenson 1995). Many of the resident herbivores feed exclusively on herbs (dicots) and determine their composition. Hence, there is a complex series of control pathways. The migrant herbivores are regulated by bottom-up flow from monocots. In contrast, the resident herbivores are regulated by top-down flow from the predators. These conclusions were reached through interpreting perturbations in the ecosystem that acted as seminatural experiments.

Yellowstone National Park, USA, is another system in which the mammalian commu-

such as drought, whereas consumers are less affected. Thus, consumers can regulate plants under stressful conditions (White 1984, Menge and Olsen 1990). The types of stress differ in these two views. The former involves adaptations to harsh environments and is long term, the latter addresses more short-term events. Both concepts are likely to be valid in different circumstances.

In aquatic systems there is evidence that indirect interactions become diluted the further along the food chain they occur. Thus, a combined top-down, bottom-up hypothesis for aquatic systems suggests that biomass is regulated from below by nutrient availability, but this effect is strongest at the plant level and becomes weaker at progressively higher levels. Equally at the top of the food web, bottom-down interactions are strong, but these effects weaken with every step down (McQueen et al. 1986, 1989, Pace and Funke 1991). Other mechanisms that reduce the efficiency of predators are interference and territoriality of predators and refuges for prey. These result in attenuation of indirect effects at lower links in the chain, and so bottom-up effects are seen at lower trophic levels (Powers 1984, Arditi and Ginsburg 1989, Arditi et al. 1991, Hanski 1991).

These concepts have all been expressed verbally and hence they are imprecise. When we discuss an effect of one trophic level on another as a result of a perturbation, we need to define what type of effect is being considered. Thus, there are immediate or "instantaneous" effects when a change in one trophic level affects another. These may be quite different from long-term consequences if the change persists and the system settles at some new equilibrium. For example, in the intertidal community on the Pacific Coast of North America, gulls feed on goose barnacles (Pollicipes spp.) and mussels (Mytilus). When gulls were excluded for 2 years from exclosure plots (Wootton 1994a), there was an increase in one prey species (Pollicipes) but a decrease in the other (Mytilus) as a result of space competition with Pollicipes. These are long-term results. The short-term result of gull removal, however, would have been an immediate increase in survival of both prey species. Thus, indirect effects can have complex results farther along the food chain and make them difficult to predict (Yodzis 1988, Wootton 1994b).

3.1.3 Case Studies in Mammalian Communities

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Ecosystem dynamics has been studied for several decades (e.g., Houston 1982, Keiter and Boyce 1991). This system was previously perturbed by the removal of the top predator, the wolf (*Canis lupus*) a century ago. As in the Serengeti, the dominant migrant herbivore, elk (*Cervus elaphus*), is regulated by food supply. Elk, in turn, determine the vegetation composition, both for the monocots and the dicots, including regenerating young trees such as aspen (Merrill et al. 1994, Singer et al. 1994). The recent reintroduction of wolves to the ecosystem will test whether this bottom-up control persists in the migrant herbivores or whether top-down control takes over in both migrant and resident mammalian herbivores.

The intertidal communities along the Pacific Coast of North America (Estes et al. 1989, Estes and Duggins 1995) show top-down control. Top predators such as sea otters (*Enhydra lutris*) determine the abundance of sea urchins (*Strongylocentrus* spp.), the dominant herbivores, and hence indirectly determine the macroalgae composition and other members of the community.

Conclusions from the intertidal system were obtained from controlled experiments. In contrast, trophic dynamics of terrestrial ecosystems have been difficult to study in controlled experiments because of the large scale and the long time needed to produce results. However, by using the small mammal community in the boreal forest of the Yukon at Kluane, we have been able to generate perturbation experiments to examine the direction of control between trophic levels. These experiments were designed to perturb each trophic level in turn. We approached the problem of predicting the effects of our perturbations in the boreal forest by considering only immediate or instantaneous effects. Then we started from first principles and considered all the possible outcomes from an experimental perturbation as the effects traveled along the food chain.

