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1	Comparison of fatty acids and elemental nutrients in periphyton, invertebrates, and
2	cutthroat trout (Oncorhynchus clarki) in conifer and alder streams of western Washington
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Abstract

27 Organism growth and reproduction are often limited by nutrient availability in freshwater 28 ecosystems where, in some cases, food webs are primarily supported by allochthonous organic 29 matter. Therefore, we hypothesized that the composition of riparian vegetation would influence 30 the variability of N, P and fatty acid content of in-stream consumers. Specifically, we predicted 31 that organisms living in alder streams would have higher levels of N, P, and polyunsaturated 32 fatty acids than organisms in coniferous streams. To determine this, we sampled fresh and aged 33 leaf litter, periphyton, invertebrates, and cutthroat trout (Oncorhynchus clarki) from 6 streams in 34 western Washington state: 3 streams had high densities of nitrogen-fixing red alder (Alnus rubra) 35 in the riparian zone, whereas 3 had high densities of conifers. We found fresh alder litter had 36 twice the total polyunsaturated fatty acid concentrations of hemlock vegetation while there were 37 few statistical differences among aged alder and aged hemlock vegetation. Multidimensional 38 plots showed fatty acid profiles were unique to vegetation and fish while periphyton and 39 invertebrates shared the same multidimensional space. We used a mixed model to determine the 40 relative importance of vegetation type (fixed factor: conifer or alder), trophic levels (fixed factor: 41 periphyton, primary consumer, or fish) and streams (random factor) on individual fatty acid 42 concentrations. Total polyunsaturated fatty acids, 16:0, 20:1, 20:3n6 and total n3 were the only 43 fatty acids influenced by stream vegetation (vegetation + stream model or full model. 67% of 44 the fatty acids were best supported by the trophic +stream model. Nitrogen, P, Ca, Fe, C:N, N:P and C:N:P were all best supported by the trophic level + stream model and Zn was the only 45 46 nutrient supported best by the full model. Correlations of n3 and n6 fatty acid concentrations 47 between periphyton and primary consumers, and primary consumers with trout indicated several 48 fatty acid metrics, such as n3:n6, showed food resources may affect relative fatty acid

abundances of consumers. Although vegetation type did not influence relative fatty acids of
stream organisms, the importance of trophic level likely indicates organisms have different
physical requirements for fatty acids. The significance of a random factor, 'stream,' suggests
that the relative abundances of fatty acids in periphyton, invertebrates and trout are more related
than similar organisms from another stream.

Introduction

56 There is a strong connection between riparian vegetation and forest stream food webs 57 (Cummins et al. 1989; Richardson 1990; Kiffney et al. 2003). In general, food webs in these 58 streams depend on biomolecules from terrestrial sources, such as leaf litter or soil runoff 59 (Vannote et al. 1980; Barlocher 1992; Webster and Meyer 1997). The availability and quality of 60 riparian leaf litter varies widely, and this variability potentially influences consumer populations 61 (Volk 2004). Therefore, quantitatively assessing the chemical quality of a stream food web might 62 be useful for predicting in-stream production and fish growth. Food quality is commonly 63 assessed using C:N and N:P ratios, but essential fatty acids are an alternative measure of food 64 quality that has recently been applied to lake and stream ecosystems (Arts 1998; Arts et al. 65 2009). Fatty acid indicators are unique in that many animals, including humans, lack the 66 desaturation enzymes that act at the n3 and n6 positions of polyunsaturated fatty acids (PUFAs). 67 These fatty acids are critical for hormone production and membrane fluidity, and since they 68 cannot be produced they are essential dietary nutrients (Sargent et al. 1999). Furthermore, high 69 dietary concentrations of PUFAs, specifically n3 and n6 fatty acids, promote growth and 70 reproductive rates for aquatic invertebrates (Ravet et al. 2003; Brett et al. 2006). Few fatty acid 71 studies have assessed fatty acid compositions across trophic levels in natural systems (Torres-72 Ruiz et al. 2007) or environmental factors that might affect fatty acid composition (e.g. Peeters et 73 al. 2004) even though general fatty acid profiles of algae, invertebrates and fish are well 74 summarized by Arts et al. (2009).

Streams in the Pacific Northwest are generally oligotrophic and, depending on underlying
geology, can be limited by N, P or NP co-limited (Volk et al. 2008; Kiffney 2008; Sanderson et
al. 2009). Red alder, *Alnus rubra*, is a common nitrogen-fixing species found along riparian

78	corridors and disturbed landscapes of the Pacific Northwest. Alder leaf litter and underlying
79	soils are rich in N and P, and our earlier research showed annual detrital inputs were about $3.5 \times$
80	higher in streams dominated by riparian red alder relative to streams bordered primarily by
81	conifers (Volk 2004). Concentrations of a number of important biomolecules (e.g., N, P) in
82	water and fluvial particulate organic matter were also higher in alder-dominated streams. Alder
83	additions occur throughout the year with a large pulse during leaf fall in autumn, resulting in
84	annual total inputs of N and P to select streams of the Olympic Peninsula, WA, USA of 8.0 and
85	0.25 g/m ² /year, respectively (Volk 2004), which are 5 to 8 × higher than inputs to nearby
86	conifer dominated streams. Others have also shown that riparian alder forests are associated with
87	variability in the trophic productivity of freshwater ecosystems in the western US (Goldman
88	1961; Compton et al. 2003; Volk et al. 2003).
89	Therefore, we hypothesized the chemistry of detrital subsidies from riparian vegetation
90	may affect the chemical composition of local aquatic biota since N, P and fatty acids are

essential for survival, growth, and reproduction (Müller-Navarra 1995; Bendiksen et al. 2003).
To test this hypothesis, we quantified concentrations of PUFAs, C, N, P, Ca, Fe, and Zn in fresh
vegetation, stream-aged leaf litter, periphyton, invertebrates, and coastal cutthroat trout
(*Oncorhynchus clarki*) from 6 independent watersheds in western Washington: three stream
riparian corridors were dominated by alder and three by coniferous vegetation. Our objectives
were to compare n3 and n6 fatty acid concentrations and elemental nutrient concentrations of: 1)

97 food webs in alder and conifer-dominated streams; and 2) among trophic levels.

