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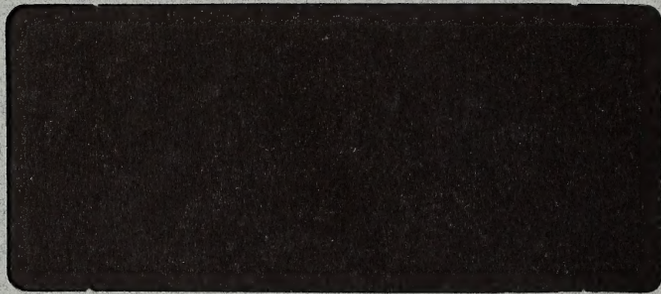
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Forest Development Research

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Manning Diversified Forest Products
Research Trust Fund
MDFP 9/97
Biodiversity in Harvested Areas in Relation to
Standing live Trees and Snags
Update 1997/98



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**Biodiversity in Harvested Areas in Relation to
Standing live Trees and Snags
Update 1997/98**

March 1998

**By Jim Schieck - Susan Crites - Lui Marinelli
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**Pub. No. T/417
ISBN: 07785-0238-4**

**Biodiversity In Harvested Areas
In Relation To The Dispersion Of
Standing Live Trees And Snags:
1997/98 Year-End Report**

Biodiversity In Harvested Areas In Relation To The Dispersion Of Standing Live Trees And Snags: 1997/98 Year-End Report

By

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Susan Crites
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Troy Sorensen
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1998

Vegreville, Alberta

DISCLAIMER

The study on which this report is based was funded by the Alberta Research Council, Millar Western Industries Ltd., Alberta Environmental Protection, Ainsworth Lumber Co. Ltd., Alberta Pacific Forest Industries Inc., and Manning Diversified Forest Products Research Trust Fund, which is a component of the Government of Alberta's Environmental Protection and Enhancement Fund. The views, statements and conclusions expressed and the recommendations made in this report are entirely those of the author(s) and should not be construed as the statements, conclusions, or opinions of members of the Manning Diversified Forest Products Research Trust Fund committee, the Government of Alberta, the Alberta Research Council, Millar Western Industries Ltd., Alberta Environmental Protection, Ainsworth Lumber Co. Ltd., Alberta Pacific Forest Industries Inc., or the Alberta Forest Research Advisory Council.

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EXECUTIVE SUMMARY

We are evaluating the extent to which biodiversity in harvested areas is affected by the size and number of tree clumps that are left standing after harvest. Twelve experimental cut-blocks were created in each of two study areas. One experimental area is in pine-dominated forests near Whitecourt, Alberta, and the other is in aspen-dominated forests near Grand Prairie, Alberta. Within each of the cut-blocks in the experimental areas, 3% of the pre-harvest merchantable trees and snags were retained in clumps. One third of the experimental cut-blocks had clumps of 0.60-0.75 ha (1.48-1.85 acres), one third had clumps of 0.14-0.15 ha (0.35-0.37 acres), and one third had clumps of 0.03-0.04 ha (0.07-0.10 acres). This research complements on-going and proposed research at the Alberta Research Council, the University of Alberta, the Network of Centers of Excellence, and forest industries throughout the Alberta by evaluating differences among dispersions of residual materials rather than evaluating differences among amounts of residual materials. All harvest patterns evaluated in this study can be implemented by forest companies that leave at least 3% of the canopy trees in some of their harvest blocks. The project started April 1997, with one year of pre-harvest biotic sampling and will continue for two years of post-harvest sampling. Although the primary focus of the project is on the biotic community, we will also evaluate the fall-down rates of trees and snags that are left standing in the experimental areas.

Within both study areas we grouped the experimental stands into four stand groups to increase the statistical power of the tests. Prior to harvest, composition and density of tree and snag species, volume of downed woody material, bird communities, and bat abundance varied among the habitat groups. However, after accounting for differences among habitat groups, forest structure and biotic communities did not vary among treatments. Thus, if we find differences after harvest they can be attributed directly to the experimental treatments. During 1998 and 1999 we will survey the forest structure and biotic communities in these experimental stands to evaluate the relative merits of leaving big versus small patches of standing trees within harvest areas.

Deliverables

As part of developing and conducting this research we have worked closely with government regulatory agencies and the forest industry to ensure that the results will be applicable to these organizations. We will continue to work with these organizations and present results (to groups and to individuals) as results become available. Specific deliverables include:

- 1) Progress reports submitted each year.
- 2) Descriptions of how the spatial pattern of standing trees and snags (i.e., variation in size and spacing of tree clumps) affects the vegetation, arthropod, bird, and bat communities within harvested areas.
- 3) Descriptions of the fall-down rates for trees and snags that are retained in harvested areas. Fall-down rates will be evaluated in relation to the size and placement of tree clumps in harvested areas.
- 4) Recommendations for clump size and clump spacing that are most suitable for meeting biodiversity goals.
- 5) Presentations of research findings in client reports, primary journals, and at conferences.
- 6) Participation in workshops, planning seminars, and committees that assess and regulate alternative harvest practices in Alberta.

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GENERAL ACKNOWLEDGEMENTS

This project has required and received strong commitments from government, academia, and the forest industry.

We appreciate the strong administrative and funding support provided by the Alberta Research Council. Special thanks are extended to Jack Nolan, Dave McNabb, Gerry Lofthaug, Duane Nakonechny, Pat Soldan, Elaine Cannan, Debby Franchuk, Sonia Hunka, Nancy Fraser Hrynyk, Susan Stecyk, Patricia Ferleyko, and Kathy Ilkiw for their assistance.

We thank our technical staff for their hard work in the field with data collection. These technicians include Laura Blonski, Erika Klausz, and Jodi Tomchyshyn. We acknowledge the editing and formatting of the document by Christine Gray. Special thanks to Len Peleshok and Kelly Sturgess for helping us get the field season started.

The following people and organizations helped with logistical support in the field, equipment storage, equipment supply, and providing a knowledge base of the area. Brian Telford, Dave Beck, Grant Williamson, Andrew Allison, and Larry LeFebvre of Ainsworth Lumber Co. in Grande Prairie; Richard Krygier, Jonathan Russell, Pat Rogers, Ray Hilts, and Colin Berg of Millar Western Industries in Whitecourt; Imperial Oil, Strathcona Refinery; Alberta Land and Forest Service, Whitecourt; and the Grande Prairie Regional College.

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CHAPTER 1. BACKGROUND AND RATIONALE

Jim Schieck

Background

As a consequence of having a thriving forest industry in Alberta, much of Alberta's boreal forest has been allocated to be harvested either for dimensional lumber or for pulp and paper. The economic benefits of this intensive forest harvesting are viewed positively by Alberta's population. The ecological ramifications of extensive forest harvest, however, are poorly understood and this lack of knowledge hampers management decisions and land use planning. Consequently, the forest industry and government organizations in Alberta have been conducting research to develop forest practices that are ecologically sound, economically feasible, and sustainable.

Short rotation practices will truncate forest succession over much of the landscape and, in some areas, may result in old forests becoming relatively scarce. In addition, the removal of most of the merchantable trees from harvested areas will alter natural floral and faunal communities in those areas and may affect biotic recovery. Finally, large live trees and snags provide suitable micro-habitats for many plants, and nesting and foraging habitats for many animals, but with conventional harvest practices, few large trees and snags will be present in harvested landscapes.

Management strategies are being developed to produce harvest patterns that are as similar as possible to natural disturbances. Since large trees and snags are abundant following natural disturbances, but are greatly reduced in harvested areas, forest managers have been recommending that as much standing residual material as is socially and economically feasible be retained within harvested areas. Clumps of standing trees and snags in harvested areas may function similar to skips in wildfire areas, and thus act as refugia for some species while the harvested forest re-grows. If residual standing live trees continue to grow, they will become large live trees and may produce some large snags in mid- and late-rotation forests. These

large live trees and snags may provide some of the structures and micro-habitats that are normally found only in post-rotation aged forests.

Given the complexity of natural ecosystems, it is not surprising that the ecological and biodiversity impacts of these alternative harvest/silviculture practices are poorly understood. Only by experimenting with new techniques in the field, and monitoring the resulting ecosystems throughout a harvest rotation, will we begin to understand the potential impacts and benefits of each technique. To this end, research has been conducted by scientists within the Alberta Environmental Protection, the University of Alberta, the Network of Centers of Excellence, and forest industries throughout Alberta's boreal forests. Preliminary results from past and on-going stand-level research indicates that biodiversity in harvested areas is positively related to the amount of standing trees and snags retained in those areas. However, only anecdotal information is available to evaluate whether the dispersion of those standing residual trees also affects the ecology and biodiversity in harvested areas. The present study will fill that gap in our knowledge by evaluating whether the biota present in harvested areas with small clumps of trees differs from that present in harvested areas with large clumps of trees. If clump size has a direct impact on biota, then by leaving trees and snags in the appropriate clump sizes, forest managers may obtain benefits to biodiversity at little additional cost. The proposed experimental treatments will be monitored throughout a harvest rotation (extended monitoring is not part of the present proposal) to enable researchers to evaluate changes in trees, understory vegetation and wildlife over time. This controlled experimental study will contribute to the adaptive management of boreal forests and will provide information that forest managers can use when designing harvest plans.

Scientific Rationale

All else being equal, it would be beneficial to make harvest areas as similar as possible to natural disturbances since native biota evolved in an ecosystem with natural disturbances (Urban et al. 1987). The dominant natural disturbance in the boreal forests of North America is wildfires (Eberhart and Woodard 1987, Hunter 1993), and cut-block size and shape can be patterned after wildfires. There are limitations, however, to matching the amount of wood remaining in post-harvest areas with that remaining post-wildfire because only a small percentage of the wood is combusted during a wildfire but most of the wood is removed during harvest (Lee et al. 1997). Given this discrepancy, it is necessary to manage the relatively few trees and snags that are retained in harvest areas to achieve the greatest benefit for biota. Following wildfires, there are groups of unburned trees and snags (fire skips) embedded within the burned areas (Eberhart and Woodard 1987). These fire skips could be emulated in harvest scenarios by leaving clumps of live trees and snags within cut-blocks. Both fire skips in burned areas, and residual clumps of trees in harvest areas, probably retain some of the structures and micro-habitats that were present in the forest prior to disturbance and consequently, may act as refugia for some native species. Small clumps may be sufficient for a few species, but larger clumps probably act as refugia for more species (Saunders et al. 1991). For example, some animals require large undisturbed areas for protection and foraging (Pomeluzi et al. 1993), and some plant species require large undisturbed areas which create suitable micro-habitats (Brothers and Spingarn 1992).

The refugia potential of residual clumps of trees is probably most important during the first few decades following harvest because many boreal forest species live at low densities in young forests (Schieck et al. 1995, Kirk et al. 1996). Thus, biota that are present in the residual clumps may disperse from these into harvest areas once the new cohort of trees is 20 to 50 years old. Distance between clumps and species dispersal abilities will affect the rate and dynamics of biotic recolonization in harvested

areas (Saunders et al. 1991, deMaynadier and Hunter 1995). The degree to which tree clumps act as refugia in harvested areas, and the appropriate dispersion of those clumps to enhance dispersal of biota, have not been evaluated in boreal forests.

In addition to the refugia potential of tree clumps, there may be management value to retaining small clumps or single live trees scattered throughout harvest areas. In areas where intensive forest harvest will decrease the amount of old forest in the landscape, it may be beneficial to produce old-forest structures and micro-habitats in mid seral stages of forests (Spies et al. 1991). Many of these old-forest structures and micro-habitats are associated with large live and dead trees (Hansen et al. 1991, Schieck et al. 1995). Consequently, if some merchantable trees are retained as standing live trees at harvest, and do not get blown down, they will become large by mid-rotation and may accelerate biotic recovery in the areas. With this management strategy, single trees or many small clumps of trees, scattered over the harvest area, may have greater impact on biota than if all standing trees and snags had been retained in few large clumps because old-forest structures and micro-habitats would then be dispersed throughout the harvest area.

The most appropriate placement of standing trees and snags differs between “managing for refugia” and “managing for the production of dispersed large trees”. Both patterns may be useful, but only by experimentally manipulating size of clumps and distance between clumps and then studying the biota throughout a rotation will we understand the biological and management implications of each. In this study, we established three treatments with different clump sizes and distances between clumps in two experimentally harvested areas to evaluate whether biota differ among the treatments. In the present study we will evaluate differences that are present immediately following harvest. The experimental areas will be maintained for the rest of the harvest rotation however, and proposals to evaluate differences among treatments will be submitted at the appropriate times (i.e., once the new cohorts of trees are

well established, at mid-rotation, and just prior to the subsequent harvest).

Objectives

In the present study, we will evaluate how clump size and distance between clumps of residual live trees and snags affects biodiversity in harvested areas. All treatment areas will be sampled before and after the experimental harvest. In this report we compared pre-harvest biotic communities among treatments to ensure that treatments were similar prior to harvest. Post-harvest biotic communities will be surveyed during 1998 and 1999 and compared among treatments to evaluate whether the biotic communities present within the first two years post-harvest are related to clump size and distance between clumps. Our results will be compared to that found in benchmark communities and to areas where other harvest prescriptions have been used.

We will evaluate: (1) whether the bird community that has specialized foraging and nesting requirements for dead trees varies among the three silviculture treatments, (2) whether the resident bird community varies among treatments, (3) whether the neotropical migrant bird community varies among treatments, (4) whether the bat community varies among treatments, (5) whether the arthropod community varies among treatments, (6) whether herb and shrub communities vary among treatments, and (7) whether fall-down rates of standing trees and snags vary among treatments.

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CHAPTER 2. STUDY AREA AND METHODS

Jim Schieck

The study is being conducted in two forest types in the boreal forest of western Alberta. The study area southwest of Whitecourt, AB (54° 10' N, 116° 20' W) is dominated by jack pine and black spruce that is approximately 110 years old and has between 2,000 and 2,200 canopy trees per ha (810-890 canopy trees per acre).

Throughout the area there is localized topographical relief resulting in pockets (<5% of the total area) of aspen and white birch.

Twelve stands that range in size from 27 to 46 ha, mean 34 ha (67 to 114 acres, mean 84 acres) were established in the study area (Fig. 2.1).

These stands were categorized into four groups of three, based primarily on the type of vegetation that surrounded them and secondarily on the tree species composition in the area prior to harvest. Within each stand group one stand was randomly assigned to each of the three treatments (Fig. 2.1).

Stands were harvested between September 1997 and February 1998. Three percent of the pre-harvest volume was left in tree clumps and as many of the non-merchantable trees and snags as was possible were retained. Clumps were not placed on roads or on cut-lines, but otherwise they were dispersed in a regular pattern. To reduce wind-throw, clumps were circular and all trees within 3 m (10 ft.) of the perimeter of each clump were topped at 4-5 m (13-16 ft.) to create stubs. We have assumed that each stub contains one third of the volume of the original tree and have added sufficient area to each clump to replace the volume that was removed from the stubs. The number of clumps in a stand was determined based on the treatment being created and the size of the stand. Based on a stand area of 30 ha (74 acres), treatment one would have two large clumps with a diameter of 79.6 m (87.0 yd.) and each clump would contain 850-935 residual trees and 145-160 stubs, treatment two would have eight medium sized clumps with a diameter of 41.7 m (45.6 yd.) and each clump would contain 200-220 residual trees and 75-80 stubs, and treatment three would have 32 small clumps with a diameter of 22.7 m (24.8

yd.) and each clump would contain 45-50 residual trees and 35-40 stubs. During the summer of 1998 the harvested areas within the stands will be re-planted with 2-year old pine seedlings. During September 1999, the stands will be sprayed with herbicide.

The study area southwest of Grand Prairie, AB (54° 50' N, 119° 20' W) is dominated by aspen trees that originated after a fire approximately 100 years ago. This study area has a few small depressions creating pockets of white spruce, balsam poplar, white birch, willow and black spruce. Within this study area there are approximately 800-1,000 canopy trees per ha (324-405 canopy trees per acre). The experimental design is similar to that for the Whitecourt study area. Twelve experimental stands were created by harvesting blocks in a checker-board pattern such that approximately half of the study area was harvested and half of the area unharvested (Fig. 2.2).

Stands range in size from 35 to 66 ha, mean 52 ha (86 to 163 acres, mean 128 acres). These stands were categorized into four groups of three, based primarily on the type of vegetation that surrounded them and secondarily on the tree species composition in the area prior to harvest. Within each group, one stand was randomly assigned to each of the three treatments (Fig. 2.2). Stands were harvested between September 1997 and February 1998. Three percent of the pre-harvest volume was left in tree clumps and as many of the non-merchantable trees and snags as was possible were retained in each stand. Clumps were not placed on roads or cut-lines, but otherwise were dispersed in a regular pattern throughout the stand. The number of clumps in a stand was determined based on the treatment being created and the size of the stand. Based on a treatment area of 50 ha (124 acres), treatment one would have two large clumps with a diameter of 97.7 m (107.0 yd.) and each clump would contain 600 - 750 residual trees, treatment two would have ten medium sized



FIGURE 2.1. Locations of sites and stands in the Whitecourt study area.



1 km

FIGURE 2.2. Locations of sites and stands in the Grande Prairie study area.
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clumps with a diameter of 43.7 m (47.8 yd.) and each clump would contain 120 - 150 residual trees, and treatment three would have 50 small clumps with a diameter of 19.5 m (21.3 yd.) and each clump would contain 24 - 30 residual trees. No post-harvest silviculture is planned for the stands.

To ensure that harvest patterns were consistent among treatments, the boundaries of the clumps of trees and snags that were retained at harvest were marked with colored plastic ribbon prior to harvest. No machine activity occurred within the residual clumps of trees during harvest.

General Survey Methods

Pre-harvest surveys of biota were conducted during the spring and summer of 1997. Ten 1-ha (2.47 acre) sites were established in the center of each stand (Figs. 2.1 and 2.2) and all surveys of forest structure and biota were conducted in those central sites. Trees, snags, downed woody material, and understory vegetation were surveyed in a series of nested plots and transects (Fig. 2.3, see Chapters 3 - 5). Breeding birds were surveyed using a combination of point counts and strip transects (Chapter 7). Bats were surveyed using automatic remote bat detectors (Chapter 8). Post-harvest surveys of biota will be conducted during the spring and summers of 1998 and 1999 (Table 2.1, see Chapters 3 - 8).

Analysis and Reporting

All field work was coordinated by Troy Sorensen a full-time researcher at Alberta Research Council. Analyses and reporting was conducted during the fall and winter by researchers at Alberta Research Council, Vegreville, AB.

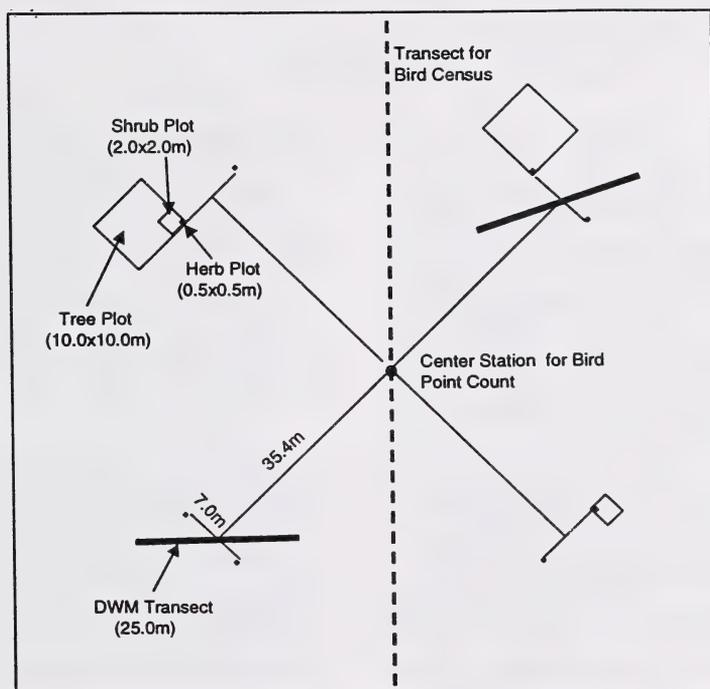


FIGURE 2.3. An example of the transect and nested-plot layout for a 1-ha (2.47 acre) site.

TABLE 2.1. Schedule by which forest structure and biota will be surveyed during the project.

Element	1997 (Pre-Harvest)	1998 (Post-harvest)	1999 (Post-harvest)
Trees and Snags	extensive survey	extensive survey	cursory survey
Down woody Material	extensive survey	cursory survey	none
Understory Vegetation	cursory survey	none	extensive survey
Arthropods	none	extensive survey	cursory survey
Birds	extensive survey	cursory survey	extensive survey
Bats	extensive survey	extensive survey	extensive survey

CHAPTER 3. TREES PRE-HARVEST CHARACTERISTICS AND COMPOSITION OF TREE AND SNAG COMMUNITIES IN GRANDE PRAIRIE AND WHITECOURT STUDY AREAS.

Susan Crites

Introduction

The natural disturbance model of ecosystem management assumes that biota are adapted to natural disturbances and are more likely to maintain viable populations if forest harvesting is as similar as possible to natural disturbances (Swanson and Franklin 1992). In Alberta's boreal mixedwood forests, fire is the dominant natural disturbance and is currently being used as a template for timber harvesting by some companies (Anonymous 1997).

Past research has demonstrated that species respond to forest structure (Stelfox 1995). Timber harvesting, which removes structure, will undoubtedly have negative consequences on biota that were present in the stand before it was harvested. Structure in the form of large trees and snags is often in abundance after fire, and its presence, mainly in the form of snags and downed woody material, is often seen long after the disturbance (Lee et al. 1997). To allow timber harvesting to more closely approximate fire, forest managers are advocating that residual merchantable material be left behind at harvest. The appropriate amount and distribution of residual material on cutblocks, however, is much debated. Clumps left behind at harvest could be compared to fire skips that serve as refugia and dispersal sites; therefore, larger clumps may function as better refugia. Conversely, residual material evenly distributed throughout a cutblock could serve to put structure into mature forests that is usually absent from harvest-origin forests until older ages, and that serves as habitat, cover, and foraging sites. Of the two patterns, it is not known which will be more beneficial to biota.

Trees species respond differently to wind depending on their structural integrity, decay properties, exposure to wind during development, and sheltering ability of

surrounding vegetation. Trees sheltered within an intact forest tend to have smaller boles, branches concentrated in the canopy, and are not as strongly rooted in the soil as trees that have been exposed to wind during development. Newly exposed residual trees left post-harvest are susceptible to the turbulent and accelerated wind in cutover areas because they have not developed the structural integrity to withstand such winds (Kimmins 1996).

