Subcanopy gaps in temperate and tropical forests

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Abstract We present a model of gaps in the vertical structure of forest vegetation. The traditional model of a forest gap assumes the existence of a ‘hole’ in the uppermost canopy layer, often extending down to near the ground. The present model extends the concept to gaps at any level, including those in lower layers below an intact canopy or subcanopy. It assumes that gaps at any level represent spaces with unused resources, especially favourable for plant growth and survival. Evidence from temperate and tropical forests indicates that gaps in the subcanopy and understorey layers below intact canopies are common, and that plants have higher growth rates in them than in non-gap sites. We also extend this model to below-ground gaps in the root zone.

Key words: below-ground gap, canopy gap, forests, light regime, soil resources, subcanopy gap, understorey gap.

INTRODUCTION

Gaps in vegetation are local sites where additional resources, both above- and below-ground, have become available due to the absence of a resident plant. As a consequence, the increased resources can allow neighbours or new recruits located in and around the gap to grow, survive and reproduce better than those outside it. In addition, gaps represent ‘regeneration niches’ (Grubb 1977), to which some species are assumed to have adapted as ‘fugitives’ from competition. The importance of gaps was first recognized in terrestrial vegetation (Watt 1947; Cousens 1974; Platt 1975; Whitmore 1975), but studies in other habitats (e.g. assemblages of any sessile organisms, including invertebrates such as corals, bryozoans etc.) have also indicated their importance (Dayton 1971; Reed & Foster 1984; Connell & Keough 1985; Sousa 1985).

However, recently some authors have suggested that the simple gap/non-gap dichotomy may be less useful in understanding community dynamics than a more complex classification of forest structure. For example, Brokaw and Scheiner (1989) stated, ‘Results ... suggest the need to explore pervasive processes unrelated to discernible gaps’; while Connell (1989) stated, ‘the selection that takes place beneath the closed canopy has a great influence on the suite of species available to take advantage of the short burst of light in a gap’. Lieberman et al. (1989) stated, ‘considering forests as a Swiss cheese of gaps and non-gaps does not even begin to do justice to the daunting complexity of real forests’.

In this paper we attempt to deal with this complexity by expanding the traditional concept of gaps to include additional types not considered previously.

We propose the following general definition of a gap: ‘a space temporarily left unoccupied in an assemblage of sessile organisms’. We include the word ‘temporarily’ in the definition because gaps will almost certainly be filled eventually by the growth of new colonists and/or survivors within and bordering the gap, unless further disturbance interrupts this process. This expectation is based upon the fact that a living individual uses local resources for maintenance, growth and reproduction, so that its death, in whole or part, releases some resources that can be used by its neighbours or new colonists as they fill the gap.

In this paper we consider only gaps in terrestrial vegetation, principally multilayered temperate and tropical forests. Our general definition of a forest gap is broader than Brokaw’s (1982): ‘a “hole” in the forest extending through all levels down to an average height of two m above ground’; or Hubbell and Foster’s (1986) definition of a gap as an opening in the uppermost canopy extending down to any height. Our definition includes these, but introduces additional types of gaps in leaf layers at intermediate or lower heights, with or without an opening in the canopy above. We first discuss these subcanopy gaps in the above-ground vegetation, then go on to discuss some theoretical ideas about gaps in the root zone.
A MODEL OF GAPS IN ABOVE-GROUND VEGETATION

Consider a model forest with just three strata of leafy vegetation: canopy, subcanopy and understorey. In this model, there are eight possible combinations of the three leaf strata (Fig. 1). The definition of Brokaw (1982) recognizes only the complete gap with holes in all three strata (type 1; Fig. 1); that of Hubbell and Foster (1986) includes this type and adds two more: a gap in the canopy alone, and a gap extending down through the subcanopy (types 5 and 2; Fig. 1). Our model adds four more types, involving the lower two strata, singly or in combination (types 3, 4, 6 and 7; Fig. 1).

