Dietary Variation in the Northern Goshawk in British Columbia

# Abstract

# Introduction

Effective wildlife conservation often requires understanding diet composition and its consequences for population demographics (Ferrer and Negro 2004, Stier et al. 2016). Specialist predators selectively consume a narrow range of prey species regardless of their abundance, whereas generalists opportunistically consume a wider range and readily switch between species in response to changes in prey abundance (Steenhof and Kochert 1988, Terraube and Arroyo 2011). Specialization increases foraging efficiency on the preferred prey at the potential cost of decreased reproductive success for the specialist when that prey is scarce (Newton 1998). However, even in a generalist predator a single key prey species can be a major driver of reproductive success (Elmhagen et al. 2000, Resano‐Mayor et al. 2016). For at-risk predators, increasing the abundance of key prey species may be a useful conservation tool (Ferrer and Negro 2004, Forsman et al. 2004, Resano‐Mayor et al. 2016).

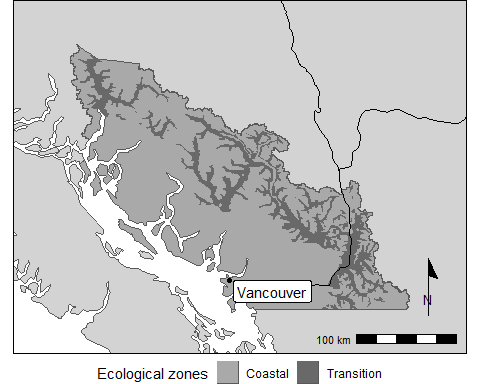
The northern goshawk (*Accipiter gentilis*) is a large forest-dwelling raptor with a Holarctic distribution. A generalist predator, the goshawk hunts a variety of small- and medium-sized mammals and birds, including squirrels, rabbits and hares, grouse, jays and crows, and pigeons (Squires et al. 2020). Despite this diverse diet, a single prey species or narrow suite of species has a strong effect on the demographics of many goshawk populations. In the Yukon goshawks depend on snowshoe hare (*Lepus americanus*) and show strong variation in productivity, mortality, and space use in response to cyclical changes in hare abundance (Doyle and Smith 1994). Goshawks in Scandinavia likewise rely on a single prey taxon and show changes in productivity and occupancy based on the annual abundance of four grouse species (subfamily Tetraoninae) (Tornberg et al. 2005). In contrast, goshawks in the American Southwest have a wide prey base and regularly take some fourteen different species (Boal and Mannan 1994). Fluctuations in goshawk productivity in this region are small and driven by total prey abundance, though the most influential single species is red squirrel (*Tamiasciurus hudsonicus*) (Salafsky et al. 2007). These examples suggest the identity and influence of key prey species in such an adaptable predator may be specific to each region.

In British Columbia, Canada, the coastal population of northern goshawks is the subject of federal and provincial management which focuses on the protection of breeding habitat to increase nest site availability (COSEWIC 2013). Like many raptors, goshawk populations are generally considered to be limited by both nest site availability and prey abundance (Reynolds et al. 2006, Rutz et al. 2006). However, current management plans do not include protections for foraging habitat or actions to increase prey populations, in part due to a lack of knowledge regarding goshawk diet and foraging behavior in this region. Goshawk diet in the coastal Pacific Northwest is variable, with hawks on Vancouver Island, British Columbia, consuming primarily red squirrels (Ethier 1999), whereas hawks in nearby southeast Alaska (Lewis et al. 2006) and western Washington (Bloxton 2002) take mostly medium and large birds. Even within British Columbia, a gradient of forest types from wet coastal to dry interior may produce dietary variation at smaller scales (Team 2008). Detailed local information on goshawk diet and its effect on reproductive success is necessary if limiting factors beyond nest site availability are to be included in management plans.

Here we describe the breeding season diet of northern goshawks in coastal British Columbia over a two-year period. We assess whether goshawk diet differs within this region between the wetter *coastal* zone and the drier *transition* zone. We further evaluate whether dietary variables (composition, diversity) influence goshawk reproductive success.