Deciduous species such as aspen and balsam poplar are more likely to respond to wind by snapping off along the bole than to be root thrown. This is due to their concentration of branches in the canopy, the depth and complexity of their rooting system, and their susceptibility to fungal attack that weakens boles (Peterson and Peterson 1992). Coniferous species such as black and white spruce respond to wind by being root thrown. This is because of their shallow rooting system and high surface area (consisting of needles and branches) exposed to wind that extends the entire length of the bole.

Site conditions also influence the ability of trees to remain standing pre- and post-harvest. In soils saturated with water, root systems are poorly anchored and trees have the tendency to fall over. In addition, frost heaving can uplift trees from the soil. Season of harvest will influence root system stability and/or tree health in areas directly affected by equipment. Timber harvesting when the ground is not frozen can lead to soil compaction, and, if compaction is very high, it can kill root systems, leading to tree death.

In this report I evaluate the pre-harvest density and characteristics (i.e., species, heights, and diameters) of trees and snags among stand groups and treatments within each study area (see Chapter 2 for experimental design). Post-

harvest, I will determine whether tree death and tree and snag fall down rates vary among treatments.

Methods

For a description of study areas and methods, see Chapter 2. Trees and snags were sampled in every second site in each stand. Each one ha sampling site was divided into four quadrants, and a 10 x 10 m (11 x 11 yd.) tree plot was placed in two randomly selected quadrants. Tree plots were located close to the centre of each quadrant (Fig. 2.3). Within each plot, density, diameter at breast height [1.4 m (4.6 ft.)], height class [1 = 1.4-5 m (4.6-16.4 ft.), 2 = 5.1-10 m (16.5-32.8 ft.), 3 = 10.1-20 m (32.9-65.6 ft.), 4 = >20 m (>65.6 ft.)], top condition (broken canopy, broken bole, or intact) and species were recorded for trees and snags that were ≥ 1.4 m (≥ 4.6 ft.) tall. For snags, decay stage was also recorded (six stages from Lee et al. 1997; stage one being a recently killed tree, stage six being a broken off, well-decayed bole). Diameter distributions of trees and snags were plotted and evaluated to separate stems of the canopy cohort from stems of the subcanopy or understory. Only stems of the canopy cohort were used in analyses. For snags, "canopy" refers to trees greater than a particular diameter (see Results), irrespective of whether the stem is actually part of the canopy (i.e., broken off).

General Linear Models (GLM, SAS Institute, Inc. 1989) were used to evaluate differences in abundances of canopy trees and snags among stand groups and treatments. Analyses were completed on log transformed data. Post hoc tests were completed using Tukey's Honestly Significant Difference test (Zar 1984; SAS Institute, Inc. 1989). All results were considered statistically significant if the probability of them occurring by chance was less than 0.05.

Results

Live Trees

A total of 2672 and 4802 trees and snags were sampled in Grande Prairie and Whitecourt, respectively. In Grande Prairie there were four tree species found during the surveys with aspen being the most common followed by balsam

poplar, paper birch, and white spruce (Table 3.1). Seven species were found during the surveys in Whitecourt, with jack pine and black spruce being the most common followed by balsam fir, paper birch, aspen, white spruce, and balsam poplar (Table 3.1).

Stem Diameter and Height

Not surprisingly, stems increased in diameter as they increased in height. For both study areas, the majority of stems were short [1.4-5 m (4.6-16.4 ft.)] and in diameter class 0-5cm (0-2 in.) (Figures 3.1 and 3.2). In Grande Prairie, the second most abundant diameter class was 15.1-20 cm (5.9-7.9 in.); therefore, canopy trees in Grande Prairie likely consisted of trees >15 cm (>5.9 in.) dbh (Figure 3.1). The majority of trees with diameters >15 cm (>5.9 in.) were >20 m (>65.6 ft.) tall (Figure 3.1). By examining the relationship between diameter and height at Whitecourt, canopy trees likely consisted of trees >10 cm (>3.9 in.) dbh (Fig. 3.2). Unlike trees in Grande Prairie, heights of canopy trees were between 10.1-20 m (32.9-65.6 ft.), with few >20 m (>65.6 ft.) (Figure 3.2).

Only canopy trees, [i.e., stems >10 cm (>3.9 in.) and >15 cm (>5.9 in.) dbh for Grande Prairie and Whitecourt, respectively] were used in subsequent analyses. Otherwise, the high frequencies of smaller trees in both study areas would dominate density calculations and interpretations.

Canopy Stem Density

Average number of canopy stems per hectare was 746.8 (± 75.0 SE) [302.3 (± 30.4 SE) stems/acre] and 1441.8 (± 123.5 SE) [583.7 (± 50.0 SE) stems/acre] for Grande Prairie and Whitecourt, respectively. In Grande Prairie, aspen dominated all stand groups (Table 3.2). When all species were combined in Grande Prairie, there were no significant differences in stems per ha among stand groups or treatments (Table 3.2). For individual species in stand groups, lower densities of aspen and higher densities of balsam poplar were found in stand group C (Table 3.2). There were no differences among treatments for individual species (Table 3.2).

TABLE 3.1. Percent species composition of trees and snags ≥ 1.4 m (≥ 4.6 ft.) in height in Grande Prairie and Whitecourt study areas. Numbers in parentheses indicate the actual number sampled.

Species	Trees		Snags	
	Grande Prairie	Whitecourt	Grande Prairie	Whitecourt
<i>Populus tremuloides</i>	81 (1937)	2 (79)	78 (222)	2 (22)
<i>Populus balsamifera</i>	8 (194)	0 (2)	16 (44)	0
<i>Betula papyrifera</i>	6 (137)	19 (726)	4 (12)	9 (87)
<i>Picea glauca</i>	5 (121)	2 (74)	1 (2)	1 (15)
<i>Pinus banksiana</i>	0	27 (1033)	0	34 (335)
<i>Abies balsamea</i>	0	22 (828)	0	2 (17)
<i>Picea mariana</i>	0	28 (1070)	0	51 (507)
unknown	0	0	1 (3)	1 (7)
Total (N)	100 (2389)	100 (3812)	100 (283)	100 (990)

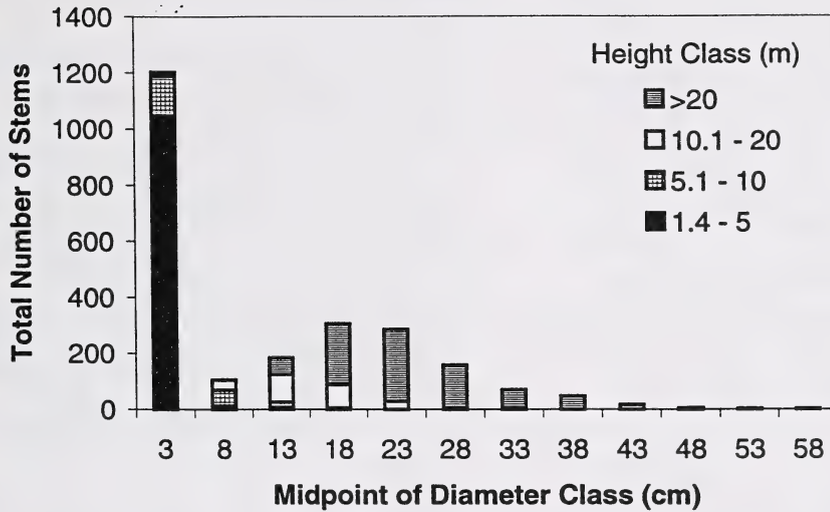


FIGURE 3.1. Total number of live trees categorized by diameter and height classes for Grande Prairie.

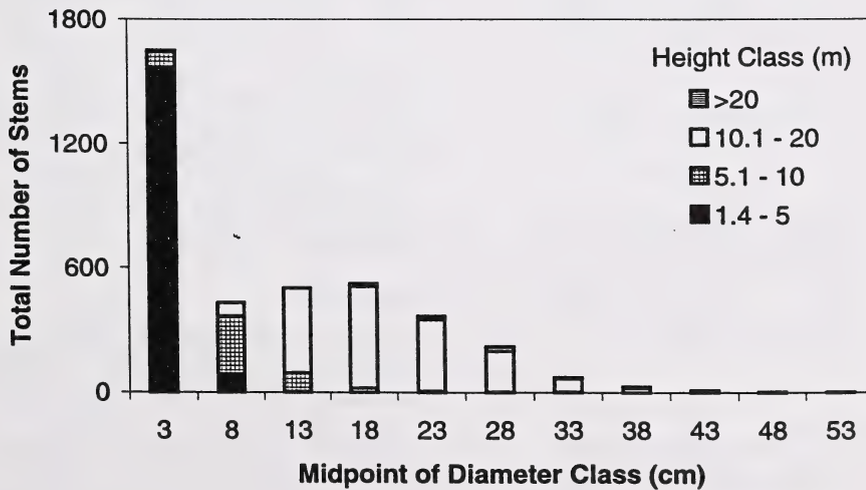


FIGURE 3.2. Total number of live trees categorized by diameter and height classes for Whitecourt.

In Whitecourt, jack pine dominated all stand groups and treatments, followed closely by black spruce. For all species combined in Whitecourt, stand group A had the highest stem densities and stand group B the lowest (Table 3.3). Jack pine and black spruce were present in higher densities in Stand group A, and in lower densities in stand group B, than in remaining stand groups. On the other hand, white spruce and balsam fir were present in higher densities in stand group B than in remaining stand groups, but these densities were relatively low (Table 3.3). When all stems were combined, there were no significant differences among treatments in Whitecourt; however, aspen was present in higher densities in treatment one than in remaining treatments, and white spruce was absent from treatment three (Table 3.3).

Snags

Of the total number of stems sampled, 11% (283) and 21% (990) were snags in Grande Prairie and Whitecourt, respectively. Dominant snag species were the same as the dominant tree species for each of the two locations. There were no paper birch snags of canopy size in Grande Prairie, and no balsam poplar snags in Whitecourt.

Snag Diameter

Diameter distributions for snags were different than for live trees for both study areas. The snag distribution in Grande Prairie peaked at 10.1-15 cm (4-5.8 in.) dbh (Figure 3.3), whereas in Whitecourt, the distribution was high from 0-10 cm (0-3.9 in.) dbh, then decreased to very few trees >30 cm (>11.8 in.) (Figure 3.4). Height measurements for snags are not presented because many snags snapped along the bole. The same diameter cut offs as used for canopy trees was used for the snag analyses (i.e., >15 cm (>5.9 in.) and >10 cm (>3.9 in.) dbh for Grande Prairie and Whitecourt, respectively).

Canopy Snag Density

There was a mean of 77.5 (± 19.2 SE) [31.4 (± 7.8 SE) per acre] and 289.3 (± 44.8 SE) [117 (± 18 SE) per acre] large snags per hectare in Grande Prairie and Whitecourt, respectively. In Grande Prairie, density of snags did not vary significantly among groups for all species

combined (Table 3.2); however, balsam poplar and birch had significantly higher densities in stand group C compared to remaining groups (Table 3.2). Density of large snags did not vary significantly among treatments (Table 3.2).

Similar to tree density, density of large snags in Whitecourt varied significantly among groups when all species were combined, with stand group B having the lowest density of snags and stand group A the highest (Table 3.3). The low density of snags in stand group B was due to the low densities of jack pine snags (Table 3.3). Density of black spruce snags was significantly higher in stand group A as compared to remaining stand groups (Table 3.3). Densities of large snags did not vary among treatments for all species combined, but for individual species, density of balsam fir snags was higher in treatment two, than in remaining treatments (Table 3.3).

Decay

In all dbh classes, snag density tended to decrease as decay state increased (Fig. 3.3 and 3.4). More than 50% of the snags in Grande Prairie have decay stage of one, with 82% of the snags being in decay stages one, two, and three (Figure 3.3). In Whitecourt 50% of the snags were within decay stages one or two and 95% of snags were in the first four decay stages (Figure 3.4).

Top Condition

Of all snags sampled, 66.2% and 67.5% had intact boles in Grande Prairie and Whitecourt, respectively. In Grande Prairie, 26.6% of snags had broken boles and 7.2% had broken canopies, compared to 16.6% and 15.9% in Whitecourt, respectively.

Discussion

Composition and characteristics of the tree and snag communities were different between the two study areas. Grande Prairie was an aspen-dominated community with canopy trees commonly reaching heights greater than 20 m (65.6 ft.) and diameters greater than 15cm (5.9 in.) dbh. The community was homogeneous with only four species and having aspen dominate all groups and treatments. The

TABLE 3.2. Mean (\pm SE) stems per hectare of canopy trees and snags [>15 cm (>5.9 in.) dbh] within each stand group and treatment for Grande Prairie study area. There were 3 and 108 degrees of freedom and 2 and 108 degrees of freedom for stand groups and treatments, respectively. Means with the same superscript letter are not significantly different. Aw = aspen, Sw = white spruce, Pb = balsam poplar, Bw = paper birch.

	TREES					SNAGS				
	All	Aw	Sw	Pb	Bw	All	Aw	Pb	Bw	
Stand Group										
A	900 (70.8)	850 ^a (75)	27 (10)	20 ^b (14)	3 (3)	63.3 (16.2)	63 (16)	0 ^b (0)	0 ^b (0)	
B	670 (71)	627 ^a (71)	17 (10)	27 ^b (14)	0 (0)	80 (21.7)	60 (21)	20 ^{ab} (9)	0 ^b (0)	
C	587 (67.4)	283 ^b (74)	17 (17)	270 ^a (52)	17 (10)	80 (19.4)	20 (7)	50 ^a (18)	10 ^a (6)	
D	830 (91.1)	770 ^a (83)	50 (21)	10 ^b (10)	0 (0)	86.7 (19.6)	77 (20)	10 ^{ab} (6)	0 ^b (0)	
Test Statistic	F 2.5 p 0.06	15.5 0.001	1.4 0.2	21.2 0.001	2.4 0.07	0.3 0.8	2.3 0.09	4.1 0.008	3.2 0.02	
Treatment										
1	758 (71.4)	645 (79)	18 (13)	85 (34)	10 (6)	72.5 (12.4)	53 (12)	15 (7)	5 (3)	
2	723 (59.1)	595 (68)	15 (9)	108 (32)	5 (5)	100 (19.9)	70 (19)	28 (11)	3 (3)	
3	760 (73.1)	658 (74)	50 (16)	53 (21)	0 (0)	60 (16.3)	43 (13)	18 (11)	0 (0)	
Test Statistic	F 0.04 p 1.0	0.2 0.8	3.5 0.05	1.7 0.2	1.5 0.2	1.8 0.2	0.7 0.5	0.6 0.6	1.1 0.3	

TABLE 3.3. Mean (\pm SE) stems per hectare of canopy trees and snags [>10 cm (>3.9 in.) dbh] within each stand group and treatment for Whitecourt study area. There were 3 and 108 degrees of freedom and 2 and 108 degrees of freedom for stand groups and treatments, respectively. Means with the same superscript letter are not significantly different. Aw = aspen, Sw = white spruce, Pb = balsam poplar, Bw = paper birch, Pj = jack pine, Sb = black spruce, Fb = balsam fir, Con = unidentified conifer species.

	TREES										SNAGS					
	All	Aw	Sw	Pb	Bw	Pj	Sb	Fb	All	Aw	Sw	Bw	Pj	Sb	Fb	Con
Stand Group A	1980 ^a (142)	37 (15)	3 ^b (3)	7 (7)	83 (29)	973 ^a (132)	860 ^a (109)	17 ^b (8)	450 ^a (71.9)	20 (14)	0 ^b (0)	57 (22)	123 ^{ab} (28)	217 ^a (63)	30 (19)	3
B	983 ^c (115)	60 (32)	53 ^a (18)	0 (0)	70 (21)	397 ^b (65)	303 ^{bc} (88)	100 ^a (41)	207 ^b (38.6)	3 (3)	17 ^a (8)	23 (8)	77 ^b (27)	70 ^b (23)	6 (0)	3
C	1527 ^{ab} (149)	67 (33)	0 ^b (0)	0 (0)	70 (22)	847 ^a (95)	533 ^b (124)	10 ^b (7)	253 ^{ab} (37.7)	17 (12)	0 ^b (0)	27 (11)	143 ^{ab} (36)	67 ^b (19)	0 (0)	0
D	1277 ^{bc} (88)	97 (41)	3 ^b (3)	0 (0)	33 (11)	105 ^a (103)	93 ^c (35)	0 ^b (0)	247 ^{ab} (31)	17 (8)	0 ^b (0)	17 (8)	183 ^a (33)	30 ^b (11)	0 (0)	0
Test Statistic F	11.1	0.4	9.1	1.0	0.7	7.4	15.6	5.6	3.7	0.6	4.2	1.15	2.8	5.6	2.4	0.08
p	0.001	0.7	0.001	0.4	0.5	0.001	0.001	0.001	0.02	0.6	0.01	0.33	0.04	0.001	0.07	0.6
Treatment 1	1560 (131)	130 ^a (43)	22 ^{ab} (12)	0 (0)	38 (14)	828 (95)	533 (107)	10 (6)	265 (28.1)	30 ^a (11)	5 (3)	15 (6)	113 (21)	100 (22)	3 ^b (3)	0 (0)
2	1270 (86)	45 ^{ab} (12)	23 ^a (8)	0 (0)	78 (21)	738 (79)	358 (66)	30 (11)	338 (57.5)	10 ^{ab} (10)	8 (6)	30 (14)	155 (33)	103 (44)	28 ^a (14)	5 (3)
3	1495 (141)	20 ^b (10)	0 ^b (0)	5 (5)	78 (20)	885 (111)	453 (101)	55 (31)	265 (39.3)	3 ^b (3)	0 (0)	48 (14)	128 (28)	85 (28)	3 ^b (3)	0 (0)
Test Statistic F	0.8	3.2	3.5	1.0	1.9	0.2	0.31	1.1	0.8	4.0	1.1	2.1	0.2	0.6	3.3	2.0
p	0.5	0.05	0.03	0.4	0.2	0.9	0.7	0.3	0.5	0.02	0.3	0.1	0.8	0.5	0.04	0.1

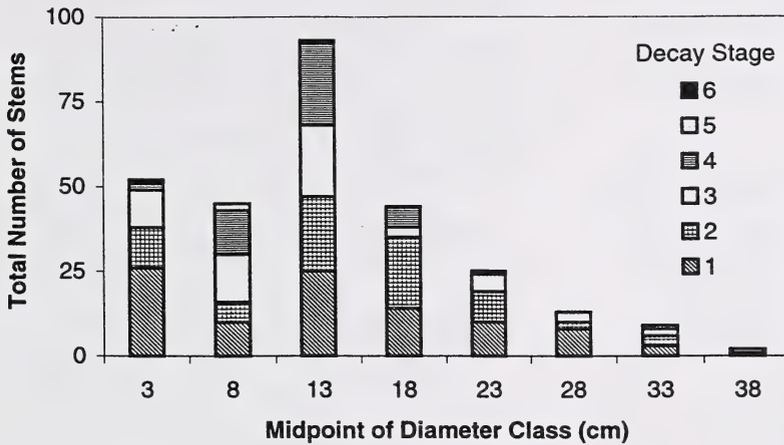


FIGURE 3.3. Total number of snags categorized by diameter class and decay stage for Grande Prairie. Decay stage one refers to a recently dead tree; decay stage six refers to a well-decayed snag that is soft in places with a broken bole.

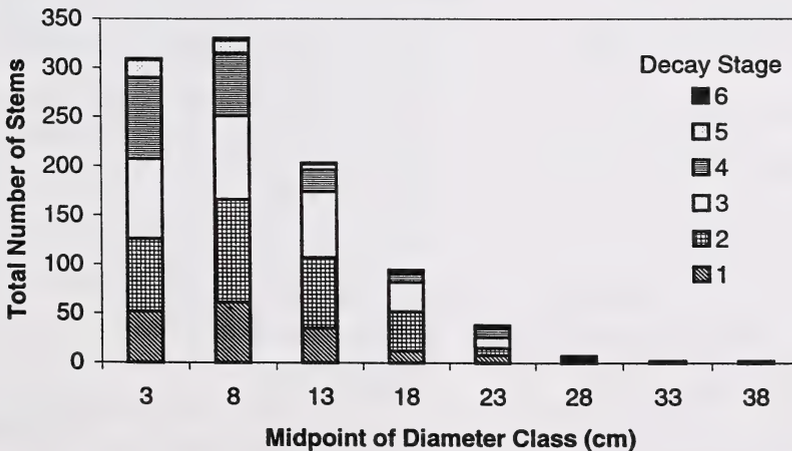


FIGURE 3.4. Total number of snags categorized by diameter class and decay stage for Whitecourt. Decay stage one refers to a recently dead tree; decay stage six refers to a well-decayed snag that is soft in places with a broken bole.

High densities of trees in small diameter classes indicates trees are continually being recruited into the understory and subcanopy. If gaps form in the canopy, these understory trees may grow to become part of the canopy. If no gaps form before these understory trees are out-competed, they will become snags.

As with trees, aspen also dominated the snag community in Grande Prairie resulting in snags with a greater frequency of broken boles than snags in Whitecourt. The self-thinning nature of aspen stands is represented in the median snag diameter of 10.1-15 cm (4-5.9 in.) in Grande Prairie, which is one diameter class lower than the median diameter class for live canopy trees [15.1-18 cm (6.0-7.1 in.) dbh]. As aspen stands grow, they compete for resources and continue to self-thin until they reach mature ages and canopy equilibrium is obtained (Peterson and Peterson 1992). As a result, snags are continuously being recruited into diameter classes smaller than the canopy live trees. Decay stage one was the most common decay stage for all snag sizes, with the exception of snags having a dbh of 5.1-10 cm (2.1-3.9 in.), and snags greater than 30 cm (11.8 in.) dbh. Surprisingly, many snags that are 5.1-15 cm (2.1-5.9 in.) dbh have been present long enough to have decayed to stages three, four, and five.

The tree community in Whitecourt was more heterogeneous than in Grande Prairie with two species, jack pine and black spruce, dominating and eight species present. Canopy trees in Whitecourt were shorter and had smaller dbh than canopy trees in Grande Prairie. As in Grande Prairie, trees in Whitecourt were abundant in small diameter classes, indicating a high recruitment of trees into the understory and subcanopy. The fate of these trees will depend on canopy gap formation and competition dynamics between tree species. Tree species richness was high in Whitecourt, leading to different potential successional patterns than found in Grande Prairie. Balsam fir and paper birch were common in the understory of many stands. Balsam fir is a late successional shade-tolerant species that, in the absence of disturbance, may eventually replace the current canopy cohort and form a 'climax' community (Kimmins 1996).