These different sorts of gaps could be created in at least two ways. First, the complete or partial death of one or more plants of any size could create any of these types. For example, the fall of a large canopy tree may smash all trees below it, creating a type 1 gap in all layers. However, if the same tree died standing, the loss of its leaves may create a type 5 gap in the canopy layer alone. Likewise, if a subcanopy or understorey tree died standing, this would create a type 6 or 7 gap, respectively. Second, the different types could occur at various stages during the filling of a type 1 gap. For example, such a gap could be filled from the bottom up, by the following sequence of gap types (shown in Fig. 1): type 1, then types 2, 5, 8, or other possible sequences. Alternatively, the gap could be filled in from the sides, if the crowns of canopy or subcanopy trees bordering the gap grew laterally into it; for example, the sequence could be: types 4, 7, 8, or types 3, 7, 8.

Fig. 1. Diagrams of the eight possible types of gaps in the above-ground vegetation in a model 3-layered forest. C, canopy layer, S, subcanopy layer, U, understorey layer.

EVIDENCE FOR THE IMPORTANCE OF SUBCANOPY GAPS

We will use the general term ‘subcanopy gaps’ for those below an intact canopy layer (i.e. types 4, 6, and 7; Fig. 1). Several studies suggest that they may be important in forest dynamics. First, Denslow et al. (1991) experimentally tested the hypothesis that, beneath a closed canopy, growth and survival of seedlings will be less if there is a layer of leaves closely overhanging them than without such a layer. Their results were consistent with this hypothesis; in particular, growing tips were killed more often with closely over-arching leaves.

Second, Brown and Parker (1994) measured the transmission of photosynthetically active radiation (PAR) in a series of 24 sites, in eastern deciduous forests in Maryland, that were in different stages of succession, ranging in age from 12 to more than 340 years. They suggested that the amount of PAR transmitted to 1 m height may be best explained by the proportion of the leaf layers that occur near the ground. This is consistent with the results of Denslow et al. (1991). Thus a gap in the understorey stratum, even with no gap in the strata above (i.e. type 7; Fig. 1), may provide as much light for seedlings as would gaps that occur only in one or more of the upper strata (types 2, 5 or 6).

Third, we have found that subcanopy gaps are usually the commonest among the seven different types of gaps in two rainforests. We estimated the proportion of these different gap types in permanent plots in a tropical and a subtropical rainforest in Queensland, Australia (see Connell et al. 1984 for a description of the sites). We estimated the height and thickness of all leaf layers above a series of points located at regular intervals along a series of transects that traversed the entire plots; 147 points were sampled in the tropical forest, 256 in the subtropical forest. We assigned the data into three strata, using the following height boundaries: canopy > 20 m, subcanopy 10–19.9 m, understorey 2–9.9 m. For each point, if no leaf layer occurred within a stratum, then it was scored as a gap in that stratum. To make the model comparable to Brokaw’s definition, we have omitted consideration of vegetation understorey 2–9.9 m. For each point, if no leaf layer occurred within a stratum, then it was scored as a gap in that stratum. To make the model comparable to Brokaw’s definition, we have omitted consideration of vegetation below 2 m in height and, for simplicity, restricted the number of strata to three layers.

An analysis of these data using the model in Fig. 1 showed that complete gaps (type 1) occupied 3% or less of the surface area in the tropical and subtropical sites (Table 1). Other types that included the canopy stratum (types 2, 3 and 5) occupied 25% or less of the areas of each site. Gaps beneath an intact canopy stratum, in the subcanopy and/or understorey (‘subcanopy gaps’, types 4, 6 and 7) occupied 38% and 48% of each site, respectively, while about a third of the area of each site had no gaps (type 8).

Fourth, Parker (1995) made the same sort of measurements of the heights of vegetation layers in a
temperate mixed deciduous forest at 39° latitude in the Appalachian mountains in the eastern United States. An analysis of these data using the model in Fig. 1 indicates that the proportions of the different types of gaps were very similar to those in the tropical forest we analysed in Queensland (Table 1). Thus ‘subcanopy gaps’ occupied the largest proportion of the area both in the two rainforests we studied and in the temperate forest studied by Parker (1995). In comparison, the classic ‘complete gap’ (type 1) was relatively rare in all three forests. Complete gaps are also rare in other rainforests; estimates usually range from 1 to 3% of the area (see review by Clark 1990).