98

99

Methods

100 Study sites

101 Six headwater tributaries to the Hoh (47° 48' 36", -124° 5' 12") and Clearwater rivers (47° 42' 7", 102 -124° 10' 23") on the western Olympic Peninsula in Washington state were used as study sites 103 (Table 1). We selected these sites because of the predominance of alder or conifer within 30 m 104 of the stream bank, low accessibility to anadromous salmon and vehicle accessibility. Bridge 105 Creek, Bull Creek and Hook Creek were classified as coniferous streams because their riparian 106 corridors were dominated by ~75 year old second-growth Sitka spruce (*Picea sitchensis*), 107 western redcedar (Thuja plicata), western hemlock (Tsuga heterophyla), and Douglas-fir 108 (Pseudotsuga mensizeii). Christmas, Shale, and Maple Creeks were classified as alder streams 109 and were predominantly vegetated with red alder within the riparian corridor. Classifications 110 were assessed by measuring litter flux to streams using 5 baskets placed within the bankfull 111 channel of a 200 m reach; alder sites were required to have over 90% of the litterfall composed 112 of alder vegetation. We calculated the catchment area upstream of sample sites (NHDPlus 113 hydrogrpahy) and then the percent of land within this catchment covered by hardwood species 114 (data from Landsat Vegetation Mapping (1998) and GAP vegetation coverages (1991)) (Table 115 1). Alder is the dominant hardwood within these coastal streams and we considered the 116 hardwood and broadleaf GIS layers a reasonable proxy for alder composition within watersheds. 117 Sample collections

In September 2003, alder and hemlock vegetation, periphyton, invertebrates, and cutthroat trout were collected from a 200m reach of each stream for C, N, P, Fe, Ca, Zn and fatty acid analyses. Freshly senesced alder leaves and hemlock needles were shaken from three trees of each species and collected. Periphyton was scraped from rocks (5 samples/stream) with a toothbrush, rinsed with deionized water, filtered onto Whatman GF/F) filters, and frozen. Baetidae ($n_{alder} = 3$ individuals from a total of 2 streams, $n_{conifer} = 7$ individuals from a total of 2

124 streams), Heptageniidae ($n_{alder} = 11$ from 2 a total of 2 streams, $n_{conifer} = 4$ from 1 stream), and 125 Glossosomatidae ($n_{alder} = 6$ individuals from a total of 2 streams, $n_{conifer} = 6$ individuals from a 126 total of 2 streams) invertebrates were hand collected and frozen. Macroinvertebrate collections 127 were limited to these predominantly herbivorous and dominant families due to a limited 128 abundance of detritivores and shredders at the time of sampling. Five to ten cutthroat trout (fork 129 length of 6-15 cm) were collected from each stream using a single pass electrofishing survey, 130 weighed (nearest 0.1 g), measured (nearest 1 mm), and frozen. Single-pass electrofishing was 131 deemed a sufficient capture method as no fish population abundance estimates were planned for 132 the study (e.g. Bateman et al. 2005). All samples were frozen at -80 °C until fatty acid extraction 133 and C, N and P and micronutrient analyses.

134 In addition to freshly senesced vegetation, we aged leaf litter in streams to simulate in-135 stream detritus. We placed 10 g packets of senesced hemlock or alder in mesh bags (15 cm 136 diameter bags with 0.4 mm mesh) and zip-tied packets to large rocks in Christmas (alder 137 vegetation type) and Hook (conifer vegetation type) Creeks (note only one stream for each 138 vegetation type was used for packet placement and we considered individual packets as 139 replicates). Packets were submerged on 9/11/2003 and after 18 days packs were recovered and 140 placed in plastic bags for biomass measurements. We submerged leaf packs long enough to 141 allow diffusion of most nutrients and partial mass loss, as most leaf litter nutrients are lost within 24 hours of submersion in water (Gessner and Schwoerbel 1989) and 40% mass loss can be 142 143 found after four weeks (Braatne et al. 2007). In the lab, invertebrates were removed from packets 144 and remaining biomass was dried (30°C) to a constant weight and weighed. All weights used 145 within analyses are total biomass and were not corrected for inorganic matter accumulation. 146 Fatty acid extraction

147 A simultaneous extraction of wax esters and total fatty acids (Kattner and Fricke 1986; 148 Doerthe C. Müller-Navarra, University of Hamburg, Hamburg, Germany, personal 149 communication) was used for extraction of n3 and n6 fatty acids. Subsamples of leaf litter (fresh 150 and aged), periphyton, fish dorsal muscle tissue and whole invertebrates were freeze dried for 4 151 hours and weighed (sample weights ranged from 0.25-1.5mg, pending tissue type). Ten µl of the 152 internal standard, 21:0 (10mg/10ml methanol; Sigma #H-5149, heneicosanoic acid) was added 153 prior to three dichloromethane: methanol (2:1v/v) extractions. The first extraction was overnight 154 (15 hours) with 5ml of dichloromethane: methanol mixture, followed by a second and third 155 extraction of the sample material with 3 and 2ml of dichloromethane:methanol solution for 2-3 156 hours and 0.5 hours, respectively. Suspension liquid was removed, set aside, and chilled (32°C) 157 between extractions of source material. After all extractions were complete, set aside suspensions 158 were recombined and then evaporated with N₂ gas (30°C heat block), and resuspended with 2ml 159 of 3% sulfuric acid in methanol and 5ml 16% n-hexane addition. This mixture was heated for 4 160 hours at 80 °C, converting all Fatty Acids to Methyl Esters (FAMEs), which are soluble in 161 hexane. FAMEs were separated from the sulfuric acid matrix with four additional extractions 162 with n-hexane (water added to solution before extraction to dilute sulfuric acid matrix and 163 facilitate hexane:sulfulric acid solution separation), then N_2 gas evaporated to dryness and 164 dissolved in 1.5ml n-hexanes. All FAMEs were frozen (-80 °C) until injection (5µl) into the gas 165 chromatograph (GC).