Based on vegetation within and surrounding stands, each stand was assigned to one of four stand groups. Stand groupings accounted for a small amount of variation in the tree and snag communities of both study areas. In addition to having fewer species, the tree and snag community in Grande Prairie exhibited fewer differences between groups than in Whitecourt, indicating that this community was more homogeneous.

There were no differences among treatments for tree and snag communities in Grande Prairie. In Whitecourt, there were differences among treatments for aspen and white spruce live trees, and for aspen and balsam fir snags. When examining the Grande Prairie community post-harvest, any differences in the tree and snag communities among the different residual patches will be attributed to the harvesting treatment, and not due to any pre-harvest differences. In Whitecourt, any post-harvest differences in the tree and snag communities will have to be evaluated after taking into account the differences among treatments prior to harvest.

Future Research

Tree and snag communities will be surveyed and monitored in 1998 and 1999. In spring of 1998, all residual trees and snags will be permanently marked within a subset of randomly selected patches. In addition, a subset of snags and trees in the cut-over area will be marked. The fate of this residual material will be regularly determined in the summer of 1998 and 1999. In 1999, I will determine which configuration of patch size and dispersion is most beneficial to the long-term persistence of standing trees and snags.

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CHAPTER 4. CHARACTERISTICS OF DOWNED WOODY MATERIAL IN RELATION TO THE DISPERSION OF RESIDUAL TREES FOLLOWING HARVEST

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Introduction

Bryophytes, lichens, fungi, invertebrates, and herbaceous plants compose the majority of biodiversity in an ecosystem. Many of these taxa are influenced by, or dependent on, the abundance, size, spatial distribution, and decomposition of downed woody material (DWM; McCullough 1948; Thompson 1980; Harmon et al. 1986; Söderström 1988a,b; Crites and Dale 1995, in press). DWM is also crucial as nest sites (Thompson 1980), food, and protection for vertebrates and invertebrates (Harmon et al. 1986). In addition to its direct influence on biodiversity, DWM functions as an important reservoir for ecosystem nutrients (Lambert et al. 1980, MacMillan 1981, Abbott and Crossley 1982, Harmon et al. 1986).

Large DWM in a wide range of decay stages is often stressed as important in maintaining the species composition of DWM-dependent taxa (Söderström 1988a,b; Lesica et al. 1991; Crites and Dale 1995, in press). Clearcutting, however, reduces the density, volume, size distributions (Abbott and Crossley 1982; Lesica et al. 1991) and decay class distributions of DWM (Söderström 1988a,b; Crites and Dale 1995, in press; Lee et al. 1995, 1997). One strategy to minimize these effects is to leave residual live trees, snags, and DWM in harvested areas (Lee et al. 1995, 1997) in an effort to promote natural size and decay class distributions of DWM, and the establishment of DWM-dependent taxa (Söderström 1988b). If forest managers are willing to leave this residual material, the question remains of how to distribute it within harvested areas. An even spatial distribution of DWM will facilitate the dispersal of DWM-dependent species, but will minimize the amount of interior-forest environment which promotes natural decomposition rates of DWM.

Currently this study has two objectives: 1) to provide pre-harvest information on the amount

and characteristics DWM in the two study areas and 2) to test for differences in DWM among stand groups prior to harvest. During 1998-99, post-harvest density, volume, spatial distribution, and decomposition of DWM will be surveyed and compared among stand treatments. Current results will be used to account for pre-harvest differences among stands.

Methods

A description of the study areas and experimental harvesting is presented in Study Area and Methods (Chapter 2). Ten 1-ha (2.47 acre) sites were established in the center [*ca.* 100 m (109 yd.) from the edge] of each stand and all surveys for biota were conducted within those sites (Fig. 2.1 and 2.2). DWM was surveyed along two 25 m (27 yd.) transects within the center of two randomly selected quadrants of each site (Fig. 2.3). The direction of each transect was random. All DWM intersecting the transect with diameters ≥ 5 cm (≥ 2 in.) were tallied; in addition, the diameter at line intersect and decay class (Table 4.1) of large pieces (≥ 10 cm [≥ 4 in.]) were measured. DWM with diameters between 5-10 cm (2-4 in.) and ≥ 10 cm (≥ 4 in.) will be called small and large DWM, respectively.

The 12 stands in both study areas were assigned into 4 groups to group stands that had the most similar tree composition. The three stands in each group were then randomly assigned a treatment of residual tree dispersion (see Chapter 2). Harvesting occurred during the winter of 1997-98.

Linear densities of DWM were calculated as the number of pieces per 5 m (16 ft.) of transect and are referred to in the rest of this chapter as densities. The volume of DWM per unit area

TABLE 4.1. Descriptions of decay classes (modified from Lee et al. 1997) assigned to large [>10 cm (>4 in.) diameter] DWM.

Decay Classes	DWM Characteristics
1	Log whole and undecayed; bark, branches, and twigs present and intact; log elevated on support points; wood hard.
2	Log sound, wood hard; twigs mostly lacking; $<50\%$ of the bark missing
3	Wood soft in places; some branches remaining; $>50\%$ bark missing; wood soft in places
4	Little to no bark remaining; no branches; wood fungi present; wood soft with small crevices and small pieces lost.
5	Large wood fragments lost; outline of trunk slightly deformed; vascular and non-vascular plants beginning to colonize.
6	Wood mostly well-decayed; log colonized by various vascular and non-vascular plants.
7	Humification nearly 100% ; hard to define as log, outline indeterminable; no evidence of hard wood.

(v) was calculated from the following formula (Van Wagner 1968):

$$v = \frac{\pi^2 \sum_{i=1}^n d_i^2}{8l}$$

where l is the transect length [25 m (27 yd.)], d is the diameter (cm) of the large pieces (>10 cm (>4 in.)) or 7.5 cm (3 in.) for small pieces [5-10 cm (2-4 in.)], and i is the number of pieces intersecting the transect. The resulting units are cubic meters per hectare. Densities and volumes were calculated for both small and large pieces and then summed for a total value. Volume was also calculated for each decay class (1-7) of the large pieces. The species composition of DWM was not analyzed because it should be directly related to the species composition of trees and snags (Chapter 3).

General Linear Models (GLM, SPSS 1997) were used to examine differences in the densities and volume of DWM between stand groups and treatments. Tukey's Honestly Significant Difference tests ($\alpha = 0.05$) were used for all post hoc tests (Zar 1984; SPSS 1997). Volumes of small DWM were not analyzed separately because they were proportionally related to small DWM densities. The spatial heterogeneity of DWM distribution was examined by using a nested design (sites within stands).

Results

Grande Prairie Study Area

DWM Densities

There was $0.99 \pm 0.08\text{SE}$ pieces of small DWM and 0.94 ± 0.06 pieces of large DWM every 5m (16 ft.) in the Grande Prairie study area. DWM density differed among stand groups for small ($F_{3,108} = 4.558, p = 0.005$), large ($F_{3,108} = 2.690, p = 0.050$), and total DWM ($F_{3,108} = 6.177, p = 0.001$; Table 4.2). Higher densities of DWM were found in stand groups near bog areas (Group A and D, Table 4.2, Fig. 2.2).

There was no significant difference in the densities of large ($F_{2,108} = 0.027, p = 0.973$) or

total DWM ($F_{2,108} = 1.930, p = 0.150$) between treatments; however, there was a significant difference between treatments in the density of small DWM ($F_{2,108} = 3.800, p = 0.025$, Table 4.2).

The spatial distribution of DWM density was heterogeneous within and among stands. Small and large DWM densities were spatially heterogeneous both within ($F_{108,120} \geq 1.641, p \leq 0.004$) and among stands ($F_{6,108} \geq 2.357, p \leq 0.035$). On the other hand, total DWM density was significantly different within stands ($F_{108,120} = 1.927, p = 0.0002$) but not among stands ($F_{6,108} = 1.757, p = 0.115$).

DWM Volume

There was $77.4 \text{ m}^3/\text{ha}$ ($\pm 5.1\text{SE}$) [$41.0 \text{ yd}^3/\text{acre}$ ($\pm 2.7\text{SE}$)] of DWM in the Grande Prairie study area. Volume differed among stand groups ($F_{3,108} = 2.952, p = 0.036$, Table 4.3) with group B having lower total DWM volume than other groups (Table 4.3); potentially due to their farther distance from bogs and wet areas (Fig. 2.2). The volume of large DWM did not differ among stand groups ($F_{3,108} = 2.487, p = 0.064$, Table 4.3). The volume of large and total DWM did not differ among stand treatments prior to harvest ($F_{2,108} \leq 0.903, p \geq 0.408$, Table 4.3).

The spatial distribution of large DWM volume was heterogeneous both within ($F_{108,120} = 1.531, p = 0.012$) and among ($F_{6,108} = 2.382, p = 0.034$) stands; however, total DWM volume was heterogeneous within stands ($F_{108,120} = 1.584, p = 0.007$) but not among stands ($F_{6,108} = 1.981, p = 0.075$).

DWM Size

The average diameter of large DWM in the Grande Prairie study area was 15.5 cm ($\pm 0.47\text{SE}$) [6.1 in. ($\pm 0.19\text{SE}$)]. However the sizes of DWM had a negative exponential distribution (Fig 4.1).

The diameter of large DWM varied among stand groups ($F_{3,144} = 4.942, p = 0.003$) and treatments ($F_{2,151} = 3.415, p = 0.035$; Table 4.4)

TABLE 4.2. Average linear density of DWM (\pm SE) among stand groups in both study areas. Superscripts denote homogeneous subsets as determined by Tukey's Honestly Significant Difference ($\alpha = 0.05$) for significant comparisons among groups/treatments.

		Linear Density (number / 5m)		
		Small DWM	Large DWM	Total DWM
Grande Prairie Area				
Group	A	1.24 \pm 0.18 ^a	1.09 \pm 0.10 ^a	2.33 \pm 0.24 ^a
	B	0.80 \pm 0.15 ^b	0.78 \pm 0.12 ^b	1.58 \pm 0.10 ^b
	C	0.90 \pm 0.06 ^b	0.89 \pm 0.11 ^{ab}	1.79 \pm 0.13 ^{bc}
	D	1.01 \pm 0.17 ^{ab}	0.99 \pm 0.11 ^{ab}	2.00 \pm 0.19 ^{ac}
Treatment	1	1.06 \pm 0.16 ^a	0.95 \pm 0.02	2.01 \pm 0.15
	2	0.82 \pm 0.09 ^b	0.93 \pm 0.06	1.75 \pm 0.15
	3	1.09 \pm 0.14 ^a	0.93 \pm 0.18	2.02 \pm 0.28
Whitecourt Area				
Group	A	1.16 \pm 0.09 ^a	1.16 \pm 0.06 ^a	2.32 \pm 0.11 ^a
	B	0.85 \pm 0.19 ^b	2.02 \pm 0.53 ^b	2.87 \pm 0.63 ^b
	C	1.35 \pm 0.17 ^a	1.42 \pm 0.20 ^{ac}	2.77 \pm 0.03 ^b
	D	1.08 \pm 0.09 ^{ab}	1.70 \pm 0.36 ^c	2.77 \pm 0.37 ^b
Treatment	1	1.18 \pm 0.17	1.38 \pm 0.32 ^a	2.56 \pm 0.29
	2	1.07 \pm 0.04	1.74 \pm 0.43 ^b	2.82 \pm 0.45
	3	1.08 \pm 0.20	1.59 \pm 0.16 ^{ab}	2.67 \pm 0.11

TABLE 4.3. Average volume per unit area of DWM (\pm SE) among stand groups in both study areas. Superscripts denote homogeneous subsets as determined by Tukey's Honestly Significant Difference ($\alpha = 0.05$).

		Volume (m^3 / ha)		
		Small DWM	Large DWM	Total DWM
Grande Prairie Area				
Group	A	17.3 ± 2.5^a	62.7 ± 6.7	80.0 ± 4.7^{ab}
	B	11.1 ± 2.0^b	47.9 ± 12.9	59.0 ± 10.8^a
	C	12.5 ± 0.8^b	74.1 ± 9.4	86.7 ± 9.2^b
	D	14.1 ± 2.4^{ab}	69.8 ± 11.3	83.8 ± 10.7^b
Treatment	1	14.7 ± 2.2^a	63.5 ± 8.3	78.2 ± 7.5
	2	11.3 ± 1.3^b	69.7 ± 4.4	81.0 ± 5.3
	3	15.2 ± 2.0^a	57.7 ± 14.1	72.9 ± 13.8
Whitecourt Area				
Group	A	16.1 ± 1.2^a	88.1 ± 6.7^a	104.2 ± 7.0^a
	B	11.8 ± 2.6^b	239.6 ± 58.4^b	251.4 ± 58.9^b
	C	18.7 ± 2.4^a	128.8 ± 19.5^c	147.5 ± 27.3^c
	D	14.9 ± 1.2^{ab}	164.3 ± 54.0^c	179.2 ± 54.0^c
Treatment	1	16.3 ± 2.4	136.8 ± 43.1	153.1 ± 42.0
	2	14.9 ± 0.5	170.8 ± 58.2	185.7 ± 58.6
	3	14.9 ± 2.7	158.0 ± 35.7	172.9 ± 33.2

with larger DWM in group C and treatment 2 (Table 4.4).

DWM Decay Classes

Very little of the large DWM was undecayed (decay class 1, Fig. 4.2). The proportion of DWM lying on the ground (76.3%) was higher for more decomposed DWM (Fig. 4.2). The decay class distribution of total DWM volume did not differ among stand groups (Fig. 4.3) nor among treatments (Fig. 4.4).

Whitcourt Study Area

DWM Densities

There was $1.11 \pm 0.08\text{SE}$ pieces of small DWM and $1.57 \pm 0.17\text{SE}$ pieces of large DWM every 5m in the Whitcourt study area. Density differed among stand groups (Table 4.2) for small ($F_{3,108} = 4.233, p = 0.007$), large ($F_{3,108} = 17.616, p = 2.22 \times 10^{-9}$), and total DWM ($F_{3,108} = 3.106, p = 0.030$). Low densities for large and total DWM occurred in stands farthest from riparian/creek areas (Group A, Table 4.2, Fig. 2.1).

Varied results occurred when comparing the densities of DWM between stand treatments (Table 4.2). There was no significant difference in the density of small ($F_{2,108} = 0.466, p = 0.629$) or total DWM ($F_{2,108} = 1.138, p = 0.324$) among treatments; however, there was a difference in the density of large DWM among treatments ($F_{2,108} = 5.646, p = 0.005$, Table 4.2). These differences are probably due to the high density of large DWM in B2 [$2.97/5 \text{ m}$ (5.5 yd.) $\pm 0.18\text{SE}$] as a result of windfall (pers. obs.).

Small, large, and total DWM densities were heterogeneous among stands ($F_{6,108} \geq 2.445, p \leq 0.030$) but only small DWM densities were spatially heterogeneous within stands ($F_{108,120} = 1.441, p = 0.026$; large and total DWM, $p \geq 0.056$).

DWM Volume

There was $170.6 \text{ m}^3/\text{ha}$ ($\pm 24.3\text{SE}$) [$90.1 \text{ yd}^3/\text{acre}$ ($\pm 12.9\text{SE}$)] of DWM in the Whitcourt study area. DWM volume differed among stand groups for large ($F_{3,108} = 20.568, p = 1.29 \times 10^{-10}$) and total DWM ($F_{3,108} = 18.858, p = 6.58 \times 10^{-10}$) with group A having the lowest volume and

group B having the highest volume (Table 4.3). Windfall in stand B2 was most likely the cause for the extremely high volume of large [$330.7 \text{ m}^3/\text{ha} \pm 20.7\text{SE}$ ($175.1 \text{ yd}^3/\text{acre} \pm 11.0\text{SE}$)] and total DWM [$346.4 \text{ m}^3/\text{ha} \pm 20.8\text{SE}$ ($183.4 \text{ yd}^3/\text{acre} \pm 11.0 \text{SE}$)]. The volume of large ($F_{2,108} = 1.956, p = 0.146$) and total DWM ($F_{2,108} = 1.759, p = 0.177$) did not differ among treatments (Table 4.3).

The spatial distribution of large and total DWM volume was heterogeneous both within ($F_{108,120} \geq 2.105, p \leq 4.01 \times 10^{-5}$) and among stands ($F_{6,108} \geq 11.160, p \leq 1.16 \times 10^{-9}$).

DWM Size

The average diameter of large DWM in the Whitcourt study area was 17.7 cm ($\pm 0.60\text{SE}$) [7.0 in. ($\pm 0.24\text{SE}$)]; however the sizes of DWM had a negative exponential distribution (Fig. 4.5). Most of the woody material (volume) constituted pieces with diameters of 10-30 cm (4-12 in.) (Fig. 4.5).

The diameter of large DWM differed among stand groups ($F_{3,144} = 15.054, p = 1.418 \times 10^{-3}$; Table 4.4) with group A having the smallest and group B having the largest DWM (Table 4.4). Diameter of large DWM did not differ among stand treatments ($F_{2,144} = 0.170, p=0.844$).

DWM Decay Classes

Most of the volume of DWM was in advanced stages of decomposition (decay class 6 and 7, Fig. 4.6). The proportion of DWM lying on the ground (75.1%) was higher for more decomposed DWM (Fig. 4.6). The decay class distribution of total DWM volume did not differ among stand groups (Fig. 4.7) or among treatments (Fig. 4.8).

Discussion

Small DWM contributes little to the total DWM volume in either study area; however, it is preferred over larger DWM by microarthropods (Abbott and Crossley 1982) and is therefore an important factor in the biodiversity of an ecosystem. Furthermore, small DWM is an important component of nutrient cycling because it quickly decomposes (MacMillan

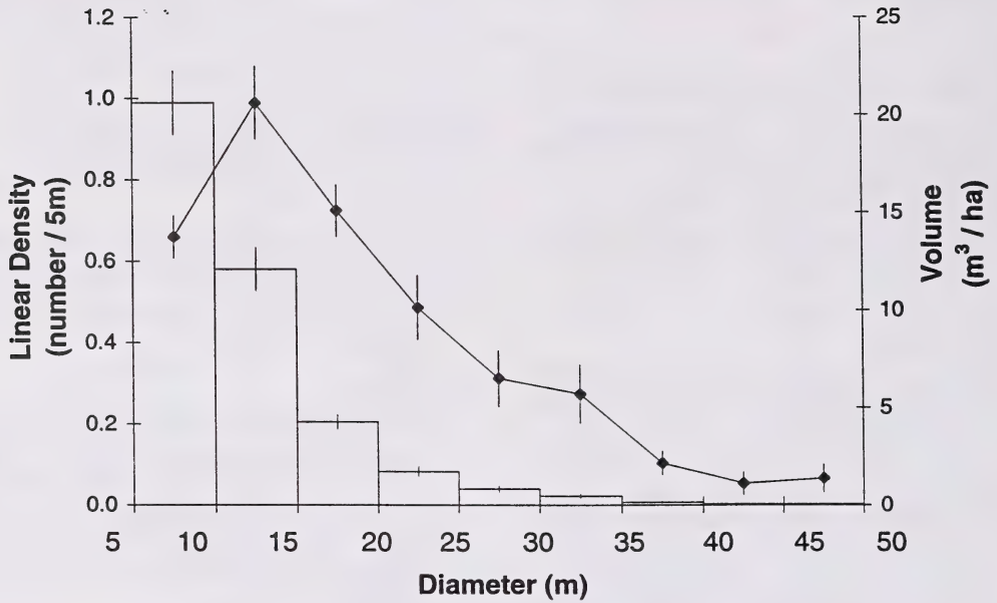


FIGURE 4.1. Size class distribution of the density (histogram) and volume (line) of DWM in the Grande Prairie study area.

TABLE 4.4. Average diameter of large DWM (\pm SE) among stand groups in both study areas. Superscripts denote homogeneous subsets as determined by Tukey's Honestly Significant Difference ($\alpha = 0.05$).

		Diameter (cm)
Grande Prairie Area		
Group	A	14.2 \pm 0.87 ^a
	B	15.0 \pm 1.39 ^{ab}
	C	16.8 \pm 0.50 ^c
	D	15.7 \pm 0.26 ^b
Treatment	1	15.5 \pm 0.80 ^{ab}
	2	16.3 \pm 0.50 ^a
	3	14.6 \pm 1.03 ^b
Whitecourt Area		
Group	A	15.5 \pm 0.21 ^a
	B	20.5 \pm 0.47 ^b
	C	17.2 \pm 0.63 ^c
	D	17.5 \pm 1.06 ^c
Treatment	1	17.9 \pm 1.09
	2	17.4 \pm 1.04
	3	17.7 \pm 1.31

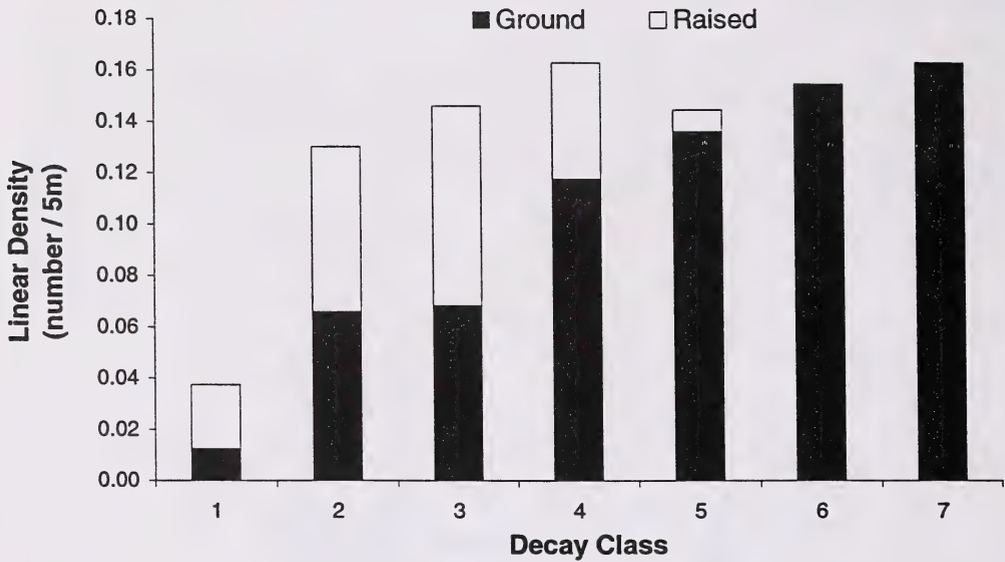


FIGURE 4.2. Average linear density of DWM in each decay class in the Grande Prairie study area distinguishing between DWM raised off the ground and DWM lying on the ground.