### 3.2 Interactions of Trophic Levels

We illustrate in figure 3.1, through a hypothetical example, the way biomass and productivity of trophic levels are related. Each box represents the biomass of a trophic level. The arrows and their numbers indicate the annual flow of biomass, or productivity. Material enters the system from outside through, for example, geological weathering, and exits the system through leaching. The remaining material cycles through the system from the soil to the predators, while each level contributes to the decomposers, which then returns material to the soil nutrients. The input and output rates to the boxes balance out.

Control of the biomass in the boxes can be either from above or below. Thus, an increase in the nutrients of the soil could result in both the flow rates and biomass increasing at each level, indicating a bottom-up control. Equally, one could envisage an increase in the flow rate of material from soil nutrients to plants and an increase in flow rate from plants to herbivores, resulting in a change in herbivore biomass but no change in plant biomass. This would occur by top-down control from herbivores to plants.

There are a number of ways we could measure the responses of one trophic level to changes in another. For example, we could measure growth rate in plants and survivorship in animals. However, to avoid confusion with interpretation, we have used the change in biomass because it can be applied to all trophic levels. We say that one trophic level limits another if changes in one trophic level are associated with changes in the biomass of the adjacent level. There are, of course, a variety of ways in which one could link species in a food web, but for simplicity, we assume direct linear interactions. We indicate these interactions in the following way:

\[ N \leftrightarrow V \leftrightarrow H \leftrightarrow P, \]

where *N* denotes nutrient concentration, *V* denotes vegetation biomass (or primary producers), *H* denotes herbivore biomass (primary consumers), and *P* denotes carnivore bio-

![Figure 3.1 Hypothetical flow diagram showing biomass and annual rates of production for nutrient pool (*N*), vegetation (*V*), herbivores (*H*), predators (*P*), and decomposers (*D*). The system has annual inputs to the soil by geochemical processes and outputs from decomposers through leaching.](image-url)
mass (secondary consumers). The arrows denote trophic-level effects, with a rightward arrow implying that an increase in resource (food) increases the rate of change of biomass of the adjacent consumer level. Similarly, a leftward arrow implies that an increase in consumer biomass decreases the rate of change of the adjacent prey biomass. A vertical arrow implies that a given trophic level has a density-dependent effect on its own rate of growth.

Some of the food web structures one could think of would be biologically implausible. For example, a terminal trophic level that has no negative feedback on its growth would grow infinitely. Accordingly, there must be an intraspecific feedback effect at the end of each plausible chain or else a link between the penultimate and terminal levels. For similar reasons, it would not be biologically plausible to have arrangements such as the following:

\[ N \rightarrow V \rightarrow H \rightarrow P \]

because there would be no negative feedback on the rate of growth in any of the trophic levels and so all would grow exponentially over time. Therefore, we assume that there must be some feedback on each level, and possible competition within the trophic levels (i.e., there is always a vertical arrow at each level implied in our models). The algebraic formulations for instantaneous change at each trophic level are illustrated in table 3.1 for three basic types of bottom-up, top-down and reciprocal control (Sinclair et al. 2000). There are, however, 27 biologically plausible models (table 3.2), all of which have negative feedbacks at every link in the web, either due to self-limited growth or an impact from the next highest level. All models comprise various combinations of the equations in table 3.1 (J. M. Fryxell, personal communication). The majority of the models are highly stable, each level equilibrating monotonically or with a few minor fluctuations. Figure 3.2 illustrates the time dynamics of three of the models. Figure 3.2a shows the stable, pure bottom-up model 1. Figure 3.2b shows the pure reciprocal model 27 with some damped fluctuation. Only three models show more marked fluctuation, and they all involve reciprocal interactions between plants and herbivores with predators imposing top-down control (models 7, 20, 22). Figure 3.2c illustrates model 20 involving double dilution effects (see table 3.2).