# Methods

### Study Area and Species



In North America, the northern goshawk ranges from boreal forests of the Yukon south to high-elevation forests of Arizona and New Mexico. Two subspecies are recognized: the widespread *atricapillus* and the limited *laingi* (Squires et al. 2020). The *laingi* subspecies was first described on the Haida Gwaii archipelago in British Columbia and is smaller and darker than the *atricapillus* subspecies found elsewhere on the continent (Taverner 1940). The range of this subspecies is limited to the west coast of North America from southeast Alaska through mainland British Columbia and Vancouver Island, possibly as far south as Washington’s Olympic Peninsula (COSEWIC 2013). *A. g. laingi* is considered a species at risk in British Columbia by both the federal and provincial governments due to significant habitat loss from industrial timber harvest (Team 2008, COSEWIC 2013).

We studied goshawks in southwestern British Columbia, a region characterized by rugged mountains interspersed with coastal fjords and low-lying valleys. The maritime climate supports temperate rainforest dominated by Douglas-fir (*Pseudotsuga menziesii*), western redcedar (*Thuja plicata*), and western hemlock (*Tsuga heterophylla*) (Meidinger and Pojar 1991). The goshawk population in southwestern British Columbia is currently classified as *A. g. laingi*, though new genetic evidence may lead to future reclassification (Geraldes et al. 2018). Within this region, goshawk managers have delineated a *transition zone* comprised of low-elevation valleys connecting the wet forests of the *coastal zone* with the dry interior forests east of the Coast Mountains. This narrow transition zone contains forest types intermediate between the western and eastern forests and likely represents an area of overlap between the coastal *laingi* population and the interior *atricapillus* population (Team 2008).

### Data Collection

We assessed goshawk diet during the 2019 and 2020 breeding seasons through a combination of egested pellets, prey remains, and nest camera photos. Active goshawk nests were located as part of long-term population monitoring conducted by the British Columbia Ministry of Forests, Lands, Natural Resource Operations and Rural Development (FLNRO) (for detailed survey methodology see McClaren (2005)). Some study sites contained active nests in both years of the study.

Prey remains and egested pellets were collected from 33 nests (2019 *n* = 18, 2020 *n* = 15). Pellets and remains were gathered from beneath active nests, from within nests after juveniles fledged, and from plucking posts located within the site. Pellets and remains were collected once during the breeding season at most sites, but some sites were visited multiple times. All prey remains and all pellets from a collection location (i.e., one nest or one plucking post) were combined into a single sample for each visit to that location.

At a subset of these nests (2019 *n* = 6, 2020 *n* = 7), cameras were installed to record prey delivered to goshawk chicks. Nest cameras are an effective and relatively unbiased method of measuring avian diet (García-Salgado et al. 2015 , Harrison et al. 2019). However, cameras may overestimate prey deliveries because goshawks cache prey items for redelivery to the nest at a later time, which creates a risk of double-counting items. Due to the discrete nature of our data we were unable to differentiate cached, re-delivered items from new items and did not attempt to account for caching in our analysis. We also did not attempt to differentiate between prey consumed by the female at the nest and prey consumed by the chicks.

Nest cameras were digital trail cameras (Reconyx brand, UltraFire and HyperFire models) mounted 2-5 meters distant from and slightly above the nest, usually in an adjacent tree. Cameras in 2019 were programmed to take three photos one second apart when triggered by motion, and an additional one photo every thirty minutes. In 2020, cameras were programmed to take five photos one second apart when triggered by motion, and an additional one photo every twenty minutes. Installation took place during the early nestling phase (between 4 June and 18 July; mean installation date 21 June) and cameras were left in place until after juveniles dispersed in the fall. We observed no nest abandonment following camera installation. Camera site selection was not random but constrained by topography, access, and timing of discovery.

Breeding chronology was not available for most sites. At 12 of the 13 nests with cameras (2019 *n* = 6, 2020 *n* = 6), chicks were aged from photos taken shortly after camera installation using a pictorial guide (Boal 1994). Productivity was defined as the number of chicks to reach 32 days age (Boal 1994, McClaren et al. 2002).