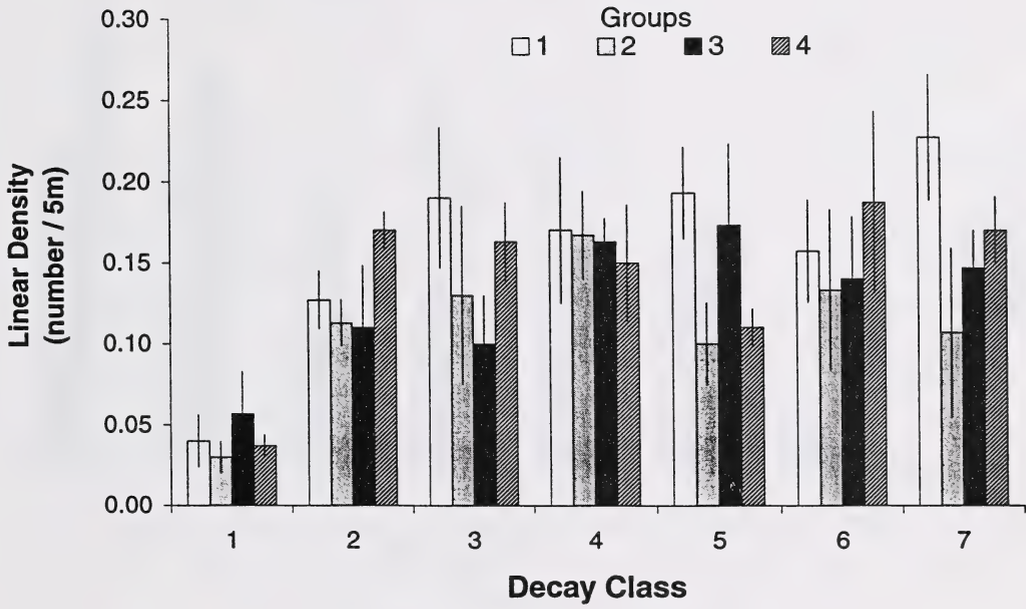


FIGURE 4.3. Average linear density (\pm SE) of large DWM in different decay stages among stand groups in the Grande Prairie study area.

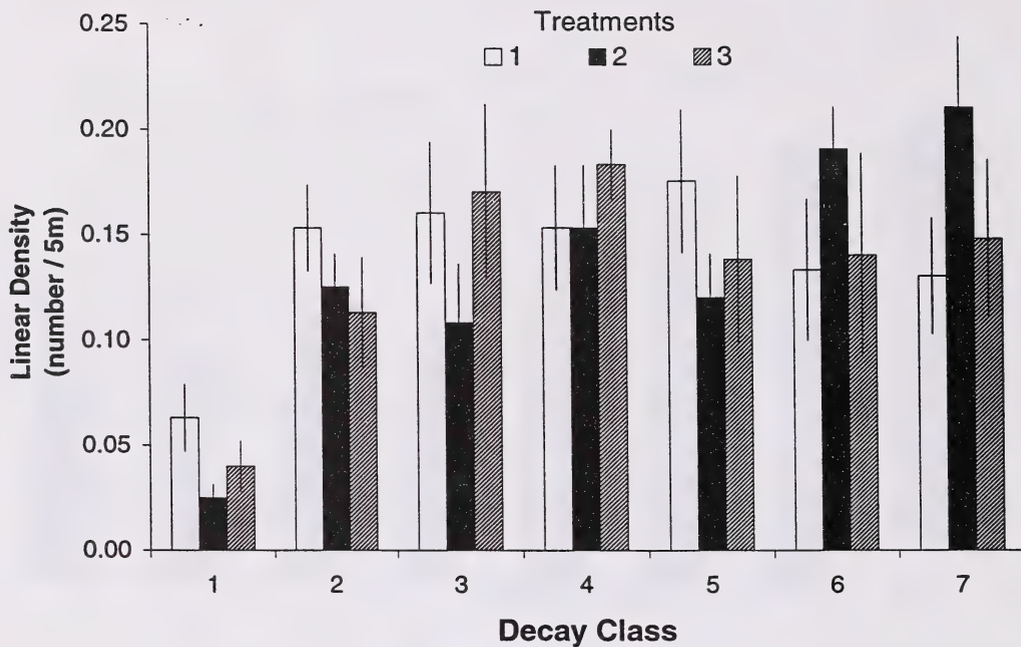


FIGURE 4.4. Average linear density (\pm SE) of large DWM in different decay stages among stand treatments in the Grande Prairie study area.

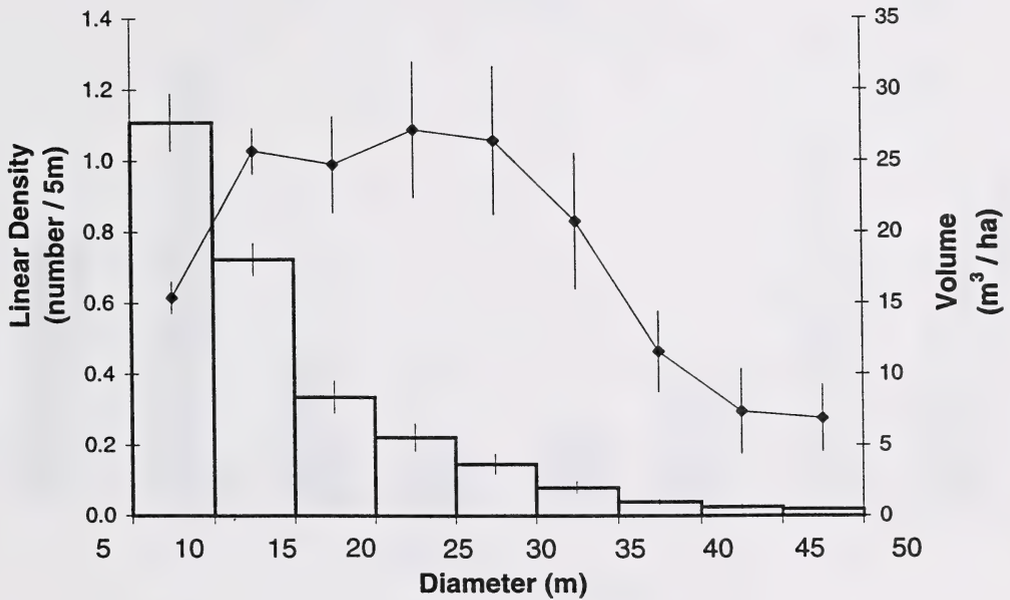


FIGURE 4.5. Size class distribution of the density (histogram) and volume (line) of DWM in the Whitecourt study area.

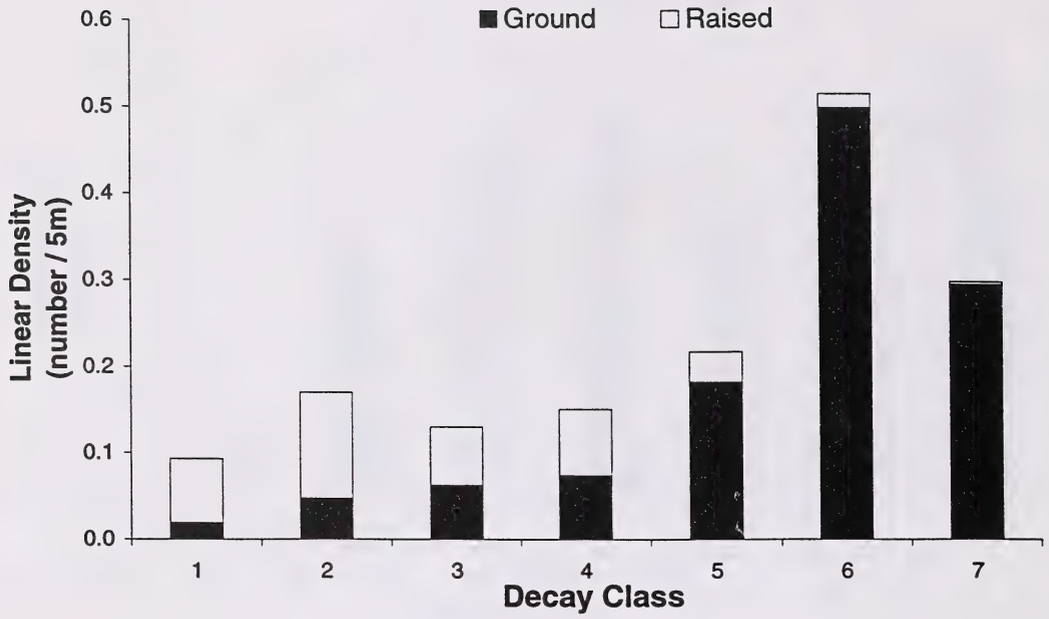


FIGURE 4.6. Decay class distribution of the density of DWM in the Whitecourt study area distinguishing between DWM raised off the ground and DWM lying on the ground.

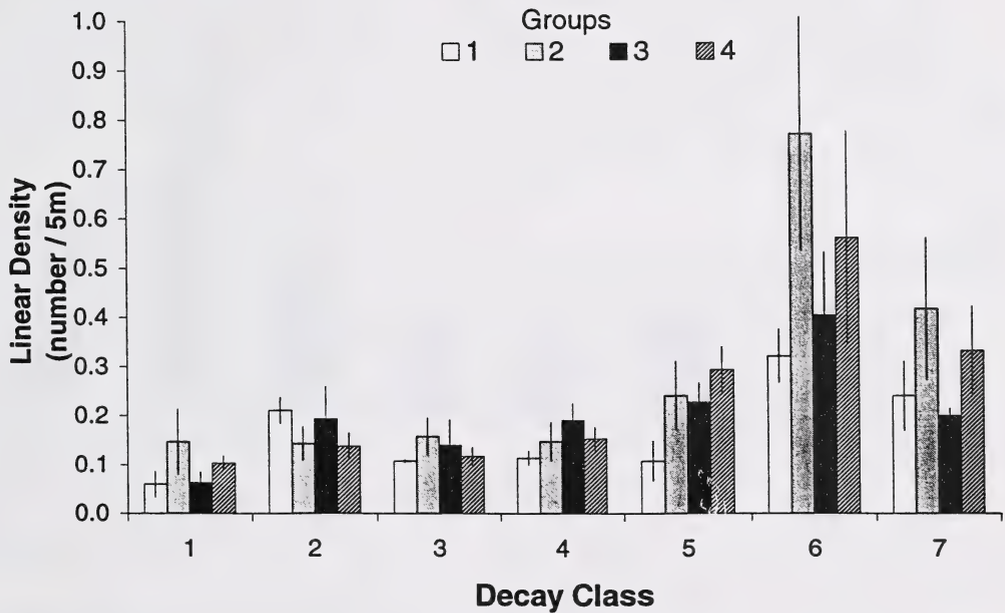


FIGURE 4.7. Average linear density (\pm SE) of large DWM in different decay stages among stand groups in the Whitecourt study area.

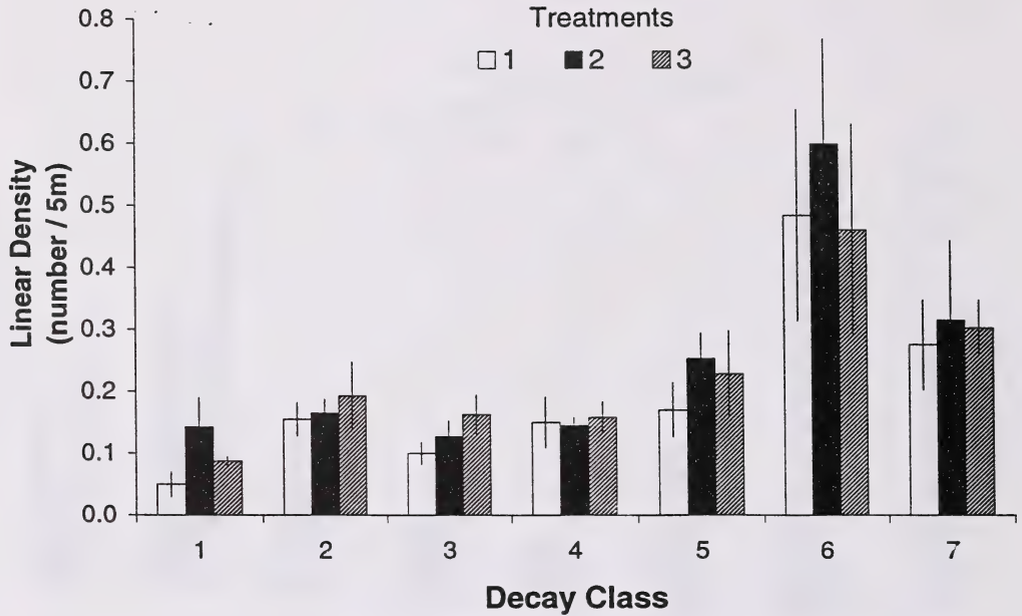


FIGURE 4.8. Average linear density (\pm SE) of large DWM in different decay classes among stand treatments in the Whitecourt study area.

1981, Abbott and Crossley 1982) returning nutrients to the soil.

The Grande Prairie study area had higher density, lower volume, and smaller sizes of DWM than other areas of >120 years-old aspen-dominated forest in Alberta (Lee et al. 1995, 1997). Densities, volumes and sizes of pine/spruce DWM could not be found in the literature.

The clumped spatial distribution of DWM within stands in both study areas suggests that leaving residual material in patches may be closer to natural spatial distributions (Lee et al. 1995, 1997) than if DWM was retained in a regular pattern. The clumped distribution of DWM in the Whitecourt study area is probably due to the nature of tree mortality; windfall usually causes clumped distributions of DWM (Harmon et al. 1986). In both study areas, DWM was more abundant near wet areas possibly because the roots of trees in wet areas facilitate windfall, especially for coniferous species (Harmon et al. 1986).

The decay class distribution of DWM in the Grande Prairie study area was very similar to that found in >120 year-old aspen dominated forests in other areas of Alberta (Crites and Dale 1995, in press; Lee et al. 1995, 1997). The onset of bark loss in year 1 for aspen logs (Miller 1983) results in DWM passing through decay class 1 quickly. Decay class distributions of DWM from pine/spruce dominated forest could not be found in the literature.

The difference in decay class distributions of DWM between the Grande Prairie and Whitecourt study areas suggests that aspen, pine, and spruce DWM may have different residence times within each decay class. The shady, cool understory of the pine/spruce forest promotes moss cover ($30.61\% \pm 1.28\text{SE}$, Chapter 5) compared to the sunny, dry understory of aspen forest ($3.27\% \pm 0.31\text{SE}$, Chapter 5; Söderström 1988a). In the pine/spruce forest, moss quickly covered intact and sound DWM therefore most DWM was classified as decay class 6 or 7. On the other hand, plants may colonized DWM at similar times in both study areas but the decomposition

rate of DWM following plant colonization (stages 6 -7) may be slower for pine/spruce DWM than aspen DWM. The time required for logs to lose half their wood density varies greatly between soft- and hardwoods: 57 years for pine (Pinus contorta, Fahey 1983), 63 years for balsam fir (Lambert et al. 1980), but only 10-14 years for aspen (Gosz 1980, Miller 1983, reviewed by Harmon et al. 1986). Hardwoods decompose faster than softwoods (MacMillan 1981, Harmon 1982) due to different wood structure and chemical composition (Wilcox 1973).

A second explanation for differences in distributions of DWM between the study areas is that there may have been a large, synchronized, mortality of pine and spruce trees in the Whitecourt study area within the time required for complete decomposition of DWM. Balsam fir logs can retain half of their volume for 100 years (Lambert et al. 1980). This cohort of DWM may be the killed trees from the last fire 105 years ago (unpublished data). If this is true, DWM dynamics not reached equilibrium before the area was harvested. Within a few decades the DWM in decay classes 6 and 7 may become indistinguishable, and the density and size distribution of DWM in the Whitecourt study area may be similar to estimates from the Grande Prairie study area which is probably at equilibrium.

Natural dynamics in old forests provide constant inputs of DWM resulting in a relatively even distribution of size classes and decay stages (Harmon et al. 1986; Söderström 1988b; Lee et al. 1995, 1997; Crites and Dale 1995, in press). On the other hand, managed stands have irregular inputs of DWM and an uneven distribution of DWM decay stages (Söderström 1988b). Clearcutting initially results in lower density and volume of large DWM (Lesica et al. 1991; Lee et al. 1995, 1997) and a pulse of undecayed small branches and twigs from felled and delimbbed trees (Abbott and Crossley 1982; Crites and Dale 1995, in press). At mid-rotation the small residual material is in advanced stages of decomposition with low input of trees from self-thinning and death of residual trees (Lee et al. 1995, 1997). In old age, large senescing trees will begin to consistently input DWM with even decay class and size distributions (Lee et al.

1995, 1997). Pulses of DWM may result in temporal heterogeneity in the abundance of species dependent on specific decay stages such as epixylics (Söderström 1988a,b; Lesica et al. 1991) and of species dependent on specific sizes of DWM such as microarthropods (Abbott and Crossley 1982, Söderström 1988a).

Many species of bryophytes, lichens, and some vascular plants are dependent on specific stages of DWM decomposition (Muhle and LeBlanc 1975; Crites and Dale 1995, in press). Söderström (1988b) suggested three requirements to maintain pre-harvest compositions of species dependent on DWM, such as lichens and bryophytes: DWM must be present in all stages of decomposition, DWM must have an even spatial distribution, and areas with DWM must be protected from drying out. A uniform distribution of DWM will allow DWM-dependent species to disperse into harvested areas (Lesica et al. 1991). Many of the rare lichens and bryophytes have limited dispersal capabilities and may require residual patches of DWM to act as dispersal centers (Söderström 1988). These same species need shaded areas with dependable humidity (Muhle and LeBlanc 1975, Larson 1984, Söderström 1988a, Lesica et al. 1991, Gustafsson et al. 1992).

The smaller, more abundant residual patches created in treatment 1 of this study will result in a more even spatial distribution of DWM and may facilitate the dispersal of DWM-dependent species into harvested areas. However, the smaller patches may have more variability in evaporation rates which suppresses nonvascular plants and DWM decomposition (Gustafsson et al. 1992, Söderström 1988a). Stands in treatments 2 and 3 with larger and wider spaced residual tree patches may retain more interior-forest habitat with a more natural humidity, temperature, shade, and natural decomposition rates of DWM. Natural decomposition rates may maintain temporally homogenous compositions of DWM-dependent species. On the other hand, the dispersal of species into the harvested areas may be restricted due to wider spaced patches of DWM.

Initially, an environment that allows DWM-dependent species to survive will be more important than an environment that facilitates dispersal. The ability of DWM-dependent species to disperse will become important once the canopy is established and the understory of harvested areas are buffered from sunlight and desiccation. Thus, the influence of DWM dispersion on biodiversity may shift from patch size to inter-patch distance as the harvested areas regenerate.

Pre-harvest characteristics of DWM provide a benchmark to which harvested areas can be compared. Stand groups were successful in accounting for pre-harvest variance of DWM, whereas there were few differences prior to harvest among randomly assigned treatments. Thus, any post-harvest differences in DWM among treatments may be attributed to treatment effects.

Future Research

Post-harvest characteristics of DWM will be surveyed in 1998 using the same methods as in 1997 with additional sampling to estimate DWM characteristics in residual tree patches. Additional transects will be sampled at random directions through the middle of residual tree patches for a total of 80 m (87.5 yd.) per stand.

If time permits, the wood density of large DWM in each decay class will be sampled so that DWM biomass can be calculated from volume estimates. Biomass is most frequently presented in the literature (Gosz 1980, Pastor and Bockheim 1984, Harmon et al. 1986). For each decay class, cross-sections ('cookies') of ten logs will be measured for volume (diameter and thickness of section), weighed, oven dried, and then re-weighed. This will also provide estimates of moisture content and biomass loss which in turn may affect bryophytes, lichens and other plants (Harmon et al. 1986).

In the winter of 1998-1999, the density, volume, spatial distribution, and decomposition of post-harvest DWM will be compared among treatment.

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CHAPTER 5. A SUMMARY OF UNDERSTORY VEGETATION IN THE BOREAL FOREST PRIOR TO HARVEST.

Christine Gray

Introduction

Due to funding constraints, only preliminary surveys were conducted for the understory vegetation during the 1997 field season. This report is a summary of that data. More detailed data collection, summary, and analyses will be conducted during 1999/2000.

Methods

In both study areas, vegetation was sampled at 5 out of the 10 possible 1 ha (2.47 acre) sites (every second site) within each stand. Each 1 ha site was divided into four equal quadrants (Fig 2.3). Two herb plots [0.5 x 0.5 m (1.6 x 1.6 ft.)] were systematically placed in all four quadrants (8 herb plots per site and 40 herb plots per stand).

For sampling purposes, vegetation was divided into two height strata. The herb stratum included all herbs regardless of height, and trees and shrubs that were <0.5 m (<1.6 ft.) in height. The shrub stratum included all shrubs ≥0.5 m (≥1.6 ft.) in height, and trees that were between 0.5 and 1.5 m (1.6–4.9 ft.) tall. Shrubs were divided into height classes of 0 = 0.5-1.39 m (1.6-4.6 ft.); 1 = 1.4-2.9 m (4.7-9.5 ft.); 2 = 3.0-4.9 m (9.6-16.1 ft.); 3 = 5.0-9.9 m (16.2-32.5 ft.); 4 = 10.0-19.9 m (32.6-65.6 ft.); and 5 = ≥20.0 m (≥ 65.6 ft.). In the Grande Prairie study area one shrub plot [2.0 x 2.0 m (6.6 x 6.6 ft.)] was systematically placed in two of the four randomly selected quadrants (2 shrub plots per site and 10 shrub plots per stand). In the Whitecourt study area a shrub plot was placed in all four quadrants (4 shrub plots per site and 20 shrub plots per stand). Herb plots were nested within shrub plots. Sampling was conducted from mid June through mid August 1997.

Percent of ground surface cover was visually estimated for litter, wood, moss, lichen, and grass, and percent of ground surface covered by a vertical projection of leaves and stems was

estimated for individual species of herbs, shrubs, and trees. In the Whitecourt study area percent of the ground surface cover by individual moss species was recorded as they were the dominant ground cover. Grasses and lichens were not identified to species at either study area. Unknown species were pressed and later identified at the University of Alberta herbarium. Nomenclature follows Moss (1992) for vascular species. For nonvascular species, nomenclature follows Vitt et al. (1988).

Results

Grande Prairie Study Area

Eighty-three species were found in the herbaceous strata, of which 55 were herbs, 25 were shrubs, and three were tree species (Table 5.1). Of the 25 shrub species found in the herb strata, 9 were not found in the shrub strata. Within the herb strata, much of the ground was covered by litter (43% cover) although herbs and shrubs were also abundant (39% and 18% cover respectively). Percent cover of grasses, mosses, and lichen were much lower (Table 5.1). Bunchberry was the most abundant herb (6% cover), with Prickly rose and Bracted honeysuckle the most abundant shrubs (5% cover for each) in the herb strata. Trees had low percent cover in both the herb and shrub strata (Table 5.1).