We have assumed in our equations the simplest (linear) density-dependent relationships. For illustrative purposes, consider interactions between plants (V) and herbivores (H). A left arrow implies that changes in herbivore density affects the rate of plant biomass change, near equilibrium, but not vice versa. Such a situation implies that herbivore growth is limited by something other than food, even though herbivores do consume plants. For example, hares feed on white spruce, and the higher the density of hares, the heavier the browsing impact on the trees. However, hare survival is little affected by spruce biomass. Similarly, a right arrow implies that changes in plant density affect the rate of change of herbivore biomass, near equilibrium, but not vice versa. Biologically this occurs when red squirrels feed on cones, hares eat senescent willow leaves, and carnivores rely on carrion. A double-headed horizontal arrow between plants and herbivores implies that herbivores respond to plant abundance, and plants, in turn, respond to herbivore abundance, forming a reciprocal relationship (Caughley 1976). Finally, a density-dependent form of intraspecific competition implies that each trophic level has its own intrinsic limits.

<table>
<thead>
<tr>
<th>Table 3.1 Minimal algebraic expressions for bottom-up, top-down, and reciprocal interactions between trophic levels.</th>
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<tbody>
<tr>
<td><strong>Bottom-up Interactions</strong></td>
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<tr>
<td>( \frac{dN}{dt} = r_0 - a_{00}N )</td>
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<tr>
<td>( \frac{dV}{dt} = a_{10}NV - a_{11}V^2 )</td>
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<tr>
<td>( \frac{dH}{dt} = a_{21}VH - a_{22}H^2 )</td>
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<tr>
<td>( \frac{dP}{dt} = a_{32}HP - a_{33}P^2 )</td>
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<tr>
<td><strong>Top-down Interactions</strong></td>
</tr>
<tr>
<td>( \frac{dN}{dt} = r_0 - a_{00}N - a_{01}NV )</td>
</tr>
<tr>
<td>( \frac{dV}{dt} = \eta V - a_{12}VH - a_{11}V^2 )</td>
</tr>
<tr>
<td>( \frac{dH}{dt} = r_2 H - a_{23}HP - a_{22}H^2 )</td>
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<tr>
<td>( \frac{dP}{dt} = r_3 P - a_{33}P^2 )</td>
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<tr>
<td><strong>Reciprocal Interactions</strong></td>
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<tr>
<td>( \frac{dN}{dt} = r_0 - a_{00}N - a_{01}NV )</td>
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<tr>
<td>( \frac{dV}{dt} = a_{10}NV - a_{12}VH - a_{11}V^2 )</td>
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<tr>
<td>( \frac{dH}{dt} = a_{21}VH - a_{23}HP - a_{22}H^2 )</td>
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<tr>
<td>( \frac{dP}{dt} = a_{32}HP - a_{33}P^2 )</td>
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\( r_i \) is the per capita rate of increase of trophic level \( i \); \( a_{ij} \) is the community matrix coefficient for trophic level \( j \) acting on trophic level \( i \). \( N \) = soil nutrient pool, \( V \) = plant biomass, \( H \) = herbivore biomass, \( P \) = predator biomass. All models use some combination of these equations.
Table 3.2 Models for instantaneous change at each trophic level.