Fatty acids were analyzed on an HP6890 series GC with an Agilent DB-WAX (30 m) + guard column (10m; 0.32 mm, 0.25 μ m film) and PTV inlet. Gas chromatography program specifics were 5 minutes at 40 °C (ramp rate = 10 °C/min), 5 minutes at 150 °C (ramp rate = 2 °C/min), 24 minutes at 220 °C (ramp rate = 2°C/min). The program was optimized for 18C fatty

acids and had a detection limit of 0.112mg dry weight. All program specifications were per the 170 171 University of California-Davis (Goldman Laboratory) specifications from Kattner and Fricke 172 (1986) and Doerthe C. Müller-Navarra (University of Hamburg, Hamburg, Germany, personal 173 communication). All fatty acids with retention times between 20 and 65 minutes were identified 174 by retention time on the chromatograph with comparison to reference standard (37 FAME, 175 Supelco Mix C4-C24). Reference peaks of interest included 10:0, 11:0, 12:0, 13:0, 14:0, 14:1, 176 15:0, 15:1, 16:0, 16:1, 17:0, 17:1, 18:0, 18:1n9, 18:2n6 cis and trans, 18:3n6, 18:3n3, 18:4n3, 177 20:0, 20:1, 20:2n*, 20:3n6, 20:4n6, 20:3n3 20:5n3 22:0, 22:1n9, 22:2n6, 23:0 24:0, 22:6n3, and 178 24:1. Once peaks were identified through comparison with the reference standard, areas for each 179 sample peak were corrected for the recovery volume of the internal standard, 21:0, and 180 multiplied by the total amount of sample (mg). No inferences on non-reference peaks were 181 made. Fatty acid analyses were replicated only within the study design (e.g. 6 detritus samples 182 per stream) and not for fatty acid analyses (e.g. 1 sample from stream extracted and run through 183 the GC multiple times). At least two blanks were included in each extraction and an additional 2 184 standards were included in each GC sample run.

185 *Elemental Nutrients*

Fresh alder and hemlock, aged alder and hemlock, periphyton, and trout muscle tissue were analyzed for C, N, P, Ca, Fe, K, and Zn. Material for elemental nutrient analyses was extracted from the same samples as fatty acids when there was ample material. Tissues were freeze dried for 4 hours and ground for C and N analysis in a CE440 Elemental Analyzer (Leeman Labs, Inc., University of Washington Oceanography Technical Services Laboratory, Seattle, WA). To determine P, Ca, Fe, and Zn concentrations, freeze dried material was digested with nitric acid for 12 hours, heated to 120 °C for 1 hour, oxidized with H₂O₂ until colorless and resuspended in 5% HCl (modified P digest from Jones et al. (1991)). Samples were run on an
ICP (Inductively Coupled Plasma Analyzer, NOAA, Northwest Fisheries Science Center,
Seattle, WA) and nutrient concentrations were calculated using standard curves for laboratory
standards. *Physical measurements*

One surface water grab sample in (September 2003) was collected for total N, total P,
ammonium, nitrite, nitrate, and phosphate analyses (Valderrama 1981). All water samples were
frozen (-80 ° C) and analyzed within 1 month of field collection. Additional physical habitat
details from a 200m survey of each stream during August-September 2003 are summarized in
Table 1 and described in detail in Volk (2004).

203 *Statistical analyses*

204 Since we assumed individual plants and leaf packs were independent samples, a one-way 205 ANOVA was used to compare nutrient composition (fatty acids or elemental nutrients) between: 206 a) fresh alder and fresh conifer vegetation and b) aged alder and aged conifer detritus. All data 207 were tested for normality (Shapiro test and Q-Q plots) and non-normal data were ln(x)208 transformed to meet assumptions of normality. Low C:N ratios and high Ca, N, P, Fe, and Zn 209 content were used as indicators of food quality for comparisons within the study. 210 Fatty acid and elemental nutrient data for fresh alder and hemlock, aged alder and 211 hemlock, periphyton, Baetidae, Glossosomatidae, Heptageniidae, and trout were compared 212 among all six streams using three mixed models in an information-theoretical approach 213 (Burnham and Anderson 1998). With fatty acids and elemental nutrient data as response metrics, 214 the models were designed such that vegetation type (classified by alder or conifer-dominated

215 vegetation) and trophic level (fresh vegetation-alder or hemlock, aged vegetation-alder or

- 216 hemlock, periphyton, invertebrates and fish) were used as predictive, fixed factors. 'Stream' was
- 217 used as a random factor to capture inherent natural differences among streams (aka sites).
- 218 Models
- 219 Model A (full model):
- 220 Response = vegetation + trophic level + stream
- 221 $y_i = \alpha_{\text{veg }i} + \beta_{\text{trophic }i} + b_i + \varepsilon_i$
- 222 Model B: Response = vegetation + stream
- 223 $y_i = \alpha_{\text{veg }i} + b_i + \varepsilon_i$
- 224 Model C: Response = trophic level + stream

225
$$y_i = \beta_{\text{trophic } i} + b_i + \varepsilon_i$$

- 226
- 227 y_i is the *i*th response data point across all streams
- 228 $\alpha_{\text{veg }i}$ has two values: alder and conifer
- 229 $\beta_{\text{trophic }i}$ has five values: vegetation, detritus, periphyton, invertebrates, and trout
- 230 $b_i \sim N(0, \sigma_{stream})$

231 $\epsilon_i \sim N(0, \sigma_{residual})$

232

233 Response metrics were averages of each fatty acid for each trophic level in a stream and were

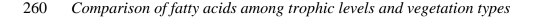
- transformed to meet assumptions of normality prior to analyses. Most fatty acids were
- transformed with log, square root, or cubed root transformations. We used arcsine-root
- transformations for C, N and P percentages. Akaike's Information Criteria corrected for small
- sample sizes (AICc) were compared among models to assess the relative importance of
- 238 vegetation type and trophic level on nutrient composition of sampled organisms. The relative

239 Akaike weight (w_i) is the relative likelihood of each model divided by the sum of all weights for 240 all models and was calculated for each model. We considered models with greater than 0.70 241 relative weights as strongly supported (AICc differences >2 from best model), relative weights 242 between 0.40 and 0.70 as moderately supported (AIC differences between 1 and 2 from best 243 model), and models with less than 0.40 relative weights as minimally supported by the dataset 244 (Burnham and Anderson 1998). Because vegetation type was a binary fixed factor (alder or 245 conifer), support for this model indicated alder and conifer response metrics are different. 246 Coefficients of the model were used to determine directionality associated with alder and conifer 247 differences.