The shrub strata contained 21 species, of which 18 were shrubs and 3 were tree species (Table 5.1). Of these, River alder and Common Labrador tea were unique to the shrub stratum. The most abundant shrubs in the shrub stratum were Bracted honeysuckle (6% cover), Canadian buffaloberry (5% cover), and Prickly rose (5% cover). Trembling aspen was the most abundant tree species (1% cover) in the shrub stratum.

TABLE 5.1. Mean percent cover (\pm SE) of understory vegetation for the Grande Prairie study area during 1997. Individual species are grouped by strata level and family.

Common Name	Species Name	Species Code	Mean Percent Cover
Litter Total Cover			43.17 \pm 0.92
Wood Total Cover			5.31 \pm 0.45
Lichen Total Cover			0.01 \pm 0.00
Moss Total Cover			3.27 \pm 0.31
Grass Total Cover			4.81 \pm 0.26
Herb Total Cover			38.92 \pm 1.01
Lycopodiaceae			
Stiff club-moss	<i>Lycopodium annotinum</i>	LYC ANN	0.06 \pm 0.02
Ground-cedar	<i>Lycopodium complanatum</i>	LYC COM	0.04 \pm 0.04
Equisetaceae			
Common horsetail	<i>Equisetum arvense</i>	EQU ARV	0.23 \pm 0.04
Woodland horsetail	<i>Equisetum sylvaticum</i>	EQU SYL	0.04 \pm 0.01
Polypodiaceae			
Lady fern	<i>Athyrium filix-femina</i>	ATH FIL	0.01 \pm 0.01
Oak fern	<i>Gymnocarpium dryopteris</i>	GYM DRY	0.01 \pm 0.01
unknown fern		UNK FER	0.08 \pm 0.08 ¹
Liliaceae			
Fairybells	<i>Disporum trachycarpum</i>	DIS TRA	0.01 \pm 0.01
Wild lily-of-the-valley	<i>Maianthemum canadense</i>	MAI CAN	0.67 \pm 0.08
Star-flowered false solomon's-seal	<i>Smilacina stellata</i>	SMI STE	0.04 \pm 0.03
Three-leaved false solomon's-seal	<i>Smilacina trifolia</i>	SMI TRI	0.02 \pm 0.01
unknown lily	<i>Smilacina</i> spp.	SMI SPP	0.02 \pm 0.02
Orchidaceae			
Rattlesnake-plantain	<i>Goodyera oblongifolia</i>	GOO OBL	0.01 \pm 0.01
Blunt-leaved bog-orchid	<i>Habenaria obtusata</i>	HAB OBT	0.01 \pm 0.01
Ranunculaceae			
Red and white baneberry	<i>Actaea rubra</i>	ACT RUB	0.32 \pm 0.06
Veiny meadow rue	<i>Thalictrum venulosum</i>	THA VEN	0.01 \pm 0.01
Saxifragaceae			
Bishop's cap	<i>Mitella nuda</i>	MIT NUD	0.78 \pm 0.08
Rosaceae			
Wild strawberry	<i>Fragaria virginiana</i>	FRA VIR	2.47 \pm 0.13
Dewberry	<i>Rubus pubescens</i>	RUB PUB	2.17 \pm 0.15
Leguminosae			
Creamy peavine	<i>Lathyrus ochroleucus</i>	LAT OCH	3.01 \pm 0.18
American vetch	<i>Vicia americana</i>	VIC AME	1.20 \pm 0.11
Violaceae			
Canada violet	<i>Viola canadensis</i>	VIO CAN	0.26 \pm 0.07
Kidney-leaved violet	<i>Viola renifolia</i>	VIO REN	0.53 \pm 0.07
Onagraceae			
Fireweed	<i>Epilobium angustifolium</i>	EPI ANG	2.65 \pm 0.19

¹ Sum of two unidentified fern species.

TABLE 5.1 cont.

Common Name	Species Name	Species Code	Mean Percent Cover
Araliaceae			
Wild sarsaparilla	<i>Aralia nudicaulis</i>	ARA NUD	1.83 ± 0.25
Umbelliferae			
Cow-parsnip	<i>Heracleum lanatum</i>	HER LAN	0.23 ± 0.09
Cornaceae			
Bunchberry	<i>Cornus canadensis</i>	COR CAN	6.11 ± 0.32
Pyrolaceae			
One-sided wintergreen	<i>Orthilia secunda</i>	ORT SEC	0.04 ± 0.03
Common pink wintergreen	<i>Pyrola asarifolia</i>	PYR ASA	0.94 ± 0.10
Northern starflower	<i>Trientalis borealis</i>	TRI BOR	0.02 ± 0.01
Gentianaceae			
Spurred gentian	<i>Halenia deflexa</i>	HAL DEF	0.06 ± 0.02
Boraginaceae			
Tall lungwort	<i>Mertensia paniculata</i>	MER PAN	1.61 ± 0.17
Labiatae			
Marsh skullcap	<i>Scutellaria galericulata</i>	SCU GAL	0.01 ± 0.01
Scrophulariaceae			
Common red paintbrush	<i>Castilleja miniata</i>	CAS MIN	0.52 ± 0.09
Rubiaceae			
Northern bedstraw	<i>Galium boreale</i>	GAL BOR	1.36 ± 0.09
Sweet-scented bedstraw	<i>Galium triflorum</i>	GAL TRI	0.06 ± 0.01
Caprifoliaceae			
Twinflower	<i>Linnaea borealis</i>	LIN BOR	2.89 ± 0.17
Compositae			
Common yarrow	<i>Achillia millefolium</i>	ACH MIL	0.26 ± 0.04
Heart-leaved arnica	<i>Arnica cordifolia</i>	ARN COR	0.36 ± 0.07
Lindley's aster	<i>Aster ciliolatus</i>	AST CIL	0.90 ± 0.13
Showy aster	<i>Aster conspicuus</i>	AST CON	3.37 ± 0.29
Purple-stemmed aster	<i>Aster puniceus</i>	AST PUN	0.03 ± 0.02
Narrow-leaved hawkweed	<i>Hieracium umbellatum</i>	HIE UMB	0.96 ± 0.17
Palmate-leaved coltsfoot	<i>Petasites palmatus</i>	PET PAL	2.38 ± 0.16
Arrow-leaved coltsfoot	<i>Petasites sagittatus</i>	PET SAG	0.14 ± 0.08
Canadian butterweed	<i>Senecio pauperculus</i>	SEN PAU	0.01 ± 0.01
Canada goldenrod	<i>Solidago canadensis</i>	SOL CAN	0.09 ± 0.04
Late goldenrod	<i>Solidago gigantea</i>	SOL GIG	0.03 ± 0.02
Common dandelion	<i>Taraxacum officinale</i>	TAR OFF	0.01 ± 0.01
unknowns		UNK	0.09 ± 0.07 ²
Shrub Total Cover (< 0.5 m)			18.21 ± 0.55
Salicaceae			
willow	<i>Salix</i> spp.	SAL SPP	0.10 ± 0.04
Betulaceae			
Green alder	<i>Alnus crispa</i>	ALN CRI	0.03 ± 0.02
Beaked hazelnut	<i>Corylus cornuta</i>	COR COR	0.07 ± 0.05
Grossulariaceae			
Skunk currant	<i>Ribes glandulosum</i>	RIB GLA	0.01 ± 0.01

² Sum of five unidentified herb species.

TABLE 5.1 cont.

Common Name	Species Name	Species Code	Mean Percent Cover
Northern black currant	<i>Ribes hudsonianum</i>	RIB HUD	0.06 ± 0.03
Black gooseberry	<i>Ribes lacustre</i>	RIB LAC	0.04 ± 0.02
Northern gooseberry	<i>Ribes oxycanthoides</i>	RIB OXY	0.16 ± 0.03
Wild red currant	<i>Ribes triste</i>	RIB TRI	0.36 ± 0.08
unknown currant	<i>Ribes spp</i>	RIB SPP	0.10 ± 0.04
Rosaceae			
Saskatoon	<i>Amelanchier alnifolia</i>	AME ALN	0.29 ± 0.08
Pin cherry	<i>Prunus pensylvanica</i>	PRU PEN	0.03 ± 0.02
Choke cherry	<i>Prunus virginiana</i>	PRU VIR	0.01 ± 0.01
Prickly rose	<i>Rosa acicularis</i>	ROS ACI	5.33 ± 0.24
Wild red raspberry	<i>Rubus idaeus</i>	RUB IDA	0.70 ± 0.10
Narrow-leaved meadowsweet	<i>Spiraea alba</i>	SPI BET	1.09 ± 0.13
Elaeagnaceae			
Canada buffaloberry	<i>Shepherdia canadensis</i>	SHE CAN	0.62 ± 0.09
Cornaceae			
Red-osier dogwood	<i>Cornus stolonifera</i>	COR STO	0.12 ± 0.05
Ericaceae			
Common bearberry	<i>Arctostaphylos uva-ursi</i>	ARC UVA	0.01 ± 0.01
Common labrador tea	<i>Ledum groenlandicum</i>	LED GRO	0.05 ± 0.03
Velvet-leaved blueberry	<i>Vaccinium myrtilloides</i>	VAC MYR	0.78 ± 0.14
Bog cranberry	<i>Vaccinium vitis-idaea</i>	VAC VIT	0.04 ± 0.02
Caprifoliaceae			
Twining honeysuckle	<i>Lonicera dioica</i>	LON DIO	0.28 ± 0.06
Bracted honeysuckle	<i>Lonicera involucrata</i>	LON INV	5.22 ± 0.36
Common snowberry	<i>Symphoricarpos albus</i>	SYM ALB	0.04 ± 0.02
Low bush-cranberry	<i>Viburnum edule</i>	VIB EDU	2.67 ± 0.29
Shrub Total Cover (> 0.5 m)			29.36 ± 1.95
Salicaceae			
willow	<i>Salix spp.</i>	SAL SPP	3.48 ± 0.51
Betulaceae			
Green alder	<i>Alnus crispa</i>	ALN CRI	1.64 ± 0.62
River alder	<i>Alnus tenuifolia</i>	ALN TEN	1.43 ± 0.83
Beaked hazelnut	<i>Corylus cornuta</i>	COR COR	0.18 ± 0.13
Grossulariaceae			
Northern gooseberry	<i>Ribes oxycanthoides</i>	RIB OXY	0.07 ± 0.04
unknown currant	<i>Ribes spp.</i>	RIB SPP	0.02 ± 0.01
Rosaceae			
Saskatoon	<i>Amelanchier alnifolia</i>	AME ALN	0.94 ± 0.27
Pin cherry	<i>Prunus pensylvanica</i>	PRU PEN	0.41 ± 0.16
unknown cherry	<i>Prunus spp.</i>	PRU SPP	0.01 ± 0.01
Choke cherry	<i>Prunus virginiana</i>	PRU VIR	0.02 ± 0.01
Prickly rose	<i>Rosa acicularis</i>	ROS ACI	5.02 ± 0.58
Wild red raspberry	<i>Rubus idaeus</i>	RUB IDA	0.14 ± 0.07
Elaeagnaceae			
Canada buffaloberry	<i>Shepherdia canadensis</i>	SHE CAN	5.24 ± 1.01
Cornaceae			
Red-osier dogwood	<i>Cornus stolonifera</i>	COR STO	0.05 ± 0.04
Ericaceae			
Common labrador tea	<i>Ledum groenlandicum</i>	LED GRO	0.08 ± 0.08

TABLE 5.1 cont.

Common Name	Species Name	Species Code	Mean Percent Cover
Caprifoliaceae			
Twining honeysuckle	<i>Lonicera dioica</i>	LON DIO	0.09 ± 0.06
Bracted honeysuckle	<i>Lonicera involucrata</i>	LON INV	6.27 ± 1.14
Low bush-cranberry	<i>Viburnum edule</i>	VIB EDU	4.28 ± 0.75
Tree Total Cover (< 0.5 m)			0.07 ± 0.04
Pinaceae			
White spruce	<i>Picea glauca</i>	PIC GLA	0.01 ± 0.01
Salicaceae			
Trembling aspen	<i>Populus tremuloides</i>	POP TRE	0.05 ± 0.04
Betulaceae			
Paper birch	<i>Betula papyrifera</i>	BET PAP	0.01 ± 0.01
Tree Total Cover (0.5 – 1.5 m)			0.87 ± 0.31
Pinaceae			
White spruce	<i>Picea glauca</i>	PIC GLA	0.04 ± 0.04
Salicaceae			
Trembling aspen	<i>Populus tremuloides</i>	POP TRE	0.74 ± 0.28
Betulaceae			
Paper birch	<i>Betula papyrifera</i>	BET PAP	0.08 ± 0.08

Whitecourt Study Area

Seventy-six vascular species (including 53 herbs, 19 shrubs, and 4 tree species) and 22 nonvascular species (19 mosses and 3 lichens) were found in the herb strata (Table 5.2). Within the herb strata moss, litter and herbs were equally abundant (31%, 30% and 28% cover respectively). Knight's plume (17% cover) and Big red stem (15% cover) were the most abundant moss species. Bunchberry was the most abundant herb (4% cover) with Common Labrador tea (3% cover) and Velvet-leaved blueberry (2% cover) the most abundant shrubs in the herb strata. Balsam fir was the most abundant tree species in both the herb and shrub strata (Table 5.2).

The shrub strata contained a total of 20 species, of which 14 were shrubs and 6 were trees (Table 5.2). Five shrub species found in the herb strata were not found in the shrub strata. Devil's club (2% cover) was the most abundant shrub species in the shrub strata. Balsam poplar and Trembling aspen were the only tree species found in the shrub strata and not the herb strata.

Discussion

The Grande Prairie study area had a well developed herb and shrub strata; shrubs were tall and abundant throughout the study area and the herbs covered a majority of the ground space. This abundance may have been due to the deciduous canopy allowing light to penetrate through to the herb and shrub strata.

The Whitecourt study area had a high density of coniferous trees (Chapter 4). The herb and shrub strata were less developed in this area, possibly due to shading. An abundant moss community was present, possibly as a result of the shaded and damp conditions.

Due to funding constraints only preliminary surveys and summary were conducted for the preharvest understory vegetation.

Future research

Surveys and analysis of post harvest understory vegetation will take place during 1999/2000. Methods will be similar to those presented above but to obtain adequate data, all 1-ha sites will be surveyed (ie. twice the intensity of the surveys during 1997). Twenty and ten supplemental herb and shrub plots respectively will be surveyed within each stand. Half of the supplemental plots will be within residual tree clumps and the other half >100 m (>109 yd.) from residual tree plots.

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TABLE 5.2. Mean percent cover (\pm SE) of understory vegetation for the Whitecourt study area during 1997. Individual species are grouped by strata level and family.

Common Name	Species Name	Species Code	Mean Percent Cover
Litter Total Cover			29.56 \pm 1.08
Wood Total Cover			4.76 \pm 0.36
Lichen Total Cover			0.08 \pm 0.04
club lichen	<i>Cladonia</i> spp.	CLA SPP	0.01 \pm 0.01
Studded leather lichen	<i>Peltigera aphthosa</i>	PEL APH	0.01 \pm 0.01
Finger felt lichen	<i>Peligeria neopolydactyla</i>	PEL NEO	0.04 \pm 0.03
Moss Total Cover			30.61 \pm 1.28
sphagnum	<i>Sphagnum</i> spp.	SPH SPP	3.74 \pm 0.68
Common hair-cap	<i>Polytrichum commune</i>	POL COM	0.39 \pm 0.12
Juniper hair-cap	<i>Polytrichum juniperinum</i>	POL JUN	0.30 \pm 0.08
Slender hair-cap	<i>Polytrichum strictum</i>	POL STR	0.49 \pm 0.12
Common northern lantern moss	<i>Cinclidium stygium</i>	CIN STY	1.46 \pm 0.20
Knight's plume	<i>Ptilium crista-castrensis</i>	PTI CRI	17.46 \pm 1.03
Big red stem	<i>Pleurozium schreberi</i>	PLE SCH	14.75 \pm 0.91
Stair-step moss	<i>Hylocomium splendens</i>	HYL SPL	0.80 \pm 0.14
Broom moss	<i>Dicranum scoparium</i>	DIC SCO	0.27 \pm 0.06
unknown mosses		MOSS	0.56 \pm 0.39 ³
Grass Total Cover			3.10 \pm 0.36
Herb Total Cover			28.24 \pm 0.95
Lycopodiaceae			
Stiff club-moss	<i>Lycopodium annotinum</i>	LYC ANN	2.61 \pm 0.21
Ground-cedar	<i>Lycopodium complanatum</i>	LYC COM	0.03 \pm 0.02
Equisetaceae			
Common horsetail	<i>Equisetum arvense</i>	EQU ARV	0.52 \pm 0.08
Woodland horsetail	<i>Equisetum sylvaticum</i>	EQU SYL	0.76 \pm 0.13
Polypodiaceae			
Lady fern	<i>Athyrium filix-femina</i>	ATH FIL	1.91 \pm 0.49
Oak fern	<i>Gymnocarpium dryopteris</i>	GYM DRY	2.53 \pm 0.21
Cyperaceae			
unknown sedge	<i>Carex</i> spp.	SED SPP	0.01 \pm 0.01
Liliaceae			
Fairybells	<i>Disporum trachycarpum</i>	DIS TRA	0.48 \pm 0.08
Wild lily-of-the-valley	<i>Maianthemum canadense</i>	MAI CAN	0.22 \pm 0.04
False solomon's-seal	<i>Smilacina racemosa</i>	SMI RAC	0.01 \pm 0.01
Star-flowered false solomon's-seal	<i>Smilacina stellata</i>	SMI STE	0.03 \pm 0.02
Three-leaved false solomon's-seal	<i>Smilacina trifolia</i>	SMI TRI	0.05 \pm 0.02
unknown lily	<i>Smilacina</i> spp.	SMI SPP	0.02 \pm 0.01
Indian hellebore	<i>Veratrum viride</i>	VER VIR	0.04 \pm 0.03
Orchidaceae			
Heart-leaved twayblade	<i>Listera cordata</i>	LIS COR	0.01 \pm 0.00

³ Sum of ten unidentified moss species.

TABLE 5.2 cont.

Common Name	Species Name	Species Code	Mean Percent Cover
Urticaceae			
Stinging nettle	<i>Urtica dioica</i>	URT DIO	0.05 ± 0.03
Ranunculaceae			
Red and white baneberry	<i>Actaea rubra</i>	ACT RUB	0.05 ± 0.03
Goldthread	<i>Coptis trifolia</i>	COP TRI	0.16 ± 0.04
Tall larkspur	<i>Delphinium glaucum</i>	DEL GLA	0.01 ± 0.01
Veiny meadow rue	<i>Thalictrum venulosum</i>	THA VEN	0.03 ± 0.01
Saxifragaceae			
Bishop's cap	<i>Mitella nuda</i>	MIT NUD	0.58 ± 0.07
Three-leaved foamflower	<i>Tiarella trifoliata</i>	TIA TRI	1.38 ± 0.16
Rosaceae			
Wild strawberry	<i>Fragaria virginiana</i>	FRA VIR	0.02 ± 0.01
Purple avens	<i>Geum rivale</i>	GEU RIV	0.02 ± 0.01
Cloudberry	<i>Rubus chamaemorus</i>	RUB CHA	0.30 ± 0.09
Five-leaved bramble	<i>Rubus pedatus</i>	RUB PED	2.90 ± 0.22
Dewberry	<i>Rubus pubescens</i>	RUB PUB	1.91 ± 0.15
Leguminosae			
Creamy peavine	<i>Lathyrus ochroleucus</i>	LAT OCH	0.01 ± 0.01
Balsaminaceae			
Spotted touch-me-not	<i>Impatiens capensis</i>	IMP CAP	0.01 ± 0.01
Violaceae			
Kidney-leaved violet	<i>Viola renifolia</i>	VIO REN	0.19 ± 0.04
Onagraceae			
Small enchanter's nightshade	<i>Circaea alpina</i>	CIR ALP	0.01 ± 0.01
Fireweed	<i>Epilobium angustifolium</i>	EPI ANG	0.93 ± 0.11
Araliaceae			
Wild sarsaparilla	<i>Aralia nudicaulis</i>	ARA NUD	2.36 ± 0.25
Umbelliferae			
Cow-parsnip	<i>Heracleum lanatum</i>	HER LAN	0.06 ± 0.06
Cornaceae			
Bunchberry	<i>Cornus canadensis</i>	COR CAN	4.34 ± 0.20
Pyrolaceae			
One-sided wintergreen	<i>Orthilia secunda</i>	ORT SEC	0.01 ± 0.00
Common pink wintergreen	<i>Pyrola asarifolia</i>	PYR ASA	0.29 ± 0.06
Green wintergreen	<i>Pyrola virens</i>	PYR VIR	0.05 ± 0.02
Boraginaceae			
Tall lungwort	<i>Mertensia paniculata</i>	MER PAN	0.29 ± 0.07
Scrophulariaceae			
Common red paintbrush	<i>Castilleja miniata</i>	CAS MIN	0.01 ± 0.01
Rubiaceae			
Northern bedstraw	<i>Galium boreale</i>	GAL BOR	0.01 ± 0.00
Sweet-scented bedstraw	<i>Galium triflorum</i>	GAL TRI	0.06 ± 0.02
Caprifoliaceae			
Twinflower	<i>Linnaea borealis</i>	LIN BOR	1.94 ± 0.15
Compositae			
Heart-leaved arnica	<i>Arnica cordifolia</i>	ARN COR	0.29 ± 0.06
Narrow-leaved hawkweed	<i>Hieracium umbellatum</i>	HIE UMB	0.03 ± 0.03
Palmate-leaved coltsfoot	<i>Petasites palmatus</i>	PET PAL	0.65 ± 0.08
Arrow-leaved groundsel	<i>Senecio triangularis</i>	SEN TRI	0.01 ± 0.01
Late goldenrod	<i>Solidago gigantea</i>	SOL GIG	0.03 ± 0.03
Common dandelion	<i>Taraxacum officinale</i>	TAR OFF	0.01 ± 0.01

TABLE 5.2 cont.