<table>
<thead>
<tr>
<th>Model</th>
<th>Models</th>
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<tbody>
<tr>
<td>Largely Bottom-up</td>
<td>Pure bottom-up</td>
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<td></td>
<td>Bottom-up, nutrient reciprocal</td>
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<td></td>
<td>Bottom-up, vegetation reciprocal</td>
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<td></td>
<td>Bottom-up, herbivore reciprocal</td>
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<td>Largely Top-down</td>
<td>Pure top-down</td>
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<td></td>
<td>Top-down, nutrient reciprocal</td>
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<td></td>
<td>Top-down, vegetation reciprocal</td>
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<td></td>
<td>Top-down, herbivore reciprocal</td>
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<tr>
<td>Herbivore Dominant</td>
<td>Herbivore dominant</td>
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<td></td>
<td>Herbivore dominant, vegetation dilution</td>
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<tr>
<td></td>
<td>Herbivore dominant, nutrient reciprocal</td>
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<td>Herbivore dominant, predator reciprocal</td>
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<tr>
<td></td>
<td>Herbivore-vegetation co-dominant</td>
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<tr>
<td>Vegetation Dominant</td>
<td>Vegetation dominant</td>
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<td></td>
<td>Vegetation dominant, herbivore dilution</td>
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<td>Vegetation dominant, nutrient reciprocal</td>
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<td>Vegetation dominant, predator reciprocal</td>
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<td>Largely Dilution</td>
<td>Herbivore dilution</td>
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<td>Vegetation dilution</td>
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<td></td>
<td>Herbivore-vegetation joint dilution</td>
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<td>Largely Reciprocal</td>
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<td>Reciprocal, predator top-down</td>
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<td></td>
<td>Reciprocal, nutrient bottom-up</td>
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<td>Reciprocal, vegetation top-down</td>
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<td>Reciprocal, vegetation bottom-up</td>
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<td>Reciprocal, herbivore bottom-down</td>
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<td></td>
<td>Pure reciprocal</td>
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</tbody>
</table>

We assume self-limitation where there is no left arrow affecting a trophic level. A left arrow indicates that one level is changed by that above, a right arrow indicates that one level is changed by that below, and a two-way arrow indicates that two levels affect each other. "Dilution" means that a level is influenced by both lower and higher levels, whereas "dominant" implies that a level affects both lower and higher levels.

N = nutrient pool; V = vegetation biomass; H = herbivore biomass; P = predator biomass.

Figure 3.2 The trajectory through time over 1000 intervals of 3 of the algebraic models described in table 3.2. (a) Pure bottom-up model, (b) pure reciprocal model, and (c) double dilution model. The compartments show nutrient pool (N), vegetation (V), herbivores (H), and predators (P).
3.2.2 Top-down Models

Top-down models have an uninterrupted direction of control from the top level downward. Thus, predators control the herbivore biomass, and herbivores control plant biomass. They all predict that nutrients are not limiting and that community structure is determined by type of predator species and predation.

These models could be applicable to aquatic systems, particularly the more eutrophic lakes with many trophic levels. Larger fish species determine the presence or absence of small fish species, which in turn determine which zooplankton occur (Menge and Sutherland 1976, Estes et al. 1977, Simenstad et al. 1978, Paine 1980, Power 1990, Menge 1992, Carpenter and Kitchell 1993, Rosemund et al. 1993, Wootton 1994a).

3.2.3 Herbivore-dominated Models

In herbivore-dominated models the herbivore level controls both the predator and plant levels. Interactions between plants and nutrients can be either top-down or bottom-up. These would apply to systems where herbivores dominate. Generally, such systems are grasslands. In tropical savannas, such as the Serengeti, large grazing mammals structure the vegetation and determine predator biomass (McNaughton 1985, Semmartin and Oesterheld 1996). Predators do not limit the dominant herbivores (Sinclair 1975, Sinclair and Arcese 1995). The dominant effects of herbivores are also seen in temperate prairies (Huntly 1991) and in arctic and antarctic tundra (Leader-Williams 1988, Albion and Langvatn 1992). In many of these systems, the large mammalian herbivores become numerous and develop seasonal migrations.

3.2.4 Vegetation-dominated Models

Vegetation-dominated models predict that soil nutrients are determined by plant biomass and that herbivores are limited by food supply. Such models would apply to forests where trees dominate the biomass. In particular, tropical forests absorb most of the nutrients in the soil, leaving the soil low in nutrients (Connell 1978). At the same time, herbivores in these forests, largely insects, consume only a small fraction of the vegetation because plants are protected by structural and chemical defenses (Reichle et al. 1973, Sinclair 1975). Similar processes occur in temperate forests (Feeny 1968).