248 We utilized a multi-dimension scaling (MDS) plot of $\ln(x+1)$ to describe the relative 249 fatty acid composition of in-stream organisms in multivariate space (Primer 6 Software, Clarke 250 and Gorley 2006). To investigate whether fatty acids are correlated between trophic levels, we 251 created a correlation matrix of stream averages of periphyton (n = 6) and primary consumers 252 Heptageniidae (n = 3), Baetidae (n = 3), Glossosomatidae (n = 4) and trout (n = 6) for each fatty 253 acid. Since not all invertebrate families were found in all streams, n values varied among 254 invertebrate families. We also correlated all primary consumers with trout similarly to 255 periphyton and invertebrates. Pearson's correlation coefficients were used to determine 256 significance of correlations.

- 257
- 258

Results



Of the three models, the trophic + stream model (Model C) best explained the variability for 67% of the fatty acids (Table 2). All monosaturated fatty acids and 7 of 9 saturated fatty acids were best supported by this model. The n3 and n6 fatty acids were of particular interest as they are essential fatty acids; half of the n3 and 5 of 6 of the n6 fatty acids were strongly supported by this model, all with relative weights greater than 0.70 (Table 2). We found similar results with elemental nutrients, where 7 of 8 elemental nutrients or nutrient ratios (e.g. N:P) were best supported by the trophic + stream model (Table 2).

268 Mean relative abundances of individual fatty acids were similar between alder and 269 conifer streams for periphyton, invertebrates and trout (Appendix 1). Although we expected 270 relative abundances of fatty acids would be influenced by riparian vegetation type, there was 271 almost no evidence that vegetation type influenced variation in fatty acid profiles, as the 272 vegetation + stream model (Model B) had low w_i values compared to the full and trophic + 273 stream models (Table 2). Two fatty acids were best supported by Model B (20:1 and 20:3n6) 274 and 3 fatty acid metrics (16:0, n3 and total PUFA) by the full model (Model A), but only five of 275 these showed relative abundances higher in alder than conifer; total PUFA was marginally higher 276 in conifer streams compared to alder streams. Despite limited support for relative abundances of 277 fatty acids, the influx of fatty acids from alder vegetation into streams may still be important to 278 stream ecosystems. We calculated the annual biomass inputs of fatty acids from senesced 279 vegetation by multiplying relative abundances of fatty acids by the estimated biomass of litterfall 280 from alder and conifer streams (Volk 2004) (Appendix 2). Annual alder biomass inputs were 281 \sim 3.5 times greater than conifer biomass inputs, this difference in detrital flux offers some 282 perspective on the total contribution of fatty acids to stream ecosystems from alder forests.

The general patterns of relative abundances of fatty acids among trophic levels can be seen in Table 3. Invertebrates and trout had higher relative abundances of n3 fatty acids than periphyton. Trout had highest levels of 22:6n3 relative to all other trophic levels. Vegetation and periphyton had higher relative abundances of n6 fatty acids than invertebrates and trout. Relative abundances of saturated fatty acids (SAFA) were also highest in vegetation and declined as trophic level increased (Table 3).

289 There was some evidence that fatty acids were conserved in these stream food webs. The 290 n3:n6 ratio was positively correlated between periphyton and consumers (r=0.67, p=0.03, n 291 =10, Figure 1) and consumers and trout (r = 0.57, p = 0.08, n = 10) (Figure 2). Furthermore, the 292 relative concentration of 18:3n3, was positively correlated between periphyton and consumers (r 293 = 0.76, p < 0.01, n = 10). Variability in 18:3n3 were positively correlated between consumers 294 and trout but this was not statistically significant (r = 0.53, p = 0.11, n = 10). Percent PUFA was 295 also positively correlated between primary consumers and trout (r = 0.63, p = 0.03, n = 10) but 296 not between periphyton and consumers (r = 0.13, p = 0.70, n = 11). The other n3 and n6 297 polyunsaturated fatty acids exhibited no correlations between different trophic levels (results not shown). 298

Results from our multidimensional plots showed fatty acid profiles of trout were tightly grouped and distinct from vegetation, periphyton and invertebrates. All vegetation (fresh alder, aged alder, fresh hemlock and aged hemlock) showed considerable overlap, but were distinct from periphyton, invertebrates and trout (Figure 3). The invertebrate families (Baetidae, Heptageniidae, and Glossosomatidae) did not separate in multidimensional space. Similarly, periphyton fatty acid profiles were highly variable and overlapped with all three invertebrate families.

307	Nutrient concentrations of decomposing alder and conifer litter in streams
308	We compared fatty acid concentrations from the two streams (1 alder and 1 conifer) after
309	18 days of decomposition (samples were used as replicates). Eighteen carbon fatty acids were
310	10-30% lower in both aged alder and hemlock compared to fresh litter concentrations, and the
311	relative abundances of 50% of measured PUFAs dropped (Table 3). N3 and n6 fatty acid
312	concentrations decreased by 30% in alder but increased by 5-25% in hemlock (Figure 4 and
313	Table 3). In alder, 18:4n3 concentrations declined by 77%, but increased by 40% in hemlock,
314	while 18:2n6 concentrations were reduced in both hemlock and alder (44 and 50%, respectively)
315	(Figure 1). 18:3n3 was the only polyunsaturated fatty acid where aged alder concentrations were
316	statistically higher than aged hemlock concentrations ($n = 12, p < 0.01$).
317	Elemental concentrations of Ca, N and P were significantly higher in fresh alder than
318	hemlock vegetation, and C:N ratios were significantly lower (percent by weight) (Table 4).
319	Nitrogen, C, and Zn were significantly higher, and C:N ratios were almost 3 times lower in aged
320	alder detritus relative to aged hemlock. Percent P was not statistically different between aged
321	vegetation types ($n = 18$, $p = 0.75$, Table 4). In alder, 36-67% of Ca was lost over 18 days
322	whereas C loss rates were twice as fast in hemlock than alder (1.00 and 0.53% per day,
323	respectively).
324	

325

Discussion

This is one of the first studies to investigate fatty acids in natural stream food webs, and factors that may affect the variability of these essential biomolecules. Overall, most of the variability in fatty acids was attributed to trophic level and stream, with vegetation type as an important covariate for only a few fatty acids. We speculate the importance of trophic level and
stream suggests organisms were not limited in essential fatty acids and that metabolic differences
likely accounted for the significance of trophic level for almost all fatty acid and nutrient
response metrics.