Common Name	Species Name	Species Code	Mean Percent Cover
unknowns		UNK	0.06 ± 0.06 ⁴
Shrub Total Cover (< 0.5 m)			10.63 ± 0.52
Salicaceae			
willow	<i>Salix</i> spp.	SAL SPP	0.01 ± 0.01
Betulaceae			
Green alder	<i>Alnus crispa</i>	ALN CRI	0.04 ± 0.02
River alder	<i>Alnus tenuifolia</i>	ALN TEN	0.01 ± 0.01
Grossulariaceae			
Northern black currant	<i>Ribes hudsonianum</i>	RIB HUD	0.02 ± 0.02
Northern gooseberry	<i>Ribes oxycanthoides</i>	RIB OXY	0.47 ± 0.09
Wild red currant	<i>Ribes triste</i>	RIB TRI	0.11 ± 0.04
unknown currant	<i>Ribes</i> spp.	RIB SPP	0.06 ± 0.02
Rosaceae			
Saskatoon	<i>Amelanchier alnifolia</i>	AME ALN	0.09 ± 0.04
Prickly rose	<i>Rosa acicularis</i>	ROS ACI	0.95 ± 0.11
Wild red raspberry	<i>Rubus idaeus</i>	RUB IDA	0.33 ± 0.09
Western mountain ash	<i>Sorbus scopulina</i>	SOR SCO	0.01 ± 0.01
Araliaceae			
Devil's club	<i>Oplopanax horridum</i>	OPL HOR	0.91 ± 0.26
Ericaceae			
Creeping snowberry	<i>Gaultheria hispidula</i>	GAU HIS	0.56 ± 0.09
Common Labrador tea	<i>Ledum groenlandicum</i>	LED GRO	2.55 ± 0.28
Black huckleberry	<i>Vaccinium membranaceum</i>	VAC MEM	0.16 ± 0.05
Velvet-leaved blueberry	<i>Vaccinium myrtilloides</i>	VAC MYR	2.12 ± 0.21
Bog cranberry	<i>Vaccinium vitis-idaea</i>	VAC VIT	0.92 ± 0.12
Caprifoliaceae			
Bracted honeysuckle	<i>Lonicera involucrata</i>	LON INV	0.60 ± 0.1
Low bush-cranberry	<i>Viburnum edule</i>	VIB EDU	0.72 ± 0.11
Shrub Total Cover (> 0.5 m)			7.00 ± 0.72
Salicaceae			
willow	<i>Salix</i> spp.	SAL SPP	0.32 ± 0.09
Betulaceae			
Green alder	<i>Alnus crispa</i>	ALN CRI	1.05 ± 0.34
River alder	<i>Alnus tenuifolia</i>	ALN TEN	0.17 ± 0.11
Grossulariaceae			
Northern black currant	<i>Ribes hudsonianum</i>	RIB HUD	0.04 ± 0.04
Northern gooseberry	<i>Ribes oxycanthoides</i>	RIB OXY	0.14 ± 0.06
Wild red currant	<i>Ribes triste</i>	RIB TRI	0.01 ± 0.01
unknown currant	<i>Ribes</i> spp.	RIB SPP	0.03 ± 0.02
Rosaceae			
Saskatoon	<i>Amelanchier alnifolia</i>	AME ALN	0.01 ± 0.01
Prickly rose	<i>Rosa acicularis</i>	ROS ACI	1.20 ± 0.18
Wild red raspberry	<i>Rubus idaeus</i>	RUB IDA	0.16 ± 0.13
Araliaceae			
Devil's club	<i>Oplopanax horridum</i>	OPL HOR	1.57 ± 0.44
Ericaceae			
Common Labrador tea	<i>Ledum groenlandicum</i>	LED GRO	1.06 ± 0.24

⁴ Sum of four identified herb species.

TABLE 5.2 cont.

Common Name	Species Name	Species Code	Mean Percent Cover
Caprifoliaceae			
Bracted honeysuckle	<i>Lonicera involucrata</i>	LON INV	0.66 ± 0.19
Low bush-cranberry	<i>Viburnum edule</i>	VIB EDU	0.59 ± 0.13
Tree Total Cover (< 0.5 m)			0.49 ± 0.10
Pinaceae			
Balsam fir	<i>Abies balsamea</i>	ABI BAL	0.40 ± 0.09
White spruce	<i>Picea glauca</i>	PIC GLA	0.01 ± 0.01
Black spruce	<i>Picea mariana</i>	PIC MAR	0.02 ± 0.01
Betulaceae			
Paper birch	<i>Betula papyrifera</i>	BET PAP	0.07 ± 0.03
Tree Total Cover (0.5 – 1.5 m)			1.78 ± 0.27
Pinaceae			
Balsam fir	<i>Abies balsamea</i>	ABI BAL	1.03 ± 0.23
White spruce	<i>Picea glauca</i>	PIC GLA	0.01 ± 0.01
Black spruce	<i>Picea mariana</i>	PIC MAR	0.16 ± 0.06
Salicaceae			
Balsam poplar	<i>Populus balsamifera</i>	POP BAL	0.02 ± 0.02
Trembling aspen	<i>Populus tremuloides</i>	POP TRE	0.03 ± 0.02
Betulaceae			
Paper birch	<i>Betula papyrifera</i>	BET PAP	0.53 ± 0.13

CHAPTER 6. ABUNDANCE AND SPECIES COMPOSITION OF CARABID BEETLES IN RELATION TO THE DISPERSION OF RESIDUAL TREES FOLLOWING HARVEST

Troy C. Sorensen

Introduction

Although arthropods compose a large proportion of the biodiversity in most ecosystems; the influence of forestry practices on these taxa is often overlooked. Recent studies have suggested that ground beetles (Coleoptera: Carabidae) may be sensitive to forestry practices and serve as ecological indicators (Thiele 1977, Butterfield et al. 1995). Arthropod communities are influenced by forestry practices because they are sensitive to temperature (Ericson 1979), moisture (Niemela et al. 1992, Niemela and Spence 1994), ground litter (Greenslade 1964), and understory and canopy cover (Niemela and Spence 1994). Furthermore, the dispersion of suitable habitat patches will influence community composition because dispersal is energetically expensive (Atkins 1969) and species differ in their ability to disperse (Thiele 1977).

Boreal communities of carabid beetles have been studied in relation to seasonal changes (Niemela et al. 1992, Digweed et al. 1995), edge effects (Halme and Niemela 1993, Spence et al. 1996), fragmentation (Halme and Niemela 1993), post-harvest regeneration (Spence et al. 1996), and post-fire regeneration (Holliday 1991, 1992; Spence et al. 1996). The main objective of this project is to determine the influence of residual tree dispersion on the species composition and abundance of carabid beetles in aspen-dominated and pine-spruce-dominated boreal forests of western Alberta. Specifically, I will examine whether the carabid community within and surrounding residual patches varies among treatments (see Chapter 2 for a description of the treatments), and (2) whether differences among treatments can be explained by microhabitat preferences or dispersal capabilities of specific species.

Future Research

Due to funding constraints, carabid beetles were not surveyed during 1997. Surveys from four stands of unharvested forest within each study area will serve as benchmarks for post-harvest data.

Carabid beetles will be surveyed with pitfall traps because the traps are effective (Obertel 1971, Luff 1975, Niemela et al. 1986, Spence and Niemela 1994, Digweed et al. 1995), and results will be comparable to previous studies (Spence et al. 1996). Furthermore, pitfall traps sample a variety of other taxa including ants and spiders that could be identified and analyzed at a later date if funding becomes available. Pitfall traps have been criticized because they sample activity rather than abundance and therefore catches over short durations are dependent on weather and the period of life cycle (Ericson 1979). However, if traps are open for weeks at a time, and sampling occurs at multiple times during the snow-free portion of a year, the effect of weather will be negligible and catches are correlated with density (Baars 1979, Ericson 1979).

Pitfall traps will consist of a removable 500ml (0.53 qt.) plastic container set inside a larger permanent container [1L (1.06 qt.)] so that repeated sampling will not disturb the soil and litter around the trap (Spence and Niemela 1994, Digweed et al. 1995). Each trap will be partially filled 2-3 cm (0.8-1.2 in.) with undiluted ethylene glycol to increase trap effectiveness and preserve the specimens (Holopainen 1992). A 15 x 15 cm (5.9 x 5.9 in.) plywood cover will be suspended with nails over each trap to prevent rain, leaves, and litter from falling into the traps (Spence and Niemela 1994). Trapping will occur during the first two weeks of each month from early May to the middle of September for a total of 5 trapping

sessions. Traps will be left open for 2 weeks at a time with a 2 week recuperation period between trapping sessions (traps closed) to prevent population depletion.

Traps will be placed along radial transects from the center of residual patches in harvested stands, and along similar transects in the unharvested stands (Fig. 6.1). Transects will begin at the center of the patch and end at the half-way point between patches in the harvested areas. The number of traps in each stand will be similar; however the number of traps used to sample each patch, and the number of patches sampled within each stand will depend on the treatment (Fig. 6.1). Single traps will be placed no less than 20m apart to insure independent catches (see Digweed et al. 1995). Previous studies caught fewer (67% less) beetles in pine-dominated than aspen-dominated forests (Spence et al. 1996); to compensate, the number of traps in each stand will be higher in the Whitecourt study area (24 traps) than the Grande Prairie area (16-18 traps). Carabid beetles will be identified to species according to Lindroth (1961-1969). Reference collections are available at the Strickland Museum, University of Alberta, and at the Northern Forestry Centre, Canadian Forest Service, Edmonton.

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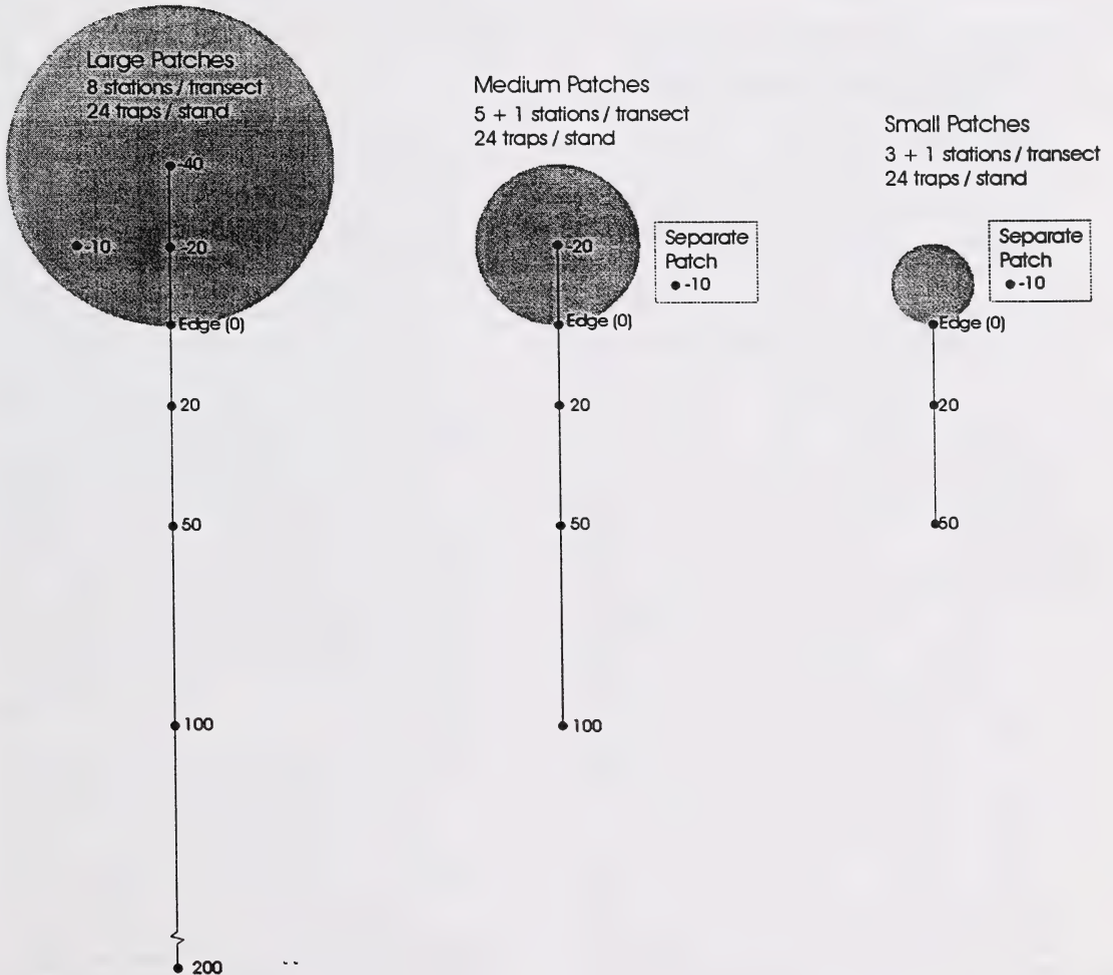


Figure 6.1. Locations of pitfall traps for the three patch sizes in the Whitecourt Area. Trapping design is similar for the Grande Prairie study area with the following distances from patch edges: -50 m, -20 m, -10 m, 0 m, 20 m, 50 m, 120 m, and 250 m (- 55 yd., - 22 yd., - 11 yd., 0 yd., 22 yd., 55 yd., 131 yd., and 273 yd.). Traps must be at least 20 m (22 yd.) apart to insure statistical independence. Traps will be placed along transects in unharvested forest with the same spacing as the traps in large patches.

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CHAPTER 7. VARIATION IN BIRD COMMUNITIES IN RELATION TO THE DISPERSION OF STANDING LIVE TREES AND SNAGS

Jim Schieck

Introduction

As a result of government regulations (e.g. British Columbia Ministry of Forests 1995) and exploratory adaptive management (e.g., Alberta-Pacific Forest Industries Inc. 1996) many forest companies have modified harvesting strategies by leaving standing trees and snags in harvested areas. By providing more complex habitat than is found after conventional clear-cut harvest, these residual trees may make the harvest areas more useable by native birds (Schieck et al. 1995, Norton and Hannon 1997). In addition, by providing inputs of large snags and downed logs throughout succession, residual trees may promote recovery of biota in harvested areas (Hobson and Schieck submitted). However, there is little information available to determine the number of residual trees and their spacing that will maximize recovery of native birds. As a result, forest managers are subjectively deciding those patterns (Alberta-Pacific Forest Industries Inc. 1996).

One possible template to follow when leaving standing trees is a natural disturbance model; all else being equal, making harvest areas as similar as possible to natural disturbances probably will be beneficial to native biota since they evolved in an ecosystem with natural disturbances (Urban et al. 1987). Wildfire, disease, and wind-throw events all create natural disturbances in forests, but the dominant natural disturbance in the boreal forests of North America is wildfire (Eberhart and Woodard 1987, Hunter 1993). Thus, wildfire is the most realistic template on which to pattern harvest in the boreal forest. Following wildfires, there are groups of unburned trees (fire skips) embedded within the burnt-over areas (Eberhart and Woodard 1987). Fire skips may retain some of the structures and micro-habitats that were present, and may act as refugia for some of the birds that lived in the forest prior to the disturbance. Fire skips could be emulated in harvest areas by leaving clumps of live trees

within cut-blocks. Small clumps may be sufficient for some native biota, but larger clumps probably act as refugia for more species (Saunders et al. 1991, MacKenzie and Steventon 1996, Seip and Parker 1997) because some animals require large undisturbed areas for protection and foraging (Pomeluzi et al. 1993, Schmiegelow et al. 1997).

There are limitations, however, to using a wildfire template for harvest because these disturbances remove different amounts of wood from the system. Only a small percentage of the wood is combusted during wildfire resulting in many snags, and when these snags fall, many pieces of downed woody material are present in post-fire areas (Lee et al. 1997). Following harvest, however, most of the wood is removed from the area and the trees that remain are live so that inputs of snags and down woody material is lower and occurs over a longer period than that found after fire. Given this discrepancy, managers should not expect to "mimic" the structure retained after fires and additional management strategies may be needed (Hutto 1995). It may be useful, on at least part of the harvest area, to retain trees in patterns that promote specific groups of native biota that would otherwise "fall through the cracks". One such pattern would be to have single live trees, or small clumps of live trees, scattered throughout the harvest area. If these trees do not get blown down, they will become large by mid-rotation with the result that many of the structures and micro-habitats associated with large live and dead trees will be present in mid-rotation forests rather than only in old forests (Spies et al. 1991). Large live and dead trees that are scattered, although uncommon following wildfires, may accelerate recovery of native biota because many birds use such structures (Hansen et al. 1991, Schieck et al. 1995). With this management strategy, single trees or many small clumps of trees, scattered throughout the harvest area, may have greater

positive impact on native biota than if all standing trees and snags had been retained as a few large clumps.

When managing for refugia, standing trees are retained in a different pattern than they are when managing for the production of dispersed large trees. Both patterns may be useful, but only by experimentally manipulating size of clumps and distance between clumps, and studying the birds living in those areas will we understand the impacts of each. In this study, we establish three treatments with different clump sizes and distances between clumps in two experimentally harvested areas to evaluate whether bird communities differ among the treatments. This project will evaluate differences that are present immediately following harvest. The experiment, however, will be maintained throughout a harvest rotation and the value of each the three treatments will be evaluated at various times in the future.

Objectives

I compared pre-harvest bird communities among stand groups and treatments to evaluate whether bird communities were similar in the treatments prior to harvest. Specifically, I evaluated: (1) whether the bird community that has specialized foraging and nesting requirements for large trees varied among treatments, and (2) whether the resident and migratory bird communities varied among treatments. This is the first year of a 3-year study. Post-harvest bird communities will be compared among the three treatments using data that will be collected during 1998 and 1999. In addition, post-harvest bird communities will be compared to the communities found in other studies within naturally disturbed forests and following different silviculture/harvest treatments.

Methods

A description of the study areas and experimental harvesting is presented in the Study Area and Methods (Chapter 2). Descriptions of the pre-harvest trees, snags, down woody materials, and understory vegetation is presented in Chapters 3 - 5.

Pre-harvest Bird Surveys

Depending on the shape of the stands, either two or three strip transects 100 m (109 yd.) wide and having a total length of 1 km (0.62 mi.), were established in each stand (Figs. 2.1 and 2.2). Strip transects were divided into ten 1-ha sites and permanent markers were located at the center of each site. To avoid censusing birds that lived in adjacent habitats, the outer edge of the strip transects were a minimum of 70 m (77 yd.), and when possible >100m (>109 yd.), from adjacent forest types. Birds were surveyed in each stand during four 12-day periods between 9 May and 26 June 1997. During each survey, the center lines of the transects were walked at a rate of 100 m per 10 min between first light and three hours past sunrise on days with little wind and no rain. The location of all birds heard or seen within the transects were recorded on maps. A 50 m (54.7 yd.) detection distance was used because this represents the maximum distance that all bird vocalizations could be detected reliably (Emlen and DeJong 1981, Wolf et al. 1995, Schieck 1997). At the center of each 1-ha (2.47 acre) site a 5-minute point count was conducted. Movements of birds were recorded on maps and summarized so that individual birds were tallied only once. When possible unknown birds were located and identified after the end of each survey. Bird vocalizations that could not be identified in the field were recorded using cassette tape recorders and parabolic microphones and identified later. The order that stands were surveyed during each period was randomly determined. Observers were trained, and methods were standardized prior to the start of the surveys.

Analyses of Pre-harvest Bird Communities

Bird species were categorized based on their migration, nesting, and foraging habits (Campbell et al. 1990, Semenchuk 1992, Gauthier and Auby 1996). Resident and migrant bird species may be affected differently by the dispersion of residual trees because residents live in boreal forests throughout the year. In addition, the dispersion of standing residual trees may have a greater affect on birds that nest or forage in large live or dead trees than those that do not. Tree Swallows (see Table 7.1 for scientific names) were included in the analyses

TABLE 7.1. Abundance (mean \pm SE), migration habits, and nesting/foraging habits of bird species that were detected in the pre-harvest forest in the Whitecourt study area and Grande Prairie study area during 1997.

Common Name	Genus And Species	Species Code	Migration Habits	Use Big Trees	Mean Abundance (number/10 ha)
Whitecourt Study Area					
Nothern Goshawk	<i>Accipiter gentilis</i>	NOGO	Resident	Yes	0.17 \pm 0.17
Ruffed Grouse	<i>Bonasa umbellus</i>	RUGR	Resident	No	0.08 \pm 0.08
Common Snipe	<i>Gallinago gallinago</i>	COSN	Migrant	No	0.17 \pm 0.11
Yellow-bellied Sapsucker	<i>Sphyrapicus varius</i>	YBSA	Migrant	Yes	0.67 \pm 0.31
Downy Woodpecker	<i>Picoides pubescens</i>	DOWO	Resident	Yes	0.08 \pm 0.08
Hairy Woodpecker	<i>Picoides villosus</i>	HAWO	Resident	Yes	0.33 \pm 0.14
Three-toed Woodpecker	<i>Picoides tridactylus</i>	TTWO	Resident	Yes	0.17 \pm 0.11
Northern Flicker	<i>Colaptes auratus</i>	NOFL	Migrant	Yes	0.50 \pm 0.26
Alder Flycatcher	<i>Empidonax alnorum</i>	ALFL	Migrant	No	0.08 \pm 0.08
Least Flycatcher	<i>Empidonax minimus</i>	LEFL	Migrant	Yes	0.08 \pm 0.08
Tree Swallow	<i>Tachycineta bicolor</i>	TRSW	Migrant	Yes	0.17 \pm 0.11
Gray Jay	<i>Perisoreus canadensis</i>	GRJA	Resident	Yes	1.58 \pm 0.5
Black-capped Chickadee	<i>Parus atricapillus</i>	BCCH	Resident	No	3.83 \pm 0.69
Boreal Chickadee	<i>Parus hudsonicus</i>	BOCH	Resident	No	2.92 \pm 0.57
Red-breasted Nuthatch	<i>Sitta canadensis</i>	RBNU	Resident	Yes	1.17 \pm 0.71
Brown Creeper	<i>Certhia americana</i>	BRCR	Migrant	Yes	0.25 \pm 0.18
Winter Wren	<i>Troglodytes troglodytes</i>	WIWR	Migrant	Yes	0.08 \pm 0.08
Golden-crowned Kinglet	<i>Regulus satrapa</i>	GCKI	Migrant	Yes	3.50 \pm 0.62
Ruby-crowned Kinglet	<i>Regulus calendula</i>	RCKI	Migrant	Yes	7.25 \pm 0.58
Swainson's Thrush	<i>Catharus ustulatus</i>	SWTH	Migrant	No	2.75 \pm 0.41
Hermit Thrush	<i>Catharus guttatus</i>	HETH	Migrant	No	1.50 \pm 0.36
American Robin	<i>Turdus migratorius</i>	AMRO	Migrant	No	0.17 \pm 0.11
Cedar Waxwing	<i>Bombycilla cedrorum</i>	CEWA	Migrant	No	0.17 \pm 0.11
Solitary Vireo	<i>Vireo solitarius</i>	SOVI	Migrant	Yes	1.58 \pm 0.43
Warbling Vireo	<i>Vireo gilvus</i>	WAVI	Migrant	Yes	0.58 \pm 0.19
Red-eyed Vireo	<i>Vireo olivaceus</i>	REVI	Migrant	Yes	0.33 \pm 0.22
Orange-crowned Warbler	<i>Vermivora celata</i>	OCWA	Migrant	No	0.25 \pm 0.13
Yellow Warbler	<i>Dendroica petechia</i>	YEWA	Migrant	No	1.08 \pm 0.34
Yellow-rumped Warbler	<i>Dendroica coronata</i>	YRWA	Migrant	Yes	10.58 \pm 0.54
American Redstart	<i>Setophaga ruticilla</i>	AMRE	Migrant	No	0.17 \pm 0.11
Ovenbird	<i>Seiurus aurocapillus</i>	OVEN	Migrant	No	0.25 \pm 0.13
Mourning Warbler	<i>Oporornis philadelphia</i>	MOWA	Migrant	No	0.33 \pm 0.14
Common Yellowthroat	<i>Geothlypis trichas</i>	COYE	Migrant	No	0.08 \pm 0.08
Canada Warbler	<i>Wilsonia canadensis</i>	CAWA	Migrant	No	0.25 \pm 0.25
Rose-breasted Grosbeak	<i>Pheucticus ludovicianus</i>	RBGR	Migrant	No	0.08 \pm 0.08
Chipping Sparrow	<i>Spizella passerina</i>	CHSP	Migrant	Yes	5.33 \pm 0.63
Lincoln's Sparrow	<i>Melospiza lincolni</i>	LISP	Migrant	No	0.08 \pm 0.08
White-throated Sparrow	<i>Zonotrichia albicollis</i>	WTSP	Migrant	No	2.25 \pm 0.62
Dark-eyed Junco	<i>Junco hyemalis</i>	DEJU	Migrant	No	4.50 \pm 0.66
Brown-headed Cowbird	<i>Molothrus ater</i>	BHCO	Migrant	No	0.25 \pm 0.13
White-winged Crossbill	<i>Loxia leucoptera</i>	WWCR	Resident	Yes	0.08 \pm 0.08
Pine Siskin	<i>Carduelis pinus</i>	PISI	Migrant	Yes	1.08 \pm 0.36

TABLE 7.1. cont.