The fifth bit of evidence comes from a study of the growth of seedlings and small saplings in the subtropical rainforest in Queensland described above. We

Table 1. Proportion of the surface area of forests comprised of different types of gaps

<table>
<thead>
<tr>
<th>Forest latitude</th>
<th>Complete gap in all 3 layers Type I</th>
<th>With gaps in only canopy and/or subcanopy layers Types 2,3,5</th>
<th>Subcanopy and understory gaps beneath an intact canopy layer Types 4,6,7</th>
<th>No gap Type 8</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tropical rainforest, 17°S</td>
<td>3</td>
<td>25</td>
<td>38</td>
<td>33</td>
</tr>
<tr>
<td>Subtropical rainforest, 28°S</td>
<td>2</td>
<td>14</td>
<td>48</td>
<td>36</td>
</tr>
<tr>
<td>Temperate deciduous, 39°N</td>
<td>3</td>
<td>26</td>
<td>43</td>
<td>28</td>
</tr>
</tbody>
</table>

See Fig. 1 for illustrations of gap types. Height limits of layers: canopy > 20 m, subcanopy 10–19.9 m, understory 2–9.9 m.

Table 2. Height of lowest leaf layer over saplings at a subtropical rainforest site

<table>
<thead>
<tr>
<th>Family</th>
<th>Species</th>
<th>Height to lowest leaf layer over sapling (m)</th>
<th>Fast-growing sapling</th>
<th>Slow-growing sapling</th>
</tr>
</thead>
<tbody>
<tr>
<td>Euphorbiaceae</td>
<td>Baloghia inophylla</td>
<td>3.4</td>
<td>0.3</td>
<td></td>
</tr>
<tr>
<td>Eupomatiaceae</td>
<td>Eupomatia laurina</td>
<td>2.7</td>
<td>7.9</td>
<td></td>
</tr>
<tr>
<td>Lauraceae</td>
<td>Cryptocarya obovata</td>
<td>9.8</td>
<td>4.0</td>
<td></td>
</tr>
<tr>
<td>Myrtaceae</td>
<td>Rhodomysurus psidioides</td>
<td>6.4</td>
<td>0.3</td>
<td></td>
</tr>
<tr>
<td>Proteaceae</td>
<td>Orites excelsa</td>
<td>17.7</td>
<td>0.5</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Stenocarpus salignus</td>
<td>14.6</td>
<td>1.5</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Stenocarpus sinatus</td>
<td>25.9</td>
<td>0.6</td>
<td></td>
</tr>
<tr>
<td>Rubiaceae</td>
<td>Randia benthamiana</td>
<td>14.6</td>
<td>0.9</td>
<td></td>
</tr>
<tr>
<td>Rutaceae</td>
<td>Acronychia pubescens</td>
<td>25.6</td>
<td>0.8</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>”</td>
<td>17.5</td>
<td>2.4</td>
</tr>
<tr>
<td></td>
<td></td>
<td>”</td>
<td>1.4</td>
<td>2.1</td>
</tr>
<tr>
<td></td>
<td></td>
<td>”</td>
<td>1.8</td>
<td>1.4</td>
</tr>
<tr>
<td></td>
<td></td>
<td>”</td>
<td>11.6</td>
<td>2.3</td>
</tr>
<tr>
<td>Sapindaceae</td>
<td>Ellatostachys nervosa</td>
<td>9.1</td>
<td>0.9</td>
<td></td>
</tr>
<tr>
<td></td>
<td>”</td>
<td>3.5</td>
<td>0.6</td>
<td></td>
</tr>
<tr>
<td></td>
<td>”</td>
<td>1.7</td>
<td>0.1</td>
<td></td>
</tr>
<tr>
<td></td>
<td>”</td>
<td>Guina semiglauca</td>
<td>11.3</td>
<td>0.6</td>
</tr>
<tr>
<td></td>
<td>”</td>
<td>Mischocarpus pyriformis</td>
<td>26.8</td>
<td>0.3</td>
</tr>
<tr>
<td></td>
<td>”</td>
<td>9.5</td>
<td>11.3</td>
<td></td>
</tr>
<tr>
<td></td>
<td>”</td>
<td>Sarcoperxy spilitata</td>
<td>6.9</td>
<td>4.0</td>
</tr>
<tr>
<td>Saxifragaceae</td>
<td>Quinlina sieberi</td>
<td>10.4</td>
<td>5.6</td>
<td></td>
</tr>
<tr>
<td></td>
<td>”</td>
<td>5.5</td>
<td>4.9</td>
<td></td>
</tr>
<tr>
<td>Sterculiaceae</td>
<td>Argyrodendron actinophyllum</td>
<td>8.4</td>
<td>1.1</td>
<td></td>
</tr>
<tr>
<td></td>
<td>”</td>
<td>4.0</td>
<td>1.5</td>
<td></td>
</tr>
<tr>
<td></td>
<td>”</td>
<td>1.4</td>
<td>0.9</td>
<td></td>
</tr>
<tr>
<td>Winteraceae</td>
<td>Tasmannia insipida</td>
<td>22.6</td>
<td>2.6</td>
<td></td>
</tr>
</tbody>
</table>