333 One of the more intruiging results was the positive correlation among the relative 334 abundance of fatty acids in different trophic levels (periphyton, consumers, and fish). The 335 positive correlation between stream trophic levels for some fatty acids suggests consumers reflect the chemical composition of their food resources. Specifically, levels of n3 and n6 fatty 336 337 acids in periphyton were positively correlated with fatty acids in primary consumers and trout, 338 suggesting these materials were conserved as they moved up the food chain. Feeding trials of 339 phytoplankton (cryptophytes, chlorophytes and cyanophytes) to *Daphnia pulex* showed high 340 correlations of 20:5n3 + 22:6n3 and 20:4n6 between food sources and *Daphnia* (Brett et al. 341 2006). Similarly, feeding trials using 18:2n6, 18:3n3, 20:4n6 or an n3 PUFA mix for laboratory 342 reared Arctic charr found a general dominance of n3 PUFA in fish muscle tissue when diets were 343 composed of 18:3n3 (Olsen et al. 1991), supporting our observations that a high PUFA content 344 in invertebrates correlated to high PUFA content in fish tissue. However, our study was 345 observational and exploratory, which limits our inference, but suggest some potential future 346 studies evaluating how fatty acids may influence food webs in stream ecosystems. For example, 347 does natural variation in essential fatty acids influence growth rate of stream fish through higher 348 growth efficiencies?

For elemental nutrients, lower C:N ratios in periphyton than terrestrial vegetation
indicates that aquatic primary producers were a higher quality food for consumers and predators
than terrestrial producers, and similar conclusions have been drawn by Reiners (1986) and Elser

352 et al. (2000). This is likely due to the high proportion of structural carbon material in terrestrial 353 vegetation (McGroddy et al. 2004). Furthermore, all else being equal, the lower C:N of algae 354 and alder may result in differences in growth efficiencies of higher trophic levels relative to 355 systems dependent on more recalcitrant sources of energy such as conifer needles, as conifer 356 needles break down more slowly than deciduous litterfall (Alberińo and Balseiro 2002). In both 357 terrestrial and lake environments, a reduction in the conversion efficiency of carbon into new 358 biomass ('gross growth efficiency') has been correlated with higher food C:N and C:P ratios 359 (Elser et al. 2000).

360 Trout, vegetation and invertebrates separated well in multidimensional space while 361 periphyton tended to overlap with vegetation and invertebrates. The high percentages of PUFA 362 (56%), and 22:6n3 (5 times higher relative abundance than other trophic levels) likely 363 contributed to the separation of trout from other trophic levels. Metabolic differences among 364 trout and invertebrates likely account for the separation of the two consumers, as the essential 365 fatty acid requirements for growth and cell structure are different (Ackman 1998). The 366 considerable overlap between periphyton and primary consumers was expected, as these 367 invertebrates were predominantly grazers, scrapers, and collectors that feed on periphyton 368 (Cummins 1973). In general, Baetidae feed by scraping algae and fine detritus from submerged 369 rocks or other submerged materials, such as woody debris. Heptageniidae are surface feeding 370 collectors or scrapers (mineral or organic scrapers) and Glossosomatidae are typically mineral 371 scrapers (Cummins 1973). The overlap between the fatty acid profiles of these invertebrate 372 families and periphyton is therefore not surprising and we assume periphyton is the dominant 373 food source for the sampled invertebrates in the study streams.

374 Results from this study suggest the relative abundance of a few fatty acids or n3:n6 ratios 375 of fatty acids can be used to assess the potential biological (e.g., growth, survival, productivity) 376 importance of different chemical components in stream food web (riparian plants, periphyton, 377 primary consumers, and trout. This suggests that streams with high relative abundance of these 378 fatty acids in periphyton will have similarly high abundances of these fatty acids in grazers and 379 trout. However, the majority of correlations among trophic levels for n3 and n6 fatty acids were 380 not significant, suggesting that some fatty acids may be better tracers than others. Further 381 studies considering feeding habits and trials in mesocosm environments with natural food 382 sources are needed especially those that quantify whether these fatty acids actually contribute to 383 variation in performance in higher trophic levels.

384 One of our main objectives of this study was to determine if leaf litter inputs from alder 385 vegetation influence the relative abundance of fatty acids of in-stream organisms because 386 primary producers are the only source of these fatty acids for higher trophic levels. We did find 387 higher abundances of PUFA, 18:3n3, 20:5n3, 20:4n6 and 22:6n3 fatty acids, and higher N and P 388 in leaf material from alder relative to hemlock vegetation (N and P results similar to Volk 2004). 389 However, the results presented here provided little evidence that fatty acids and elemental 390 nutrients in periphyton, invertebrates and trout were strongly influenced by vegetation type. We 391 suggest three potential reasons for this result: 1) the relative abundance of only 5 fatty acids was 392 higher in alder than conifer vegetation, suggesting there are very few fatty acids where we would 393 have expected to see influences from alder vegetation; 2) low sample sizes, especially for 394 invertebrates, did not provide enough power of detection for this study; and 3) polyunsaturated 395 fatty acids are not limiting food webs in these small streams and therefore relative abundances do 396 not change with the presence of additional fatty acid resources.

397 Instead, our model results indicated trophic level and streams were important covariates 398 predicting variation in biomolecules. The importance of trophic level likely indicates organisms 399 at different trophic levels have different metabolic requirements for fatty acids, particularly 400 essential fatty acids. The significance of a random factor, 'stream,' suggests that the relative 401 abundances of fatty acids in periphyton, invertebrates and trout are stream-specific and are 402 responding to local environmental or communal variables. Physical aspects or food resources 403 unique to each stream could influence the fatty acid profiles of these food webs. For example, 404 this 'stream effect' may be linked to differences in the composition of in-stream primary 405 producers, litterfall, other inputs of biomolecules (e.g. plant reproductive structures) or feeding 406 relationships. However we investigated a limited number of food resources and physical aspects 407 of a site that may contribute to this random variation in fatty acid profiles.