Common Name	Genus And Species	Species Code	Migration Habits	Use Big Trees	Mean Abundance (number/10 ha)
Grande Prairie Study Area					
Ruffed Grouse	<i>Bonasa umbellus</i>	RUGR	Resident	No	0.33 ± 0.14
Common Snipe	<i>Gallinago gallinago</i>	COSN	Migrant	No	0.08 ± 0.08
Yellow-bellied Sapsucker	<i>Sphyrapicus varius</i>	YBSA	Migrant	Yes	3.25 ± 0.41
Downy Woodpecker	<i>Picoides pubescens</i>	DOWO	Resident	Yes	0.17 ± 0.17
Hairy Woodpecker	<i>Picoides villosus</i>	HAWO	Resident	Yes	0.33 ± 0.19
Northern Flicker	<i>Colaptes auratus</i>	NOFL	Migrant	Yes	0.67 ± 0.36
Pileated Woodpecker	<i>Dryocopus pileatus</i>	PIWO	Resident	Yes	0.08 ± 0.08
Alder Flycatcher	<i>Empidonax alnorum</i>	ALFL	Migrant	No	0.83 ± 0.21
Least Flycatcher	<i>Empidonax minimus</i>	LEFL	Migrant	Yes	8.33 ± 1.04
Tree Swallow	<i>Tachycineta bicolor</i>	TRSW	Migrant	Yes	0.17 ± 0.17
Gray Jay	<i>Perisoreus canadensis</i>	GRJA	Resident	Yes	0.25 ± 0.25
Common Raven	<i>Corvus corax</i>	CORA	Resident	Yes	0.25 ± 0.18
Black-capped Chickadee	<i>Parus atricapillus</i>	BCCH	Resident	No	2.58 ± 0.58
Boreal Chickadee	<i>Parus hudsonicus</i>	BOCH	Resident	No	0.25 ± 0.13
Red-breasted Nuthatch	<i>Sitta canadensis</i>	RBNU	Resident	Yes	0.58 ± 0.29
Golden-crowned Kinglet	<i>Regulus satrapa</i>	GCKI	Migrant	Yes	0.08 ± 0.08
Ruby-crowned Kinglet	<i>Regulus calendula</i>	RCKI	Migrant	Yes	0.42 ± 0.26
Veery	<i>Catharus fuscescens</i>	VEER	Migrant	No	0.08 ± 0.08
Swainson's Thrush	<i>Catharus ustulatus</i>	SWTH	Migrant	No	3.08 ± 0.43
Hermit Thrush	<i>Catharus guttatus</i>	HETH	Migrant	No	1.08 ± 0.26
American Robin	<i>Turdus migratorius</i>	AMRO	Migrant	No	1.67 ± 0.53
Cedar Waxwing	<i>Bombycilla cedrorum</i>	CEWA	Migrant	No	0.25 ± 0.18
Solitary Vireo	<i>Vireo solitarius</i>	SOVI	Migrant	Yes	0.25 ± 0.18
Warbling Vireo	<i>Vireo gilvus</i>	WAVI	Migrant	Yes	1.25 ± 0.37
Red-eyed Vireo	<i>Vireo olivaceus</i>	REVI	Migrant	Yes	3.92 ± 0.69
Tennessee Warbler	<i>Vermivora peregrina</i>	TEWA	Migrant	No	3.17 ± 0.82
Orange-crowned Warbler	<i>Vermivora celata</i>	OCWA	Migrant	No	0.58 ± 0.23
Connecticut Warbler	<i>Oporornis agilis</i>	COWA	Migrant	No	1.08 ± 0.29
Yellow Warbler	<i>Dendroica petechia</i>	YEWA	Migrant	No	3.83 ± 0.74
Magnolia Warbler	<i>Dendroica magnolia</i>	MAWA	Migrant	No	0.58 ± 0.23
Yellow-rumped Warbler	<i>Dendroica coronata</i>	YRWA	Migrant	Yes	9.50 ± 0.87
Black-throated Green Warbler	<i>Dendroica virens</i>	BTGW	Migrant	Yes	0.33 ± 0.22
Blackpoll Warbler	<i>Dendroica striata</i>	BPWA	Migrant	No	0.08 ± 0.08
Black and white Warbler	<i>Mniotilta varia</i>	BWWA	Migrant	No	0.08 ± 0.08
American Redstart	<i>Setophaga ruticilla</i>	AMRE	Migrant	No	4.67 ± 1.11
Ovenbird	<i>Seiurus aurocapillus</i>	OVEN	Migrant	No	5.92 ± 0.85
Northern Waterthrush	<i>Seiurus noveboracensis</i>	NOWA	Migrant	No	0.33 ± 0.22
Mourning Warbler	<i>Oporornis philadelphia</i>	MOWA	Migrant	No	0.67 ± 0.28
Common Yellowthroat	<i>Geothlypis trichas</i>	COYE	Migrant	No	0.17 ± 0.11
Canada Warbler	<i>Wilsonia canadensis</i>	CAWA	Migrant	No	0.08 ± 0.08
Western Tanager	<i>Piranga ludoviciana</i>	WETA	Migrant	Yes	2.17 ± 0.55
Rose-breasted Grosbeak	<i>Phœcticus ludovicianus</i>	RBGR	Migrant	No	3.83 ± 1.04
Chipping Sparrow	<i>Spizella passerina</i>	CHSP	Migrant	Yes	5.50 ± 0.96
Song Sparrow	<i>Melospiza melodia</i>	SOSP	Migrant	No	0.17 ± 0.11
Lincoln's Sparrow	<i>Melospiza lincolnii</i>	LISP	Migrant	No	1.83 ± 0.53
White-throated Sparrow	<i>Zonotrichia albicollis</i>	WTSP	Migrant	No	15.92 ± 1.40
Dark-eyed Junco	<i>Junco hyemalis</i>	DEJU	Migrant	No	4.58 ± 0.81
Red-winged Blackbird	<i>Agelaius phoeniceus</i>	RWBL	Migrant	No	0.33 ± 0.19
Brown-headed Cowbird	<i>Molothrus ater</i>	BHCO	Migrant	No	3.08 ± 0.93
Pine Siskin	<i>Carduelis pinus</i>	PISI	Migrant	Yes	1.08 ± 0.67

when detected flying within the transect. This species, unlike the other forest birds, does not land in trees or on the ground while foraging (Godfrey 1986, Semenchuk 1992). Other species that flew over during the surveys were not included in the analyses.

For each bird species, an index of density in each 1 ha (2.47 acres) site was calculated as the maximum number of individuals detected during the four surveys. Differences in bird species richness and total abundance were evaluated among stand groups using ANOVA for all bird species together and subsequently and for each of the four categories of bird species (residents versus migrants, and species that use big trees versus those that do not). Since bird species richness and total abundance differed among stand groups (see Results), differences in both were evaluated among treatments after first adjusting for differences among stand groups (Type III sum of squares in a 2-way ANOVA; SAS Institute Inc 1989). In all analyses, results were considered statistically significant if the probability of them occurring by chance was less than 0.05.

Differences in bird species richness and abundance do not describe all the differences among bird communities because species composition may differ even when richness and abundance are similar. I used correspondence analyses and canonical correspondence analyses (CA and CCA respectively, Halvorsen 1996) to evaluate differences in bird communities among stand groups and treatments. In all CAs and CCAs, the option to down-weight rare species was used (ter Braak 1992) because chance events may have resulted in these rare species being observed in non-typical habitats. Differences among stand groups and treatments were tested by coding dummy variables for the appropriate categories, including these dummy variables as environmental variables in the analyses, and then testing for explained variation using a bootstrap Monte Carlo test with 99 iterations (ter Braak 1992). To evaluate whether variation in bird communities was related to variation in vegetation, I included 16 characteristics of live and dead vegetation in the CCA. Live trees were categorized into three groups: the dominate species (pine in

Whitecourt study area, and aspen in Grande Prairie study area), the second most dominant species (black spruce in Whitecourt study area, and white spruce in Grande Prairie study area), and all other tree species. Within each of these three categories of trees, the densities of trees with DBH < and > 20 cm (7.9 in.) was calculated. In addition, in both study areas I calculated densities of snags with DBH < and > 20 cm (7.9 in.), volume of downed woody material per ha, and percentage of the ground surface covered by shrubs <0.5 m (<1.6 ft.) high, shrubs >0.5 m (>1.6 ft.) high, herbs, grass, moss, and dead vegetation.

Results

Whitecourt Study Area

Forty-two species and 901 individuals were detected during the bird surveys (Table 7.1). This bird community was dominated by species that migrate with only 24% of the species being year-round residents. Most residents (70% of 10 species), but fewer migrants (40% of 32 species) nested or foraged in large live or dead trees. More bird species (Table 7.2) and individuals (Table 7.3) were observed in stand group B than in the other three stand groups. That difference was largely due to more resident bird species that use large trees being present in stand group B. Species richness and abundance were similar among treatments for all categories of birds after adjusting for differences among stand groups (Tables 7.2 and 7.3).

When evaluating differences in bird communities among sites, the first and second functions of the CA had eigen values of 0.07 and 0.04, respectively. Combined, these two functions accounted for only 3.4% of the variation in bird species density. However, even with this small amount of explained variation, bird communities were statistically different among stand groups (Fig. 7.1; $F = 1.8$, $P < 0.001$). After adjusting for differences among stand groups, bird communities did not differ among treatments (during the bootstrap Monte Carlo test permutations were restricted to within the appropriate stand group; $F = 1.2$, $P = 0.10$). In addition, all categories of bird species were present in all stand groups (Fig. 7.2). Variation in the bird communities was not related to

TABLE 7.2. Mean (\pm SE) species richness within sites within each stand group and treatment. Species were categorized by whether or not they used large trees or snags and their migration habitats. There were 3 and 121 degrees of freedom for the tests involving stand groups and 2 and 119 degrees of freedom for the tests involving treatments. Groups with different superscripts are statistically different (Student-Neuman-Keuls range test).

Categories	Test Statistic	Use Big Trees		Do Not Use Big Trees		Total
		Resident	Migrant	Resident	Migrant	
Whitecourt Study Area						
Stand Group A		0.2 \pm 0.1 ^b	2.9 \pm 0.2	0.6 \pm 0.1	1.1 \pm 0.1	4.8 \pm 0.3 ^{ab}
Stand Group B		0.5 \pm 0.1 ^a	3.0 \pm 0.2	0.6 \pm 0.1	1.5 \pm 0.2	5.6 \pm 0.4 ^a
Stand Group C		0.2 \pm 0.1 ^b	2.8 \pm 0.2	0.7 \pm 0.1	1.5 \pm 0.2	5.1 \pm 0.3 ^{ab}
Stand Group D		0.2 \pm 0.1 ^b	2.3 \pm 0.3	0.6 \pm 0.1	1.4 \pm 0.2	4.5 \pm 0.4 ^b
	F	3.5	1.6	1.6	0.9	2.49
	P	0.02	0.19	0.22	0.49	0.05
Treatment 1		0.2 \pm 0.1	2.5 \pm 0.2	0.8 \pm 0.1	1.2 \pm 0.2	4.7 \pm 0.3
Treatment 2		0.3 \pm 0.1	2.9 \pm 0.2	0.5 \pm 0.1	1.5 \pm 0.2	5.3 \pm 0.3
Treatment 3		0.3 \pm 0.1	2.8 \pm 0.2	0.6 \pm 0.1	1.4 \pm 0.2	5.0 \pm 0.3
	F	0.6	0.8	0.3	1.2	1.3
	P	0.94	0.43	0.77	0.32	0.27
Grande Prairie Study Area						
Stand Group A		0.1 \pm 0.1	3.2 \pm 0.2	0.3 \pm 0.1 ^{ab}	4.1 \pm 0.2 ^b	7.7 \pm 0.4 ^b
Stand Group B		0.1 \pm 0.1	2.7 \pm 0.2	0.4 \pm 0.1 ^a	4.0 \pm 0.2 ^b	7.3 \pm 0.4 ^b
Stand Group C		0.2 \pm 0.1	3.4 \pm 0.2	0.1 \pm 0.1 ^b	5.3 \pm 0.3 ^a	9.0 \pm 0.5 ^a
Stand Group D		0.1 \pm 0.1	2.8 \pm 0.2	0.3 \pm 0.1 ^{ab}	5.0 \pm 0.3 ^a	8.2 \pm 0.4 ^a
	F	0.2	2.5	3.2	6.4	4.6
	P	0.92	0.06	0.03	0.001	0.001
Treatment 1		0.1 \pm 0.1	3.0 \pm 0.2	0.3 \pm 0.1	4.5 \pm 0.2	7.9 \pm 0.04
Treatment 2		0.1 \pm 0.1	3.2 \pm 0.2	0.2 \pm 0.1	5.0 \pm 0.3	8.5 \pm 0.06
Treatment 3		0.1 \pm 0.1	2.9 \pm 0.2	0.3 \pm 0.1	4.3 \pm 0.3	7.6 \pm 0.04
	F	0.6	0.8	0.3	2.3	2.2
	P	0.94	0.44	0.77	0.10	0.14

TABLE 7.3. Mean (\pm SE) abundance per ha within each stand group and treatment. Species were categorized by whether or not they used large trees or snags and their migration habitats. There were 3 and 121 degrees of freedom for the tests involving stand groups and 2 and 119 degrees of freedom for the tests involving treatments. Groups with different superscripts are statistically different (Student-Neuman-Keuls range test).

Categories	Test Statistic	Use Big Trees		Do Not Use Big Trees		Total
		Resident	Migrant	Resident	Migrant	
Whitecourt Study Area						
Stand Group A		0.4 \pm 0.2	3.3 \pm 0.2	0.6 \pm 0.1	1.2 \pm 0.1	5.5 \pm 0.4
Stand Group B		0.6 \pm 0.1	3.5 \pm 0.3	0.7 \pm 0.1	1.6 \pm 0.2	6.3 \pm 0.5
Stand Group C		0.3 \pm 0.2	3.2 \pm 0.3	0.7 \pm 0.1	1.6 \pm 0.2	5.9 \pm 0.4
Stand Group D		0.2 \pm 0.1	2.7 \pm 0.3	0.7 \pm 0.1	1.5 \pm 0.2	5.1 \pm 0.4
	F	1.2	1.8	0.1	0.8	2.3
	P	0.31	0.14	0.95	0.46	0.08
Treatment 1		0.3 \pm 0.1	2.9 \pm 0.2	0.8 \pm 0.1	1.2 \pm 0.1	5.2 \pm 0.4
Treatment 2		0.5 \pm 0.1	3.4 \pm 0.3	0.5 \pm 0.1	1.7 \pm 0.2	6.2 \pm 0.5
Treatment 3		0.3 \pm 0.1	3.2 \pm 0.2	0.7 \pm 0.1	1.4 \pm 0.2	5.6 \pm 0.4
	F	0.4	1.3	0.5	1.7	2.0
	P	0.65	0.29	0.61	0.19	0.11
Grande Prairie Study Area						
Stand Group A		0.1 \pm 0.1	3.9 \pm 0.3 ^{ab}	0.3 \pm 0.1 ^{ab}	5.0 \pm 0.3 ^c	9.3 \pm 0.5 ^b
Stand Group B		0.1 \pm 0.1	3.4 \pm 0.3 ^{ab}	0.6 \pm 0.2 ^a	4.7 \pm 0.3 ^c	8.8 \pm 0.4 ^b
Stand Group C		0.2 \pm 0.1	4.4 \pm 0.4 ^a	0.1 \pm 0.1 ^b	7.4 \pm 0.5 ^a	12.0 \pm 0.6 ^a
Stand Group D		0.2 \pm 0.1	3.1 \pm 0.2 ^b	0.3 \pm 0.1 ^{ab}	6.2 \pm 0.4 ^b	9.8 \pm 0.5 ^b
	F	0.3	3.3	3.2	11.8	9.7
	P	0.86	0.02	0.03	<0.001	<0.001
Treatment 1		0.1 \pm 0.1	3.5 \pm 0.2	0.3 \pm 0.1	5.6 \pm 0.2	9.6 \pm 0.5
Treatment 2		0.1 \pm 0.1	4.3 \pm 0.4	0.3 \pm 0.1	6.2 \pm 0.3	10.7 \pm 0.6
Treatment 3		0.2 \pm 0.1	3.3 \pm 0.3	0.3 \pm 0.1	5.5 \pm 0.4	9.6 \pm 0.5
	F	0.4	25	0.1	1.5	2.0
	P	0.66	0.09	0.87	0.22	0.11

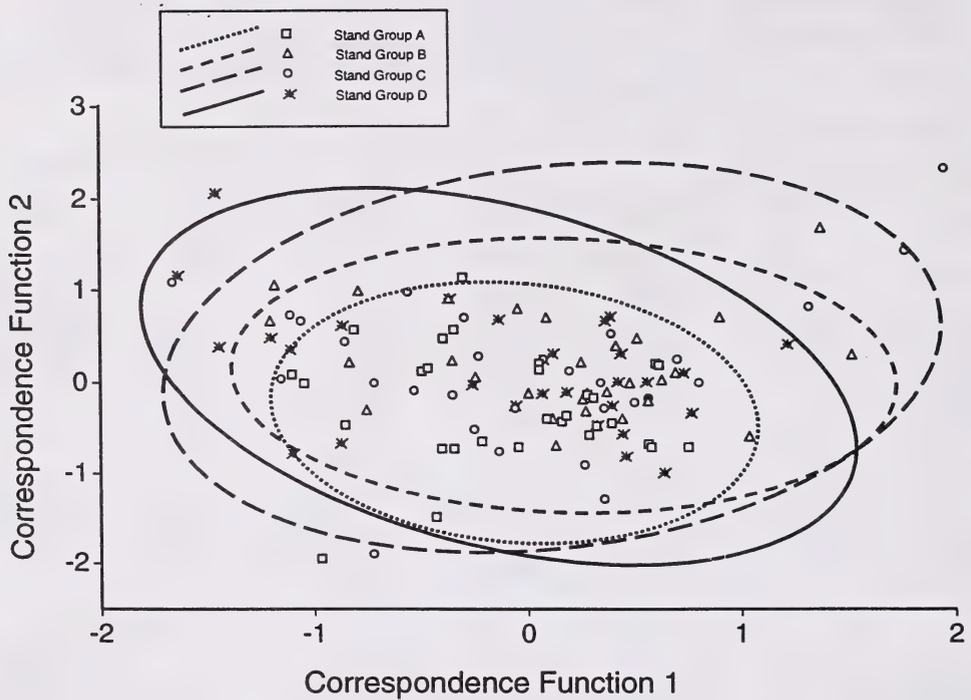


FIGURE 7.1. Correspondence analyses based on bird densities in the Whitecourt study area during 1997. Sites that are close together within the correspondence plot have similar bird communities and sites that are far apart have dissimilar bird communities. Sites were categorized based on stand group. For each stand group bivariate ellipses that enclose 90% of the sites were presented (Systat Inc. 1989).

variation in vegetation characteristics among sites ($F = 1.1$, $P = 0.32$).

Grande Prairie Study Area

Fifty species and 1592 individuals were detected during the bird surveys (Table 7.1). This bird community was dominated by species that migrate with only 18% of the species being year-round residents. Most residents (67% of 9 species), but fewer migrants (34% of 41 species) nested or foraged in large live or dead trees. Both species richness (Table 7.2) and number of individuals (Table 7.3) differed among stand groups. Differences in species richness were mainly due to differences for birds that did not use large trees and snags, whereas differences in abundance were due to differences for birds that did not use large trees and snags and differences for birds that migrate and use large trees. After adjusting for differences in richness and abundance among stand groups, richness and abundance were similar among treatments for all categories of birds.

When evaluating differences in bird communities among sites, the first and second functions of the CA had eigen values of 0.07 and 0.04, respectively. Combined, these two functions accounted for only 4.1% of the variation in bird species density. However, even with this small amount of explained variation, bird communities were statistically different among stand groups (Fig. 7.3; $F = 1.9$, $P < 0.01$). After adjusting for differences among stand groups bird communities did not differ among treatments ($F = 1.2$, $P = 0.14$). In addition, all categories of bird species were present in all stand groups (Fig. 7.4). Variation in the bird communities was not related to variation in vegetation characteristics among sites ($F = 1.2$, $P = 0.18$).