### Diet Quantification

We reconstructed prey from pellets and prey remains following a modification of the protocol used by Lewis et al. (2004). Within each sample, remains were identified to the lowest possible taxonomic category and the minimum number of individuals counted (i.e. 3 hare femurs = 2 *Lepus americanus*). Intact pellets and broken but reassembled pellets were analyzed individually within each sample, while fragmented pellets were combined within each sample. Pellets were dissected and feathers, fur, and hard parts (bones, teeth, claws) were identified to the lowest taxonomic level. We counted the minimum number of individuals represented within the pellet or pellet collection. Prey items from pellets and remains were additionally categorized by size (small = sparrow- or vole-sized, medium = jay- or squirrel-sized, and large = grouse- or hare-sized).

Prey items identified to species were assigned mass using data from the literature. We assigned mass to mammals from Nagorsen (2002) and to birds from Billerman et al. (2020), using the geographically closest estimates available and averaging the mass of males and females. We treated some relatively homogenous genera (such as *Eutamias* and *Myotis*) as a single species. For these genera, we assigned mass by averaging the masses of all possible species, based on range maps. Red squirrels (*Tamiasciurus hudsonicus*) were only present at a single site within our study area; when unable to distinguish between the two members of the genus *Tamiasciurus* we assigned the item to the more common *T. douglasii*. Unidentified grouse were common among remains; these were assigned the mean mass of the two grouse species present in our study area (*Bonasa umbellus* and *Dendragapus fulignosus*). Juveniles were assigned 50% of adult mass. Unidentified items were assigned mass by averaging the masses of the identified species in that size category and taxonomic class.

Data from prey remains and egested pellets are known to be biased indices of diet. Some authors have found combining data from both sources to produce relatively unbiased results that can serve as a helpful supplement to nest camera data (Simmons et al. 1991, Lewis et al. 2004). After testing for differences between pooled pellets-and-remains data and camera data we found significant differences between these two sources. We therefore report results from pellets, pooled pellets-and-remains, and cameras separately. We do not report results from prey remains alone, as diet composition estimates from prey remains are highly biased and infrequently used in raptor diet studies.

Nest camera photos were reviewed and each new prey item delivered to the nest was recorded and identified to species when possible. When items could not be identified to species, they were identified to the lowest possible taxonomic level. Prey items identified from photos were assigned a size category and biomass by the same method used for remains and pellets. Partial items were assigned the average mass for that size category and taxonomic class.

We quantified goshawk diet across the entire study area in several ways using data from pellets, pooled pellets-and-remains, and nest cameras. For ease of comparison, we simplified prey items into eight broad categories: tree squirrels (genus *Tamiasciurus*), hares (genus *Lepus*), all other mammals, grouse (subfamily Tetraoninae), thrushes (family Turdidae), corvids (family Corvidae), all other birds, and unidentified items. We calculated the relative proportion of avian and mammalian biomass, as well as the relative proportion of biomass composed of tree squirrels (genus *Tamiasciurus*), which are known to be an important source of prey for goshawks in British Columbia (Ethier 1999). For nests with cameras, we additionally quantified diet at the level of the individual nest and further calculated diet diversity with Simpson’s Diversity Index (Simpson 1949) using counts of items identified to genus or better. We report all diet quantification as percent of biomass or mean percent biomass the standard deviation except where counts or percents of items are explicitly specified.

### Statistical Analysis

Sites were classified as either coastal or transition based on whether the site was centered within the transition zone defined by Team (2008). We used counts of items assigned to the eight broad prey categories to assess differences in goshawk diet between the coastal and transition zones. We combined all data within each zone and tested each data source separately using a chi-squared test with simulated Monte Carlo *p*-values (2000 permutations) due to small sample sizes (Hope 1968). For nests with cameras, we also calculated the proportion of squirrel biomass and diet diversity at the individual nest level and compared these between the zones using a *t*-test. Finally, we tested for differences in productivity between the two zones using a *t*-test.

To determine the potential reproductive consequences of dietary variation, we examined how two aspects of diet, diet diversity and the proportion squirrel biomass in the diet, influenced productivity using linear regressions. We included all available nests in this analysis, including one site which contained an active nest in both years of the study. The presented results are not altered if variables for this site are averaged across years or a single year is randomly selected for inclusion. Because productivity data were available only from sites with nest cameras and nest-level diet data from pellets and prey remains were sparse, we performed this analysis using only diet data from nest cameras. All analyses were performed in R (Team 2020). We used a significance level of *P* = 0.05 for all tests.