Mean 10.5 2.3
SE 1.6 0.5
N 26 26

Paired $t$-test

$P = 4.76$  
$P < 0.001$

*More herbivore damage on the slow-grower; bSlow-grower very close to a large tree.*
observed that, while most seedlings and saplings had grown very little over a 6 year period, some had grown appreciably; many of the fast-growers were not in 'complete gaps' as defined by Brokaw (1982). To discover the reasons for the fast growth outside complete gaps, we identified the 100 fastest-growing small saplings (0.3–2 m height at the start of the study), and in the field examined them for condition and local neighbourhood, recording proximity to larger trees, evidence of herbivory and distance to the nearest leaf layer above. As a control, we matched each fast-grower with a slow-growing conspecific sapling that: (i) had grown more slowly during the study interval (< 25% of the growth rate of the fast-grower), (ii) was about the same size at the start of the interval (within ± 25% of the height of the fast-grower), and (iii) was located nearby (< 30 m apart). As we were interested in growth outside complete gaps, we eliminated any sapling that had occurred in such a gap during the 6 year period of measurement. We found 26 pairs of saplings that satisfied all these criteria.

Our survey indicated that the fast-growers tended to be located in sites with a greater distance to the first leaf layer above than was the case for the slow-growing member of the matched pair. In 17 of the 26 pairs, the fast-grower had at least twice the open space above it as the slow-grower; this difference was significant (paired t-test; Table 2). The other variables noted were apparently less important: we found only two saplings (both slow-growers) that were very close to larger trees (Table 2); and instances of obvious herbivory occurred in only four pairs, always with more leaf damage in the slow-grower. In sum, the fast-growers tended to be in subcanopy and/or understory gaps (types 4, 6, or 7), with significantly more distance to the leaf layers above them than the slow-growers had.

We suggest that the mechanism producing this difference is the degree of obstruction of solar radiation arriving from various angles within the hemisphere of space centred on the seedling. Canham et al. (1990) have shown that beneath the canopy of multilayered forests in both tropical and temperate latitudes, most light arrives at the ground from low angles, further than 20° away from the zenith. A low layer of leaves overhanging a seedling at a short distance above it will occlude more of the light that arrives from these low angles, than would the same leaf layer situated further above the same seedling. Therefore a gap in the understory should allow more low-angle light to reach a seedling or small sapling than a gap in a site with an intact understory layer of leaves. Canham et al. (1990) noted that in a temperate Douglas fir–hemlock forest, a low subcanopy layer of hemlock probably occluded much of the light arriving from low angles.