One model (13) has both vegetation and herbivores as codominants. This would predict high vegetation biomass, but its structure would be strongly influenced by herbivores because of its high palatability. Such special conditions are not obviously identified in nature, but may represent subarctic and alpine systems where vegetation and herbivores dominate (Oksanen et al. 1981, Batzli 1983, Keith 1983, Fretwell 1987, L. Oksanen 1990, Moen et al. 1993, Marquis and Whelan 1994, Krebs et al. 1995).

3.2.5 Dilution Models

Dilution models imply that both the lower and higher levels limit the intermediate trophic level. Thus, there are both bottom-up and top-down effects (figure 3.2c). Such a process may be seen in aquatic systems, where both nutrient content of lakes and the

The 27 possible ways trophic levels can interact directly with each other can be conveniently grouped by the dominant direction of control in the system and by the dominant trophic level. Although other groupings can be produced just as easily, we prefer the six groups described next and shown in table 3.2.

3.2.1 Bottom-up Models

Bottom-up models have an uninterrupted direction of control from lower to higher levels (figure 3.2a). They all predict that nutrient availability determines the biomass in the higher levels and thus assume that supply of nutrients is limiting.

Such models, therefore, would be most relevant in nutrient-poor environments where plants have difficulty obtaining nutrients. In terrestrial systems this would apply to polar tundra (L. Oksanen 1988, 1990), subarctic boreal forests with acidic soils (this study), and nutrient-poor sclerophyll forests in Australia. The soils of much of Australia are so poor that nutrient content of leaves in individual trees and the densities of mammalian folivores reflect the local nutrient condition of the soil the trees grow in (Braithwaite et al. 1983, Braithwaite 1996). In aquatic systems nutrients are low in high-latitude, cold oligotrophic lakes, in acidic marshes and peat bogs, and in warm open-ocean systems.
3.2.6 Reciprocal Models

Reciprocal models suggest that there are two-way interactions between most of the trophic levels. The last model (27; figure 3.2b) predicts reciprocal effects at all levels. These models, therefore, could apply to most ecosystems.

3.3 Experimental Perturbations in the Boreal Forest

We tested the models described above using experimental perturbations of the boreal forest food web. The models can be distinguished by unique sets of predictions produced from seven perturbations that systematically reduced or enhanced each trophic level. The predictions were concerned with the subsequent direction of change in the populations or biomass of other trophic levels. The main components of the system are described elsewhere, but in essence the plant level is characterized by herbaceous dicots, woody shrubs such as willow and birch, and white spruce. The herbivore level is dominated by the snowshoe hare, which exhibits a 10-year cycle of abundance, and by ground squirrels, red squirrels, and various vole species. The predator level is composed of carnivores such as lynx and coyote and various raptors, notably the great horned owl.

Predictions from the models on the changes in biomass are indicated in table 3.3 as an increase, decrease, or no change. However, where two perturbations are applied simultaneously, both acting in the same direction, then an additional prediction can be made on the magnitude of change relative to either single perturbation. The seven experiments are explained below. Details of how they were set up are outlined in chapter 4.

3.3.1 Experiment 1: Application of Fertilizer

Fertilizer was applied from the air to two 1-km² blocks of forest. This should increase the soil nutrient pool (N). Models that suggest that plants are responsive to the nutrient pool (i.e., those with a double arrow between N and V) predict increases in biomass of higher trophic levels to varying lengths of the food chain. Models with a left arrow predict that plants are not limited by nutrients and should not respond to fertilizer inputs.

3.3.2 Experiment 2: Addition of Rabbit Chow

Commercial rabbit chow was applied ad libitum to two areas. This food is eaten readily by hares and ground squirrels. The treatment has the effect of artificially increasing the food supply for herbivores independently of the natural food. Models that propose herbivores respond to food supply predict an increase in herbivores and a subsequent decrease in natural food. Some of these models also predict that predators should increase. The remaining models assume that herbivores are not limited by food supply and do not respond to food addition.
Table 3.3 (Continued)

<table>
<thead>
<tr>
<th>Model</th>
<th>Exp. 1</th>
<th>Exp. 2</th>
<th>Exp. 3</th>
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</tbody>
</table>

N = nutrient pool, V = vegetation biomass, H = herbivore biomass, P = predator biomass.