# Results

## Goshawk Diet

|  | | | | Camera | | Pellets | | Remains | |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| Class | Common name | binomial | N | % items | % biomass | % items | % biomass | % items | % biomass |
| Aves | Unknown |  | 157 | 14.22 | 4.76 | 25.19 | 11.10 | 48.76 | 54.86 |
| Aves | Northern goshawk | Accipiter gentilis | 2 | - | - | - | - | 1.65 | 1.81 |
| Aves | Mallard | Anas platyrhynchos | 1 | - | - | - | - | 0.83 | 1.60 |
| Aves | Ruffed grouse | Bonasa umbellus | 4 | 0.67 | 1.65 | - | - | 0.83 | 0.74 |
| Aves | Swainson's thrush | Catharus ustulatus | 7 | 1.33 | 0.23 | 0.74 | 0.16 | - | - |
| Aves | Northern flicker | Colaptes auratus | 14 | - | - | 6.67 | 7.75 | 4.13 | 1.11 |
| Aves | Northwestern crow | Corvus caurinus | 1 | - | - | - | - | 0.83 | 0.55 |
| Aves | Steller's jay | Cyanocitta stelleri | 18 | 1.33 | 0.81 | 2.96 | 2.80 | 6.61 | 1.35 |
| Aves | Sooty grouse | Dendragapus fulignosus | 8 | 0.67 | 3.33 | - | - | 4.13 | 6.69 |
| Aves | Downy woodpecker | Dryobates pubescens | 3 | 0.67 | 0.11 | - | - | - | - |
| Aves | Hairy woodpecker | Dryobates villosus | 4 | 0.44 | 0.20 | 1.48 | 0.88 | - | - |
| Aves | Pileated woodpecker | Dryocopus pileatus | 3 | 0.22 | 0.36 | - | - | 1.65 | 0.81 |
| Aves | Varied thrush | Ixoreus naevius | 19 | 2.00 | 0.90 | 1.48 | 0.87 | 6.61 | 0.84 |
| Aves | Mew gull | Larus canus | 1 | - | - | - | - | 0.83 | 0.55 |
| Aves | Band-tailed pigeon | Patagoienas fasciata | 4 | 0.67 | 1.44 | - | - | 0.83 | 0.53 |
| Aves | Gray jay | Perisoreus canadensis | 5 | 1.11 | 0.44 | - | - | - | - |
| Aves | Spotted towhee | Pipilo maculatus | 3 | - | - | 1.48 | 0.46 | 0.83 | 0.06 |
| Aves | Red-breasted sapsucker | Sphyrapicus ruber | 5 | - | - | 1.48 | 0.64 | 2.48 | 0.25 |
| Aves | Barred owl | Strix varia | 2 | 0.22 | 0.90 | - | - | 0.83 | 1.01 |
| Aves | American robin | Turdus migratorius | 4 | 0.89 | 0.35 | - | - | - | - |
| Aves | TOTAL |  | 265 | 24.44 | 15.50 | 41.48 | 24.65 | 81.82 | 72.75 |
| Mammalia | Unknown |  | 38 | 2.89 | 6.75 | 17.04 | 14.34 | 1.65 | 2.03 |
| Mammalia | Mountain beaver | Aplodontia rufa | 1 | 0.22 | 1.27 | - | - | - | - |
| Mammalia | Flying squirrel | Glaucomys sabrinus | 3 | 0.67 | 0.59 | - | - | - | - |
| Mammalia | Snowshoe hare | Lepus americanus | 17 | 1.11 | 8.46 | - | - | 9.92 | 22.65 |
| Mammalia | Bat | Myotis sp | 1 | 0.22 | 0.01 | - | - | - | - |
| Mammalia | Chipmunk | Neotamias sp | 8 | 1.56 | 0.59 | 0.74 | 0.36 | - | - |
| Mammalia | Bushy-tailed woodrat | Neotoma cinerea | 2 | 0.22 | 0.47 | - | - | 0.83 | 0.53 |
| Mammalia | Rat | Rattus sp | 19 | 4.22 | 6.47 | - | - | - | - |
| Mammalia | Shrew | Sorex sp | 1 | 0.22 | 0.01 | - | - | - | - |
| Mammalia | Douglas squirrel | Tamiasciurus douglasii | 240 | 39.78 | 46.00 | 40.74 | 60.65 | 4.96 | 1.72 |
| Mammalia | Red squirrel | Tamiasciurus hudsonicus | 10 | 2.00 | 2.55 | - | - | 0.83 | 0.32 |
| Mammalia | TOTAL |  | 340 | 53.11 | 73.17 | 58.52 | 75.35 | 18.18 | 27.25 |
| Unknown | Unknown |  | 101 | 22.44 | 11.34 | - | - | - | - |
|  | TOTAL |  | 706 | 100.00 | 100.00 | 100.00 | 100.00 | 100.00 | 100.00 |