Aging leaf litter in streams reduced the relative abundances of n3 and n6 fatty acids, and this may be due to an accumulation of inorganic material on aged leaf litter. This hypothesis may be supported by the large increase in Fe between fresh and aged material, as Fe is common element in inorganic minerals. Other changes in n3 and n6 fatty acid content between fresh and aged material may have been due to algal or microbial colonization, but these communities were not directly studied.

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Conclusions

Relationships between 1) periphyton and invertebrate primary consumers and 2)
invertebrate primary consumers and fish suggest that the relative abundance of these resources
were conserved. Further observational and experimental studies are needed to improve our
understanding of the nutritional ecology of freshwater ecosystems, because this understanding

420	may help us conserve and restore ecologically and economically important fish species and their
421	ecosystems. Although the widespread abundance of riparian red alder in the Pacific Northwest
422	provides particulate and dissolved nutrient resources, it is difficult to discern the role of these
423	resources for primary and secondary consumers unless the limiting resources of the local
424	ecosystems are known. Moreover, we need a better understanding of the linkages between
425	watershed and riparian conditions that may affect chemical constituents potentially important in
426	the trophic productivity of freshwater food webs because this understanding may improve our
427	restoration and management of forested watersheds with economically and ecologically
428	important fish (Wipfli and Baxter 2010).
429	
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430 431	Significant assistance was provided by Anne Liston and the University of California Limnology
430 431 432	Significant assistance was provided by Anne Liston and the University of California Limnology Laboratory, Michael Brett (University of Washington), and Brice Semmens (Scripps Institute of

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- salmonid watersheds. Fisheries 35: 373-387

556 Table 1. Physical stream habitat characteristics of study sites

5<u>57</u>

	Stream					
	Maple	Shale	Christmas	Bridge	Hook	Bull
Vegetation	Alder	Alder	Alder	Conifer	Conifer	Conife
Alder in watershed, %	16.0	25.0	62.0	46.0	0	(
Alder leaf litter in riparian area, %	100.0	97.0	91.0	10.0	34.0	18.0
Alder leaf litterfall in riparian area, g/m^2	15.2	59.8	81.5	1.8	12.5	8.2
Conifer leaf litterfall in riparian area, g/m^2	0	2.5	6.9	7.9	16.4	12.
Stream gradient, %	2.7	2.4	1.2	5.1	3.6	1.
Discharge, L/s	16.9	21.6	17.4	1.3	153.0	1.
7d average T, °C	12.0	11.5	12.0	11.8	11.7	11.
TP, μg/L	26.0	35.0	25.0	25.0	29.0	28.
TN, μg/L	102.0	169.0	85.0	219.0	161.0	165.
$PO_4, \mu g/L$	4.4	4.8	2.9	7.5	4.0	2.
$NO_3, \mu g/L$	64.0	30.0	13.0	195.0	84.0	81.
NH_4 , $\mu g/L$	6.7	5.7	8.0	4.4	1.4	5.

Table 2. AIC scores and relative model weights (w_i) for the full, vegetation and trophic models. NA = Data not analyzed using mixed models due to high zero counts in data that led to violations in normality assumptions. Response metrics are relative fatty acid abundance or elemental nutrient concentrations per sample. * values calculated using subset of data for model due to abundance of zeros within trophic levels. ⁺ data did not conform well to assumptions of normality. ⁺⁺ 3 models not completed due to limited amount of non-zero data in all trophic levels. 565

	AICc				Wi	
Response	Full model	Vegetation	Trophic	Full	Vegetation	Trophic
metric + stream		+ stream	model	+ stream	+ stream	
	(Model A)	(Model B)	(Model C)			
14:0	-26.29	28.49	-31.49	0.07	0.00	0.93
15:0	-181.63	-178.34	-187.56	0.05	0.01	0.94
16:0	756.96	799.82	757.76	0.60	0.00	0.40
17:0	-194.43	-172.64	-201.57	0.03	0.00	0.97
18:0	-47.20	-70.42	-52.86	0.00	1.00	0.00
20:0	133.15	165.79	129.53	0.14	0.00	0.86
22:0	191.45	207.96	187.86	0.14	0.00	0.86
23:0	NA	NA	NA	NA	NA	NA
24:0*	112.65	127.95	109.29	0.16	0.00	0.84
SAFA	-218.74	-173.69	-225.29	0.04	0.00	0.96
14:1++	NA	NA	NA	NA	NA	NA
15:1++	NA	NA	NA	NA	NA	NA
16:1	707.06	788.95	708.77			
17:1 ⁺	164.85	194.02	161.01	0.13	0.00	0.87
18:1n9	256.17	263.12	252.70	0.15	0.00	0.85
20:1	0.048	-6.60	-4.44	0.03	0.00	0.25
22:1n9 ⁺⁺	NA	NA	NA	NA	NA	NA
24:1 ⁺⁺	NA	NA	NA	NA	NA	NA
MUFA	-122.45	-88.43	-129.29	0.03	0.00	0.97
18:3 n 3	305.95	350.38	304.45	0.32	0.00	0.68
18:4 n 3	-40.38	-44.40	-45.83	0.04	0.00	0.64
20:5 n 3	43.93	170.12	38.62	0.07	0.00	0.93
20:3 n 3*	60.82	62.77	57.21	0.13	0.00	0.92
20:5 n 3 ⁺	203.65	469.65	201.25	0.13	0.00	0.82
n3	848.63	1009.01	850.91	0.25	0.00	0.24
18:2 n 6	-106.36	-47.67	-111.17	0.08	0.00	0.92
18:2 n 6	117.40	125.15	113.06	0.10	0.00	0.92
20:3 n 6 ⁺	68.43	63.51	65.085	0.10	0.00	0.90
20:3 n 6 20:4 n 6	-65.06	23.04	-69.87	0.08	0.03	0.30
20:4 n 6 22:2 n 6*	-03.00 NA		-09.87 NA	0.08 NA	0.00 NA	0.92 NA
		51.90				
n6	-97.02	-92.27	-102.92	0.05	0.00	0.95
n3:n6	-85.82	14.96	-92.06	0.04	0.00	0.96
PUFA	871.74	989.69	874.67	0.81	0.00	0.19
C ⁺	-153.33	-74.43	-160.53	0.03	0.00	0.97
N	-244.45	-96.48	-252.08	0.02	0.00	0.98
P	64.00	119.59	60.82	0.17	0.00	0.83
Ca	70.23	91.02	67.35	0.19	0.00	0.81
Fe	86.26	218.36	82.42	0.13	0.00	0.87
Zn	110.21	114.95	110.25	0.48	0.04	0.47
C:N	-188.55	-76.31			0.00	0.93
N:P	36.50	46.72	34.48	0.27	0.00	0.73
C:N:P	38.25	123.95	32.74	0.06	0.00	0.94

569 Table 3. Summary of fatty acids in aquatic food web. Values are relative percentages of fatty

- 570 acid (or fatty acid ratio) averaged across all streams. N indicates the number of replicates (total
- 571 for all streams). Baetidae not collected from Hook Creek and Glossomatidae not collected from
- 572 Christmas Creek.