3.3.3 Experiment 3: Exclusion of Carnivores

Carnivores were excluded from a 1-km² area by wire fencing. This fence is permeable to hares and squirrels through small holes in the fence. We also covered a small part of this area (10 ha) with an overhead monofilament screen. This had the effect of partially deterring great horned owls from depredating animals below the screen. For our purposes the fence reduced the predator trophic level and allowed us to see how the effects traveled along the food chain.

3.3.4 Experiment 4: Exclusion of Carnivores and the Addition of Rabbit Chow

A similar 1-km² area was used for experiment 4. Carnivores were excluded and ad libitum rabbit chow provided. The removal of predators is predicted to increase herbivores. In addition, for those models where herbivores are responsive to their food supply, we predict an additional increase in herbivore biomass relative to that change from experiment 3. The remaining models predict a change in herbivores similar to that in experiment 3.

3.3.5 Experiment 5: Exclusion of Hares

Hares are the dominant herbivore biomass in the system, and they were excluded by fencing a 4-ha area. This has the effect of removing the top two trophic levels. The predictions are that vegetation biomass should increase and soil nutrients decrease.

3.3.6 Experiment 6: Exclusion of Hares and the Addition of Fertilizer

This experiment is similar to experiment 5 but with the addition of fertilizer as in experiment 1. The two perturbations are predicted to have opposite effects on soil nutrients: fertilizer adds nutrients, but removal of hares results in less nutrients. Because the outcome could be any value depending on absolute amounts of inputs and outputs, no qualitative prediction for soil nutrients can be made. However, for vegetation biomass one can predict two alternative responses. In models that assume that plants respond to nutrient levels, there should be a greater increase in plant biomass relative to experiment 5. In contrast, the other models predict no difference in plant biomass changes between experiments 5 and 6.

3.3.7 Experiment 7: Removal of Vegetation

In ten 1-m² plots on each of five sites, vegetation was killed by herbicide and left in situ. The perimeter was cut with a spade so as to kill roots from outside the plot. Soil nutrient levels were measured relative to an equal number of immediately adjacent control plots with intact vegetation. Soil nutrients should increase in plots where the vegetation has been killed.

In summary, 21 models make unique sets of predictions for these experiments. There are also three pairs of unique sets (5 and 8, 12 and 19, and 6 and 26). The three pairs of
models with the same predictions occur because one perturbation is missing from our boreal forest experiments (for logistical reasons): the direct addition of herbivores. If this experiment had been achieved, then all these models could have been discriminated.

3.4 Discussion

3.4.1 Limitations of the Models

We employed both removal and addition experiments. Removal experiments are more powerful because, to analyze the effect of a factor, it must be removed. Addition experiments are not the reverse of removals (Royama 1977), but they are necessary in this context to determine the presence of right-arrow regulatory effects. The double perturbation experiments where both addition and removal were applied allow us to evaluate the relative strengths of simultaneous top-down and bottom-up effects.

In general, the most informative experiments are those that influence higher rather than lower trophic levels. This is because perturbations at lower levels do not necessarily feed up—only some models predict this result. In contrast, perturbations on higher levels should always feed down to lower levels.

Our assumptions of the effect of food addition may be incorrect in two ways. One possibility is that rabbit chow could act more as an attractant, causing hares and other species to commute to the food from outside the experimental area. Such commuting would have the effect of increasing the herbivore trophic level directly through a behavioral response instead of through a reproductive response to food. A second possibility is that food addition would attract herbivores to eat more of the artificial food and so would decrease the impact of herbivores on the vegetation. In both cases predictions can be altered accordingly, and the number of unique sets does not alter.