We identified a total of 9 unique species from pellets collected at 25 nests. Of the 135 prey items obtained from pellets, 58% could be identified to genus or better and all items were identified at least to class. There were 18 species identified from remains collected at 30 nests, for a total of 21 unique prey species from the pooled pellets-and-remains sample. Of the 121 prey items obtained from remains, 51% were identified to genus or better and all items were identified at least to class.

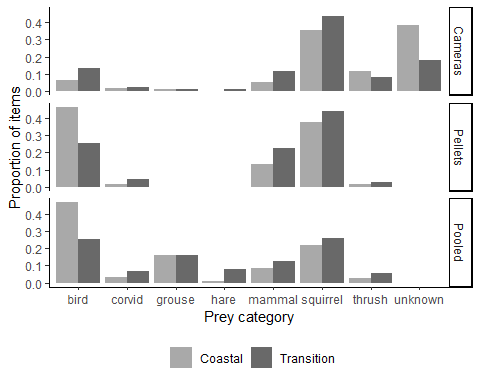
The majority of prey (75% of biomass) identified from pellets was mammalian. Squirrels made up 61% of biomass, while other birds (neither grouse, corvids, nor thrushes) made up 21% and other mammals (neither tree squirrels nor hares) made up another 15%. The remaining 4% of biomass was made up of corvids and thrushes.

In contrast, the majority of prey (63% of biomass) identified from the combined pellets-and-remains sample was avian. The largest prey group was grouse (37%), followed by other birds (23%) and hare (18%). Squirrels made up only 14% of the combined sample. The remaining 3% of biomass was made up of corvids and thrushes.

We identified a total of 22 unique species from 518 prey deliveries recorded on 13 nest cameras. After excluding 68 deliveries that were completely obscured from the cameras, each nest contributed an average of 35 items (range 1 - 69). We were able to identify 60% to genus or better, and 78% at least to class. Small and medium birds were disproportionately represented among items identified only to class, frequently arriving at the nest already plucked and decapitated. Variability in the number of items recorded and the rate of identification was due to differences in camera placement and sensitivity settings.

The majority of prey (73% of biomass, mean = 71 12) identified on nest cameras was mammalian. Birds accounted for only 15% (mean = 16 10), and the remaining 11% could not be identified to class (mean = 20 27). The high proportion of mammalian biomass was driven by the dominance of squirrels (49% of biomass, mean = 50 14) and other mammals (16%, mean = 17 12). Unidentified items accounted for 11% of biomass (mean = 20 27). The remaining biomass was composed of hare (8%, mean = 31 2), other birds (7%, mean = 7 8), grouse (5%, mean = 9 5), thrushes (3% , mean = 4 3) and corvids (1%, mean = 3 3). Overall diet diversity for the study area, based on counts of items identified to genus or better, was moderate (0.56). Diet diversity of individual nests was highly variable, ranging from 0 to 0.72 (mean = 0.46 0.21.)