Discussion

To achieve a rigorous experimental design, we chose two study areas that were large and relatively homogenous. Within these study areas, the stands (i.e., experimental harvest units) were similar to each other prior to harvest, but as expected within natural forests, the stands were not identical to each other. We

created four groups of stands in each study area based on vegetation in and around the stands (see Chapter 2). This resulted in a study design where some of the variation in vegetation and structure could be accounted for statistically in a stratified analysis (Chapters 3 and 4). Similarly, within both study areas, variation among bird communities was related to differences among stand groups. That was true for both the simple analyses of bird species richness and abundance, and for the more complex community analyses. However, variation among stand groups accounted for only a small amount of the variation in bird communities and the bird communities overlapped greatly among the stand groups. This was expected since there were only slight differences in vegetation among stand groups.

After controlling for the differences among stand groups, bird communities did not differ among treatments prior to harvest. Consequently, if bird communities differ among treatments after harvest, those differences can be attributed unambiguously to differences in the size and dispersion of tree patches that were retained in these experimental areas during harvest. Thus, as planned, this study will provide a rigorous comparison of bird communities in among the three dispersions of standing trees and snags. The bird communities found in the stands prior to harvest are benchmarks from which I will evaluate differences after harvest.

Future Research

Bird communities will be surveyed during 1998 and 1999. However, most of the post-harvest bird surveys will be conducted during 1999 because birds may crowd into the remaining unharvested habitat immediately after harvest and those high densities probably will decrease during the next year (Lovejoy et al. 1986, Schmiegelow 1997). Bird surveys during 1999 will be an extended version of those conducted prior to harvest. Within each stand, the 10 ha (24.7 acres) study area (Figs. 2.1 and 2.2) will be surveyed using a combination of strip transects and point counts (see Methods). In addition, eight point counts will be centered in residual

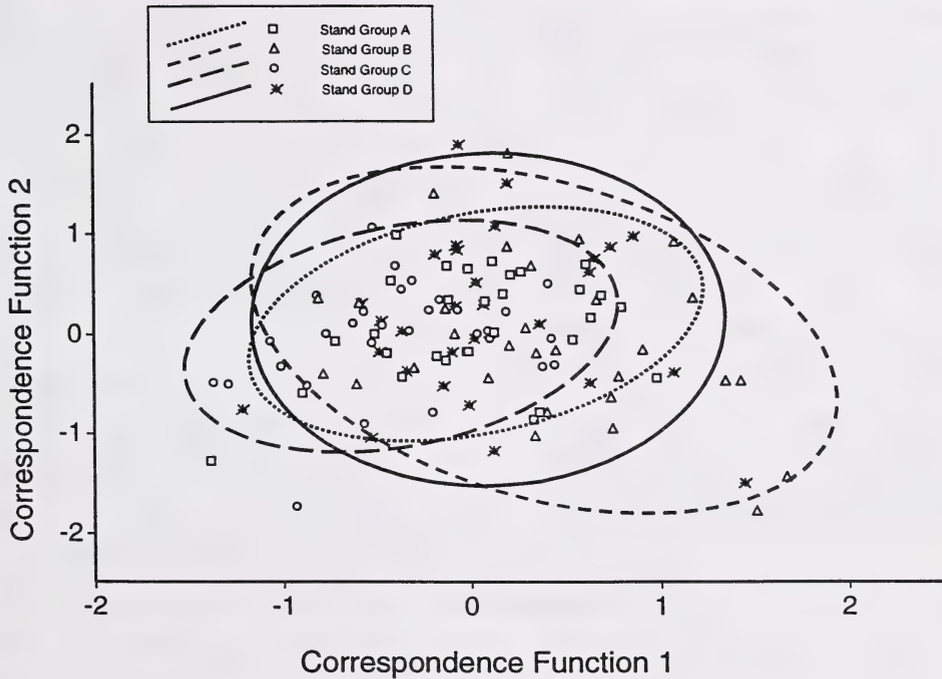


FIGURE 7.3. Correspondence analyses based on bird densities in the Grand Prairie study area during 1997. Sites that are close together within the correspondence plot have similar bird communities and sites that are far apart have dis-similar bird communities. Sites were categorized based on stand group. For each stand group bivariate ellipses that enclosed 90% of the sites were presented.

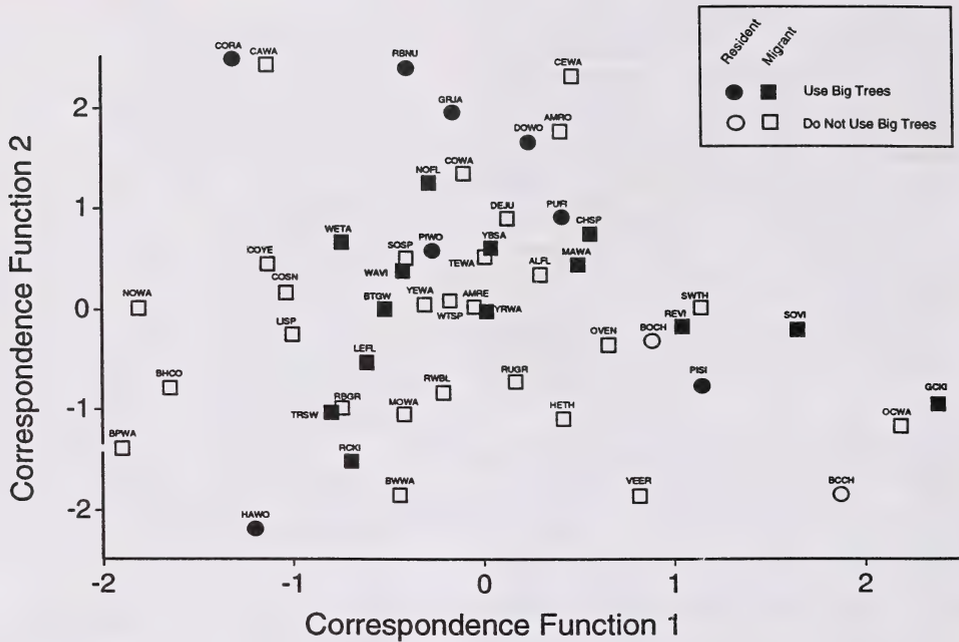


FIGURE 7.4. Predicted location for each bird species based on the correspondence analyses of bird densities in the Grande Prairie study area during 1997. Species that are close together have strong positive covariance in abundance among sites whereas species that are far apart have strong negative covariance in abundance among sites. Species were categorized based on their migration and nesting/foraging habits.

clumps within each stand and four counts will be located >100 m (>109 yd.) from residual clumps in each stand. Surveys will be conducted in all treatments once between 15 and 30 April, and three times at bi-weekly intervals between 15 May and 30 June. During the reduced 1998 survey, only 40 point counts will be conducted in each study area. Ten of these point counts will be centered within small residual patches, ten centered within mid-sized residual patches, ten centered within large residual patches, and ten >100 m (>109 yd.) from residual patches. Results from these bird surveys will be analyzed and presented during fall and winters of 1998 and 1999.

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CHAPTER 8. THE USE OF RESIDUAL PATCHES BY BATS IN THE BOREAL FOREST OF NORTHWESTERN ALBERTA

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Introduction

Timber harvesting strategies are restructuring the boreal forest mosaic in Alberta. With the recent introduction of aspen into harvesting plans, forests are being reshaped at an even faster rate. Holling (1995) characterized traditional resource management as controlling the variability of ecosystems so that a steady flow of goods and services could be produced for humans. This traditional view of forest value is limited to the monetary worth of present and future timber resources. Emerging environmental, ecological, and social concerns are forcing a much broader perspective of resource-use strategies. Strategies of ecosystem management are being developed (e.g., Alberta Forest Conservation Strategy 1997), encouraging private companies to reassess the value of the forest and modify traditional harvesting practices (e.g., Alberta Pacific Forest Industries Inc. 1996).

Harvesting practices are evolving to include leaving standing trees and snags in harvested areas in an effort to retain pre-harvest species of flora and fauna. Such forest structure may make harvested areas more desirable to native bats that use the forest environment to forage, roost, and breed (Thomas 1988). Currently, the pattern in which residual trees and snags are left on the landscape is decided subjectively (Alberta Pacific Forest Industries Inc. 1996). Objective decision-making requires a greater understanding of the effect on biotic communities of retaining residual elements and the manner of their dispersion.

The boreal forest of Alberta supports five species of bats (van Zyll de Jong 1983). These bats show a fair degree of fidelity to roost sites in the forest when they return from hibernaculum or migration. During the spring and summer, cavity-roosting colonial species use a wide variety of cavities including spaces

behind exfoliated bark and natural or excavated cavities in large, old snags (Thomas and La Val 1988). Within Douglas-fir forests of southern Washington State, old forests were considered critical for bats because they contained large, decayed snags, which provide crevices for cavity-roosting bat species (Thomas 1988). The two foliage-roosting solitary species (*Lasiurus cinereus* and *Lasionycteris noctivagans*) generally do not exploit cavities but roost in fixed or shifting locations in foliage of large, live trees (Barclay 1984). In some species, males and females use different habitats (Barclay 1991). Males do not maintain high body temperatures during the day and may select cool roosts to minimize their metabolic rates and diurnal energy expenditures (Thomas 1988). Females have a reproductive constraint on their ability to exploit the energy saving of torpor because fetal growth rates and presumably milk production are temperature-dependent processes (Racey 1982). During pregnancy and lactation, females maintain body temperatures well above ambient during the day to maximize fetal growth rates (Kunz 1982b, Racey 1982) and as such tend to select warm roost sites. Thus, retaining bats in forest patches will depend upon requirements of different bat species and sexes.

In this study, three treatments with different clump size and distances between clumps in two experimentally harvested areas will be established to evaluate how bat communities respond to harvested forests. We will concentrate on differences that are detected in the first two years following harvest, however, throughout the forest rotation the opportunity exists to evaluate the value to bats of the patches that are created in the experiment.

Objectives

I will compare bat communities among three different harvesting treatments (Chapter 2) to evaluate which pattern of dispersion of residual

trees maximizes bat diversity and abundance. Pre-harvest sampling will be used to assess whether bat communities were similar among treatments prior to harvest. Post-harvesting bat communities will be compared among the three treatments using data that will be collected during 1998 and 1999.

Methods

A description of the study areas and experimental harvesting is presented in the Study Area and Methods section (Chapter 2). Descriptions of the pre-harvest trees, snags, downed woody materials, and understory vegetation are presented in Chapters 3 - 5.

Bat detectors were used to estimate species richness and abundance in each stand in both Whitecourt and Grande Prairie study areas in July and August 1997. Ultrasonic detectors do not provide data that can be translated directly into estimates of population density (Thomas and La Val 1988); they do however, provide a relatively unbiased index of the levels of use or animal abundance among habitats (Thomas 1988). Detectors consisted of an Anabat IV ultrasonic bat detector (Titley Electronics, AUS), a delay switch, a cassette recorder, and a 12-volt battery, all of which were housed in a 1.5 litre (1.6 gal.) rubbermaid container. A 2 cm (0.8 in.) hole was cut into the side of the container to accommodate the microphone. The battery was used to run the detector and the delay switch. The delay switch was set to turn the detector on at dusk and off at dawn. The sensitivity setting on the detectors was set between 6 and 7 to minimize insect noise and to eliminate detection of bats flying in adjacent stand groups. To allow the bat call to be recorded by a cassette recorder, the frequency of each call was reduced from the ultrasonic to the audible range by setting the division ratio on detectors to 16.

Bats were surveyed from three points, approximately 300 m (328 yd.) apart, in each stand. Two stands in each study area were sampled simultaneously. Detectors were attached to trees 1.5 m (5 ft.) off the ground and oriented such that the microphone was pointing 45-degree above horizontal. Detectors were run

for three consecutive nights in each stand and were checked each morning to ensure proper functioning. After three days, detectors were moved to new stands.

Bats were identified by analyzing their search-phase echolocation calls. Each sequence of one or more echolocation pulses with <1 s between sequential pulses was recorded as a pass by a bat (Fenton 1970). Calls were analyzed using Anabat and Analook software (Titley Electronics, AUS). Frequency range and call shape from the calls we recorded were compared with a library of calls from known species.

Differences in bat abundance were evaluated among stand groups and treatments using one-way ANOVAs. In all analyses, results were considered statistically significant if the probability of them occurring by chance was less than 0.05. Canonical correlation analysis was used to evaluate the relationship between the bat abundance and characteristics of trees in each stand. In both areas, I included the following tree variables in the analyses: density of snags with a diameter at breast height (DBH) of <20 cm (<7.9 in.), snags >20 cm (>7.9 in.) DBH, and total volume of down woody material. In the Whitecourt study area, density of pine and black spruce <20 cm (<7.9 in.) DBH and density of pine and black spruce >20 cm (>7.9 in.) DBH were included in the analyses. In the Grande Prairie study area, density of aspen and white spruce <20 cm (<7.9 in.) DBH and density of aspen and white spruce >20 cm (>7.9 in.) DBH were included in the analyses.

Results

Based on 756 hrs of data collection in each area almost twice as many bat calls were detected in the Grande Prairie study area (n = 120) as in the Whitecourt study area (n = 67). Of the bats detected (n = 187), the majority were identified as little brown bats (*Myotis lucifugus*). Hoary bats (*Lasiurus cinereus*) were detected three times in the Whitecourt study area. Accordingly, I restricted the analyses to detections of the little brown bats.

TABLE 8.1. Mean (\pm SE) number of little brown bat calls detected within each stand group and treatment in Whitecourt and Grande Prairie study areas.

		Whitecourt	Grande Prairie
Stand group	1	2.3 (1.5)	14.0 (4.4)
	2	2.7 (1.5)	7.3 (2.0)
	3	2.7 (1.2)	1.7 (0.7)
	4	8.0 (6.5)	2.3 (1.2)
Treatment	1	1.8 (1.1)	6.0 (2.3)
	2	2.3 (1.0)	7.5 (5.1)
	3	7.8 (4.5)	5.5 (2.2)

Bat detections in the Whitecourt study area did not differ among treatments ($F_{2,11} = 1.50$, $P = 0.27$) or among stand groups ($F_{3,11} = 0.62$, $P = 0.62$; Table 8.1). In the Grande Prairie study area no significant difference was detected among treatments ($F_{2,11} = 0.09$, $P = 0.92$). However, among stand groups, some differences were significant ($F_{3,11} = 5.23$, $P = 0.03$). A Tukey post-hoc test suggested that difference between stand group 1 and stand groups 3 and 4 are responsible for the significant difference.

The density of trees and snags accounted for only a small amount of the variation in the bat numbers in both study areas (4% in Grande Prairie, and 5% in Whitecourt). In the Grande Prairie study area, aspen density [both < and > 20 cm (7.9 in.) DBH] was the best predictor of bat abundance (Fig. 8.1). Snag density [>20 cm (>7.9 in.) DBH] was the best predictor of bat abundance in the Whitecourt study area and was positively correlated with bat abundance (Fig. 8.2). The graphical depiction of the relationship between bat abundance and tree variables (Fig. 8.1 and 8.2) also shows the lack of differentiation among the stand groups and treatments in both study areas.

Discussion

Stands in each area were categorized into four stand groups based on observed similarity in vegetation in and around the stands (see Chapter 2). Little brown bats were equally abundant in all stand groups in the Whitecourt area, but were more abundant in stand group 1 in the Grande Prairie area. Accordingly, if the density of little brown bats in the Whitecourt area differs among stands after harvest, those differences can be attributed to differences in the size and dispersion of tree patches that were retained at harvest. However, in the Grande Prairie study area, post-harvest differences will have to be evaluated after taking into account pre-harvest differences.

Aspen trees in the Grande Prairie study area and snags in the Whitecourt study area appeared to be important in predicting bat abundance. Bats rely on crevices in trees or raised bark to roost during the day. Accordingly, snags in the

Whitecourt study area may be important because of the roost sites they offer. Bats can also roost in live trees, using naturally occurring cracks or excavated cavities. Bats may also select an area based on the dispersion of live trees. Although bats do not appear to spend a significant portion of their time feeding in the forest (Thomas 1988), when they do, they require a corridor to fly and feed. Mature aspen in the Grande Prairie study area may provide roost sites and/or the proper corridors for feeding.

Future Research

Bat communities will be surveyed during 1998 and 1999 to evaluate the effect of the harvest treatments on bats. Three bat detectors units will be positioned in a stand each night; one at the center of a residual patch, one at the patch edge, and one within the clear-cut area. Bat calls will be recorded for three consecutive nights. Flying insect communities will be sampled concurrently at these same locations using insect intercept traps that will be suspended at a height of 2 m (6.6 ft.). Insect collection will be limited to the time when bats are most active, from sunset to three hours after sunset, by using automated traps that will be controlled by photocells, timers, and motors. Results from bat surveys will be analyzed and presented during fall and winters of 1998 and 1999.

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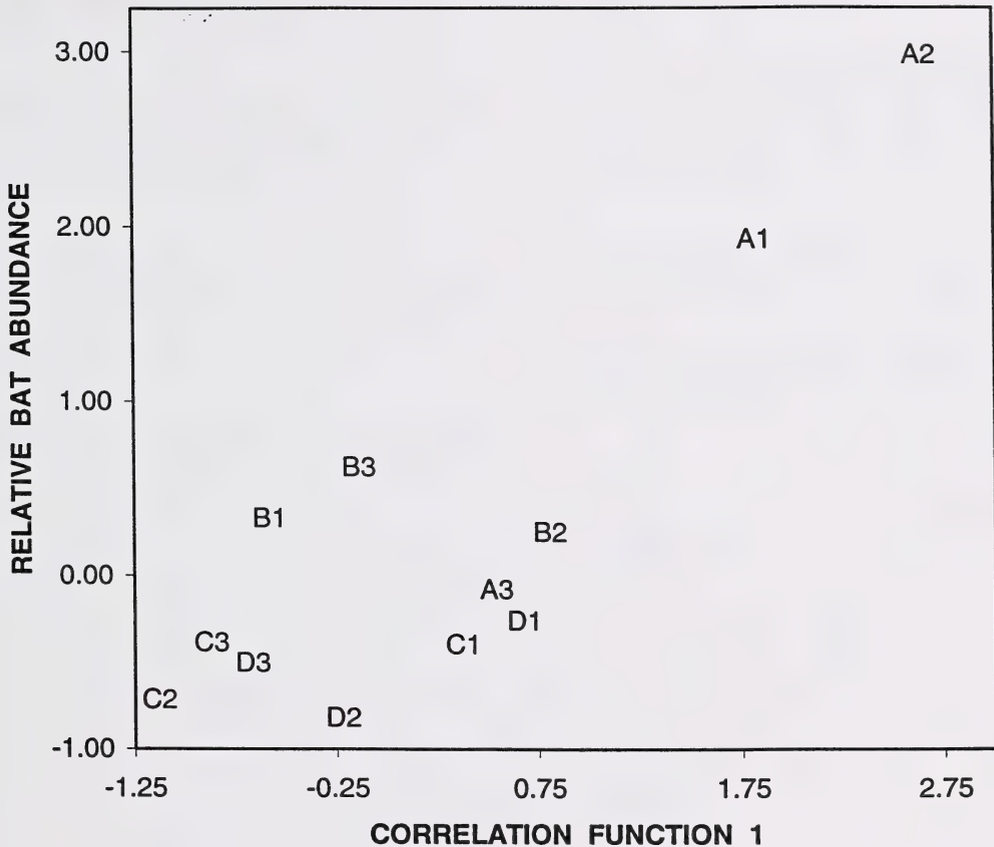


FIGURE 8.1. Canonical correlation analyses based on abundance of little brown bats in boreal forest stands near Grandè Prairie, AB during 1997. The first correlation function describes an increasing density of aspen trees. The symbols represent each stand identified by stand group (A-D) and treatment (1-3).

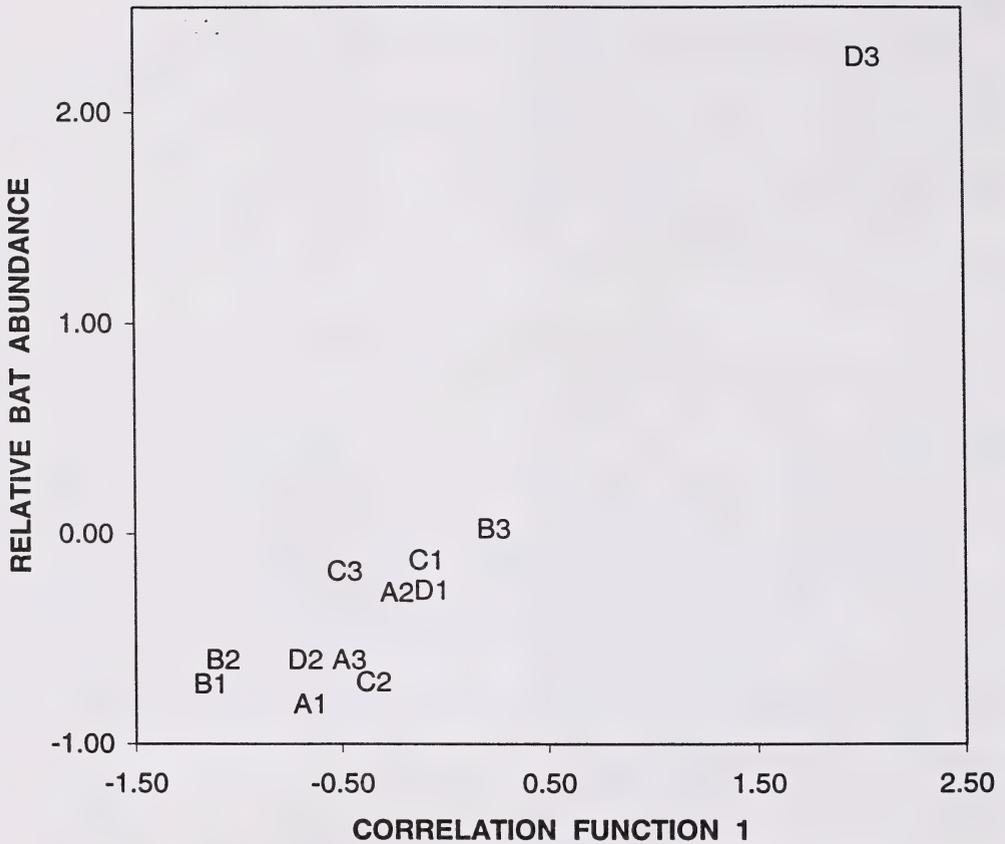


FIGURE 8.2. Canonical correlation analyses based on abundance of little brown bats in boreal forest stands near Whitecourt, AB during 1997. The first correlation function describes an increasing density of snags with a DBH of >20 cm (>7.9 in.). The symbols represent each stand identified by stand group (A-D) and treatment (1-3).

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