These studies (Denslow et al. 1991; Brown & Parker 1994; Parker 1995; Canham et al. 1990; Tables 1, 2) suggest that (i) gaps in the subcanopy and/or understory layers of dense, multilayered forests (types 4, 6 or 7) are more common than gaps that include the upper canopy layers (types 2, 3 or 5; see Table 1), and (ii) subcanopy gaps that include the understory (types 4 or 7) probably allow as much or more light to reach seedlings and small saplings (which should promote their growth) than gaps that occur only in the canopy and/or subcanopy layers (types 2, 5 or 6).

**Below-ground Gaps**

We have concentrated on the effect of changes in above-ground vegetation on the availability of light to smaller plants. However, there is strong evidence that, as well as the above-ground portions, many roots die in gaps. This should increase availability of soil resources in these 'below-ground gaps', analogous to the increase in light in above-ground gaps. For example, the biomass of fine roots per unit volume was significantly lower in type 1 gaps than in shaded understory in a tropical rainforest (Sanford 1989, 1990) and in a temperate broad-leaved forest (Wilczynski & Pickett 1993). The soil water content in samples from 0 to 15 cm depth was also higher in type 1 gaps than in shaded understory in a tropical rainforest (Vitousek & Denslow 1986). Seedling recruitment, growth and survival have been enhanced in below-ground gaps produced experimentally by trenching, in temperate forests and tree plantations (Fricke 1904; Toumey & Kienholz 1931; Korstian & Coile 1938; Lutz 1945; Shirley 1945; Horn 1985; Cristy 1986) but less so in tropical rainforests (Connell 1971; Fox 1973).

Given this evidence, we hypothesize that below-ground gaps occur whenever a plant of any size dies, with the consequence that the soil resources formerly taken up by that plant become available to other plants in and near the gap. If this hypothesis is true, we suggest that the shorter plants (seedlings and saplings) in subcanopy and/or understory gaps should have access to additional soil water and nutrients, as has been demonstrated for complete gaps in the studies referred to above. The reasoning behind this suggestion is that because seedlings and small saplings have shallow roots, they will only have access to additional soil resources released by the death of shallow roots. As most trees, including those in the subcanopy and understory, have shallow (as well as deep) roots, the fine-root biomass at shallow soil depths should be reduced below a recently dead medium-sized or a small tree, as it would below a large recently dead tree. If so, seedlings or small saplings in subcanopy gaps (types 4, 6 and 7; Fig. 1), should have access to additional soil water and nutrients.

**Discussion**

Our aim in the models of above-ground vegetation (Fig. 1) is to focus attention on the role of gaps in the lower layers of forest vegetation, which usually have
been considered only when the layers above them were missing. We argue that when a tree of any size dies, the resources (light and soil nutrients and water) that it formerly used will then be available to its neighbours, which should respond and fill the gap.

Our results (Table 2), and those of Canham et al. (1990), Denslow et al. (1991), and Brown and Parker (1994), suggest that a small plant in the shaded understorey of a dense, multilayered forest will receive more light, and probably grow and survive better, the greater the amount of unobstructed space it has in the hemisphere of space above it.

In regard to below-ground gaps, previous studies have shown that the fine-root biomass at shallow soil depths is reduced in complete gaps (type 1; Fig. 1). We suggest that root biomass will also be reduced at shallow soil depths in subcanopy gaps beneath an intact canopy. Therefore, seedlings and small saplings, which have shallow roots, will have access to additional soil nutrients and water in these subcanopy gaps (types 4, 6 and 7; Fig. 1).

These ideas do not diminish the importance of the classic complete gap to the recruitment, growth and survival both of young stages of trees and of adults of subcanopy and understorey species. However, the proportion of area occupied by complete gaps is usually small, whereas subcanopy gaps, both in the rainforests studied here and in a temperate forest (Parker 1995), occupied a much larger proportion of the area. Therefore most seedlings, saplings and smaller understorey adults will spend the majority of their lives below an intact upper canopy layer, often in subcanopy and understorey gaps. Our aim here is to point out that, in addition to traditional complete gaps, the subcanopy gaps that exist beneath an intact upper canopy may also be important to the dynamics of broad-leaved multilayered forests, because of their relative frequency, and because the resources they release are of particular importance to the success of seedlings and saplings, which are the age classes that determine the future species composition and diversity of these forests.

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