## Difference between ecological regions

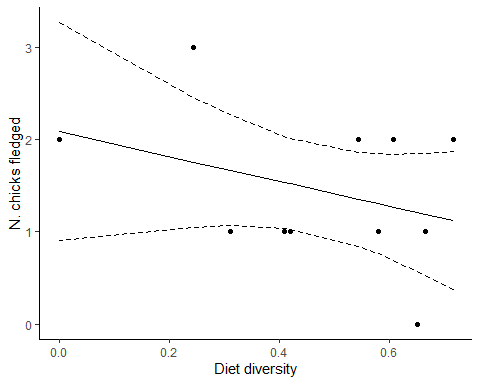
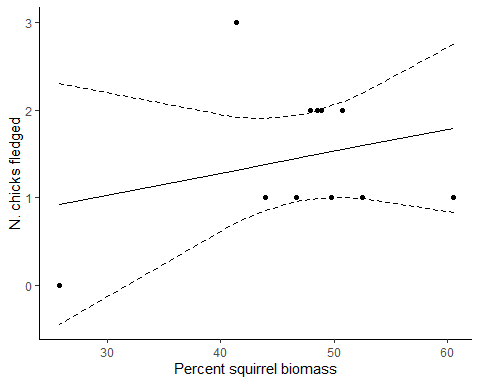


We observed differences in goshawk diet between the coastal and transition zone, although these differences were more pronounced in the camera data (2 = 23.84, *P* = 0.002) and pooled pellets-and-remains data (2 = 41.75, *P* << 0.05) than in data from pellets alone (2 = 7.53, *P* = 0.11). However, we observed no significant differences in diet diversity (*t* = -1.9, df = 4.13, *P* = 0.13) or the proportion of squirrel biomass (*t* = 1, df = 3.75, *P* = 0.38) between the zones.

## Productivity

We were able to measure productivity for 12 of 13 [**this will be updated to 14 when GRV data comes in**] nests monitored with nest cameras. Productivity data were not obtained from two nests because the camera memory cards filled prior to fledging.

Goshawks successfully fledged young from 11 of 12 nests, producing an average of 1.36 0.81 chicks per active nest (range 0-3) and 1.55 0.69 chicks per successful nest (range 1-3). Siblicide was common, accounting for two of the three deaths in the failed nest and one death in each of four other nests.



We found little evidence to suggest goshawk productivity was impacted by either the proportion of diet composed of squirrel biomass (*F*1 = 0.65, df = 9, *P* = 0.44) or diet diversity (*F*1 = 1.31, df = 9, *P* = 0.28). There was also no significant difference in goshawk productivity between the coastal and transition zones (*t* = 0.93, df = 4.71, *P* = 0.4).

# Discussion

Northern goshawks are considered generalist predators and consume a wide variety of prey species. However, a single key prey species often contributes disproportionately to goshawk diet, and the identity of this species may be specific to each goshawk population (Doyle and Smith 1994, Ethier 1999, Tornberg et al. 2005). In coastal British Columbia we found goshawks consumed 29 different species during the breeding season. These prey ranged in size from tiny bats (~ 6 grams) to grouse larger than the goshawks themselves (~ 1000 grams). However, the majority of their diet was composed of tree squirrels of the genus *Tamiasciurus*. Across the entire study area this single taxa made up approximately half of the total biomass recorded. Individual nests varied in the degree to to which they specialized on tree squirrels, with some nests consuming as little as 26% squirrel biomass and others as much as 86%. Despite the clear importance of tree squirrels in the diet, the proportion of squirrel biomass delivered to the nest did not influence its success in fledging chicks.

Goshawk survival, migration, reproductive success, and other demographic parameters are often related to the abundance of a key prey species (Doyle and Smith 1994, Tornberg et al. 2005, Rutz and Bijlsma 2006). Although we lacked data on prey abundance within our study area, other authors have found goshawk diet to reflect site-level prey abundance (Lewis et al. 2006, Rogers et al. 2006). We observed significant variation in the proportion of tree squirrel biomass delivered to each nest, which may reflect differences in squirrel abundance between sites. However, we did not find evidence to support an effect of this variation on goshawk productivity. When key prey abundance is low, high diet diversity may indicate a reliance on alternate prey and associated negative reproductive consequences (Resano‐Mayor et al. 2016), but we likewise found no evidence of an effect of diet diversity on goshawk productivity. However, it is notable that the only nest in our study to experience a complete breeding failure received the smallest proportion of squirrel biomass out of any nest. Given the strength of evidence from other studies and the clear importance of tree squirrels in the diet of this population, information on factors that influence the abundance of tree squirrels and their relationship to goshawk demography seems crucial.