		Vege	etation						
Fatty acid	Fresh	Fresh	Aged	Aged	Periphyton	Baetidae	Hept	Gloss	Trout
	Alder	hemlock	alder	hemlock					
n =	6	6	6	6	27	10	15	12	36
14:0	3.00	3.75	3.92	4.59	4.52	1.25	0.92	3.29	0.80
15:0	0.38	0.66	0.49	0.41	0.78	0.26	0.69	033	0.18
16:0	25.88	20.10	24.45	22.37	30.25	27.66	20.25	20.96	21.29
17:0	0.56	0.85	0.69	0.55	0.49	1.19	1.53	0.84	0.63
18:0	6.88	11.98	5.61	8.74	8.06	9.32	13.14	4.32	7.03
20:0	3.69	4.08	3.80	3.72	1.08	0.49	1.88	0.45	0.20
22:0	6.53	2.75	2.53	4.02	1.02	0.39	2.28	0.43	0.15
23:0	0.43	3.02	0.42	0.15	0.02	0	0	0	0
24:0	3.73	3.97	3.99	7.49	1.44	0.01	0.06	0	0.01
SAFA	51.09	51.17	45.89	52.05	47.67	40.59	40.74	30.63	30.29
14:1	0.35	0.32	0.07	0.16	0.02	0	0.01	0.40	0
15:1	0	0.81	0	0	0	0	0.01	0.02	0
16:1	1.48	2.38	2.13	1.33	11.9	5.98	11.23	10.53	3.21
17:1	0.13	0.84	0.33	0	1.32	0.30	0.31	1.67	0.08
18:1n9	12.68	14.76	14.10	14.87	10.53	8.49	10.21	10.49	9.58
20:1	0.30	0	0	0	0.12	0.05	0.01	0.06	0.08
22:1n9	0	0.26	0	0.49	0.02	0.05	0	0	0.01
24:1	0	0	0	0	0.19	0	0	0.20	0.02
MUFA	14.63	19.37	16.63	16.85	24.11	14.88	21.78	23.01	12.98
18:3 n 3	18.98	8.79	21.17	10.37	8.01	19.18	8.26	17.37	7.41
18:4 n 3	1.78	0.74	0.67	0.64	1.38	2.21	0.98	3.64	1.21
20:5 n 3	0.78	0	1.14	3.43	7.16	14.08	19.39	16.62	11.87
20:3 n 3	0.22	0.29	0.21	0	0.06	0.18	0.35	0.23	0.70
22:6 n 3	0.27	5.38	0	3.38	1.07	0.29	0.22	0.23	28.41
n3	22.03	15.20	23.19	17.82	17.67	35.95	29.20	38.09	49.61
18:2 n 6	10.90	11.20	7.71	8.36	6.21	6.85	3.99	3.83	3.99
18:3 n 6	0.09	0.32	0	0	0.42	0.23	0.19	0.52	0.07
20:3 n 6	0	0	0	0.38	0.02	0.07	0.01	0.34	0.25
20:4 n 6	0.44	0	0.50	0	1.69	14.08	3.62	3.51	2.66
22:2 n 6	0	0	3.46	4.25	1.26	0	0	0	0.06
n6	11.43	11.52	11.67	12.99	9.70	8.34	7.81	8.20	7.03
n3:n6	2.09	1.52	2.32	1.73	2.26	4.57	3.96	5.71	7.51
PUFA	34.20	28.92	37.21	30.82	27.94	44.53	37.30	46.30	56.71
SAFA: (MUFA+PUFA)	1.05	1.06	0.85	0.92	0.95	0.78	0.61	0.44	0.43

575 Table 4. Summary of elemental nutrient concentrations for fresh and aged alder and hemlock

576 litter. Significance values are for ANOVA analyses comparing fresh alder to fresh hemlock and

aged alder to aged hemlock ($\alpha = 0.05$). Fe and Zn values not available for fresh hemlock due to

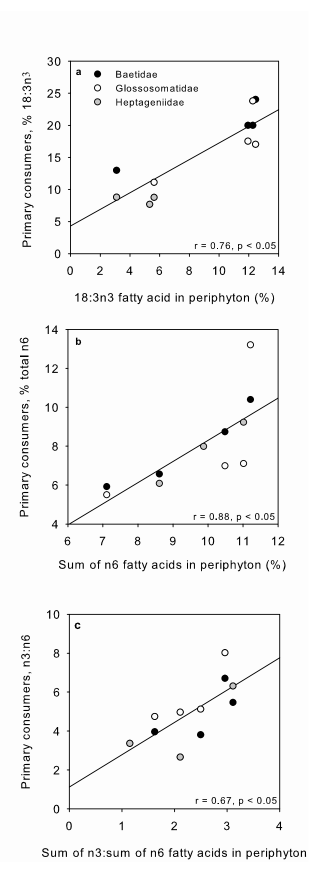
578 sample limitations.