We started this analysis of models and their predictions by assuming simple linear, first order interactions. Many other models, including nonlinear interactions, such as saturating functional responses, could also be suggested, and this would be the next step. We consider that these more complex models would lead to basically similar predictions. We started with experimental tests of simple models because they are easier to reject than complex models. We should first test simple models with our results before turning to complex models (Hairston and Hairston 1997). Density manipulation experiments in these different ecosystems is the most effective way of determining the generality of these top-down and bottom-up models (Dwyer 1995).

3.4.2 Models of Trophic-level Interactions in Different Ecosystems

Although we cannot yet assign individual models to particular ecosystems, we can speculate on the classes of models that may be applicable in different ecosystems. Nutrient-poor sclerophyll forests such as those of Australia have strong bottom-up effects (Braithwaite et al. 1983, Braithwaite 1996) and would be largely represented by right-arrow models such as 1–4, 16, 20–23, and 25. Tropical forests, where vegetation dominates (Connell 1978, 1983), could be represented by models 14–17, 21–22, and 24–25.

In subarctic tundra systems, where vegetation and herbivores dominate (Oksanen et al. 1981, Freiwald 1987, L. Oksanen 1990, Moen et al. 1993, Marquis and Whelan 1994, Krebs et al. 1995), models 9–17 could apply. As one moves to lower latitudes, such as in the temperate grasslands of the North American prairies (Huntly 1991) and in the tropical savannas of the Serengeti (Sinclair 1975, McNaughton 1985, Sinclair and Arcese 1995), herbivores dominate, and models with arrows leading from the herbivores could apply (e.g., models 9–13, 21, 23–24, and 26).

Both bottom-up and top-down (dilution) effects at the herbivore level could operate in insect herbivores (models 15–18, and 20) (Hairston et al. 1960, McQueen et al. 1986, Pace and Funkie 1991, Harrison and Cappuccino 1995). However, insect-dominated systems may also be controlled by predators or parasitoids (Lawton and Strong 1981, Strong et al. 1984, Price et al. 1990, Gomez and Zamora 1994, Spiller and Schoener 1994, Floyd 1996, Moran and Hurd 1998), and models 5–8, 22, 24, and 26 might apply.

In aquatic ecosystems, models with mainly left arrows (largely top-down effects; e.g., 5–8, 22, 24, and 26) may apply (Menge and Sutherland 1976, Estes et al. 1977, Simenstad et al. 1978, Paine 1980, McQueen et al. 1986, Power 1990, Menge 1992, Carpenter and Kitchell 1993, Rosemund et al. 1993, Wootton 1994a). Indirect effects are well known in aquatic and marine systems (Schoener 1993, Wootton 1993, 1994b, Menge 1997) but far less evident in terrestrial systems. The most general of the models is 27, the pure reciprocal model, because it could represent many if not all ecosystems.

3.5 Summary

Models of community organization involve variations of the top-down (predator control) or bottom-up (nutrient limitation) hypotheses. Verbal models, however, can be interpreted in different ways, leading to confusion. Therefore, we predict from first principles the range of possible trophic-level interactions and define mathematically the instantaneous effects of experimental perturbations. Some of these interactions are logically and biologically unfeasible. The remaining set of 27 feasible models is based on an initial assumption, for simplicity, of linear interactions between trophic levels. Many more complex and nonlinear models are logically feasible but, for parsimony, simple ones are tested first.

We have described a series of seven experiments designed to test the predictions of the models and distinguish between them. These experiments were conducted on the vertebrate community in the boreal forest at Kluane, Yukon. With these experiments we can distinguish 21 models and 3 pairs of models, giving 24 unique sets of predictions. In chapter 17 we apply these models to the results discussed in detail in the intervening chapters of this book.

Literature Cited


Menge, B. A. 1997. Detection of direct versus indirect effects: were experiments long enough? American Naturalist 149:801–823.