Across much of North America, the key goshawk prey species is usually mammalian, often from the family Leporidae or Sciuridae (Boal and Mannan 1994, Doyle and Smith 1994, DeStefano et al. 2006, Rogers et al. 2006, Miller et al. 2014). However, in the coastal temperate rainforests of the Pacific Northwest, goshawk diet generally contains more birds than mammals and the key prey is usually a species of grouse (subfamily Tetraoninae) (Watson et al. 1998, Thraikill et al. 2000, Bloxton 2002, Lewis et al. 2006). Despite inhabiting coastal rainforests, goshawks on Vancouver Island, British Columbia, consume primarily red squirrels (*T. hudsonicus*), with this single species making up 69% of the diet (Ethier 1999). Our results from the coastal mainland of British Columbia are consistent with findings from Vancouver Island and more broadly with results from the interior of North America, but stand in contrast to findings from elsewhere in the Pacific Northwest.

Within our study area, low-elevation mountain valleys bridge the wet forests of the coast and the dry forests of the interior, creating a narrow region of intermediate habitat types where coastal and interior goshawk populations may overlap (Team 2008). At this smaller scale, we found evidence of a difference in diet composition between the coastal and transition zones. However, the identity of the key prey, tree squirrels, was the same in both zones and comprised an equally large proportion of the diet in each. We likewise found no difference in diet diversity between the zones. While the large-scale variation in goshawk diet across North America likely reflects regional abundance and availability of prey species, the small-scale variation within our study area may reflect the fine differences in prey availability between forest types.

Raptor diet is studied through a variety of indirect methods, such as the collection of pellets and prey remains, and direct methods, such as nest cameras and observation from blinds. Nest cameras are considered one of the least biased methods for measuring diet at the nest in raptors (Tornberg and Reif 2007, García-Salgado et al. 2015, Harrison et al. 2019). Cameras in this study provided significantly more data at a finer resolution than either pellets or prey remains, which could only be collected during the infrequent surveys each site received. However, the cost, effort, and logistical challenges of camera installation restricted the number of sites from which camera data could be collected. Additionally, technical issues relating to camera settings and placement resulted in a loss of data at some sites. Despite these limitations, we believe nest cameras provided the most accurate and complete picture of goshawk diet within out study. Compared to cameras, pellets were relatively unbiased in measuring coarse diet composition, but severely underestimated prey species richness. The pooled pellets-and-remains sample captured a much greater prey richness, including several species not detected on nest cameras, but greatly overestimated the proportion of avian biomass relative to camera data. Measuring diet composition by counts or biomass adds further uncertainty, with measurements of counts overestimating avian prey relative to measurements of biomass. These complex results highlight the importance of clearly reporting the source and measurement of raptor diet data. Because these methods have all been used in past studies we believe there is value in reporting the results of each for ease of comparison. However, we advocate for future diet studies to prioritize the collection of data via cameras, either video or still images, rather than physical specimens.

Our study addresses a fundamental question regarding the basic ecology of an at-risk population of the northern goshawk. This population is currently considered part of *A. g. laingi*, a subspecies restricted to the coastal Pacific Northwest from southeast Alaska through British Columbia and northwest Washington. In portions of the subspecies’ range the diet is dominated by avian prey (Bloxton 2002, Lewis et al. 2006), and in others by mammalian prey, specifically tree squirrels [Ethier (1999); this study]. This taxa clearly plays a key role in the diet of some *laingi* populations, including the population of Haida Gwaii (Roberts 1997, cited in COSEWIC (2013)), where red squirrel is an introduced species. Genetic evidence indicates goshawks on this isolated archipelago may be distinct from goshawks on the mainland coast and Vancouver Island (Sonsthagen et al. 2012, Geraldes et al. 2018). Regardless of their taxonomic relationship, dietary evidence suggests the goshawk populations of Haida Gwaii, Vancouver Island, and southwestern British Columbia are more similar to each other in foraging habits than to other putative *laingi* populations. Ecological similarity may therefore prove a better guide than genetic similarity when when incorporating foraging habitat or prey populations into management plans.

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