579

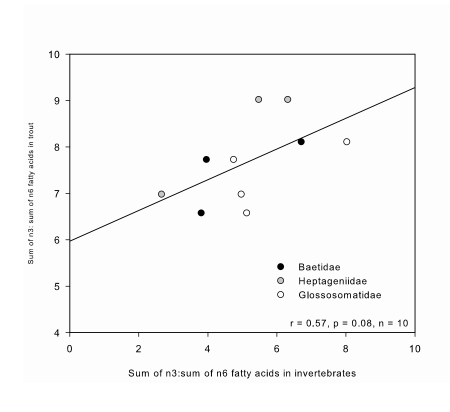
		Fresh				Aged			
	n	Alder	Hemlock	р	Alder	Hemlock	р		
C (%)	12	47.00	59.08	0.20	36.35	41.60	0.30		
N (%)	12	2.37	0.99	0.02	3.06	1.41	0.03		
P (%)	18	0.27	0.13	<0.01	0.32	0.22	0.75		
C:N (molar)	12	23.13	69.61	<0.01	14.38	34.28	<0.01		
C:P (molar)	12	459.81	1344.46	<0.01	6.03	23.89	0.04		
C:N:P (molar)	12	2.7×10^{6}	1.9×10^{6}	<0.01	2266.05	14261.83	0.04		
$Ca(\mu g/g)$	18	22846.00	4704.00	0.02	15302.00	5817.00	<0.01		
Fe $(\mu g/g)$	18	322.23	na	na	3957.00	2333.00	0.21		
$Zn (\mu g/g)$	18	252.08	na	na	4976.00	148.00	0.03		

580

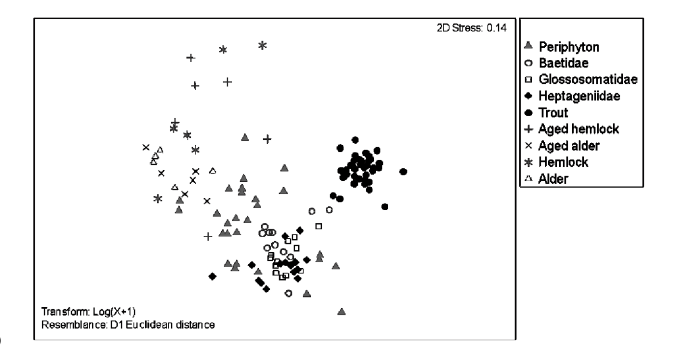
582	Fig 1 Correlations between periphyton and primary consumers for a) 18:3n3; b)n6 and c) n3:n6
583	fatty acids
584	
585	Fig 2 Correlation between primary consumers and trout for n3:n6 fatty acids
586	
587	Fig 3 Fresh and aged leaf litter, periphyton, invertebrates and trout fatty acids in
588	multidimensional space
589	
590	Fig 4 Major fatty relative abundances in fresh and aged vegetation. $n= 6$ for each series. sum n6
591	= sum of 18:2n6 (cis and trans) 18:3n6, 20:3n6, 20:4n6, and 22:2n6; SAFA = saturated fatty
592	acids; MUFA= monosaturated fatty acids, PUFA = polyunsaturated fatty acids. Bars indicate
593	standard error, * indicates significance between fresh alder and fresh hemlock at $p < 0.05$ and +
594	indicates significance between aged alder and aged hemlock at $p < 0.05$
595	

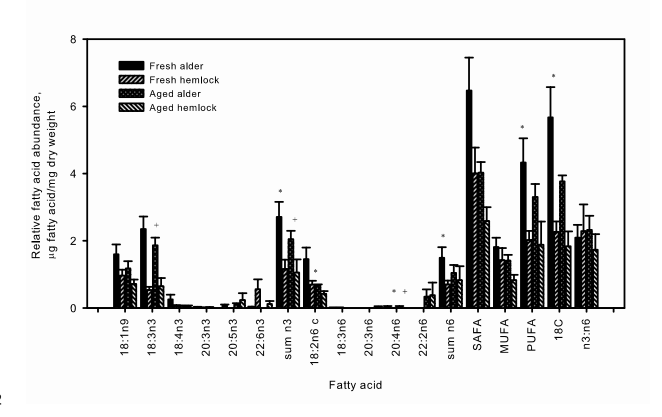












Appendix 1. Fatty acid summary for alder and conifer streams. Values are relative percentages of
fatty acid (or fatty acid ratio) averaged across all streams. N indicates the number of replicates
(total for all streams). Baetidae not collected from Hook Creek and Glossosomatidae not

- 606 collected from Christmas Creek.

Fatty	Periphyton	Periphyton	Baetidae	Baetidae	Hept	Hept	Gloss	Gloss	Trout	Trout
acid	Alder	Conifer	Alder	Conifer	Alder	Conifer	Alder	Conifer	Alder	Conifer
	14	13	3	7	11	4	6	6	13	23
<u>n</u> =										
14:0	5.82	3.13	1.59	1.21	0.92	0.92	3.68	2.90	0.77	0.83
15:0	0.97	0.57	0.23	0.27	0.79	0.40	0.29	0.37	0.17	0.19
16:0	31.47	28.94	26.84	27.74	19.97	21.02	22.30	19.63	21.64	21.01
17:0	0.56	0.41	0.90	1.22	1.61	1.31	0.70	0.99	0.57	0.67
18:0	8.93	7.13	6.22	9.64	13.68	11.65	3.78	4.87	6.63	7.35
20:0	1.38	0.75	0.28	0.51	1.96	1.65	0.37	0.53	0.15	0.25
22:0	1.23	0.79	0.15	0.42	2.40	1.96	0.39	0.47	0.10	0.19
23:0	0.05	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00
24:0	1.45	1.44	0.00	0.02	0.08	0.00	0.00	0.00	0.03	0.00
SAFA	51.86	43.16	36.21	41.02	41.41	38.91	31.50	29.75	30.05	30.50
14:1	0.04	0.00	0.00	0.00	0.01	0.00	0.06	0.02	0.00	0.00
15:1	0.00	0.00	0.00	0.00	0.01	0.00	0.03	0.01	0.00	0.00
16:1	11.61	12.22	10.08	5.57	12.50	7.76	10.41	10.66	2.84	3.51
17:1	1.37	1.27	0.47	0.28	0.31	0.30	1.68	1.65	0.07	0.08
18:1n9	9.22	11.94	7.62	8.59	10.27	10.04	10.85	10.14	10.65	8.70
20:1	0.20	0.04	0.00	0.05	0.00	0.04	0.10	0.02	0.10	0.07
22:1n9	0.03	0.00	0.00	0.06	0.00	0.00	0.00	0.00	0.01	0.01
24:1	0.16	0.22	0.00	0.00	0.00	0.00	0.00	0.40	0.04	0.00
MUFA	22.64	25.68	18.17	14.55	23.10	18.13	23.13	22.90	13.71	12.37
18:3 n 3	7.41	8.66	19.99	19.10	8.07	8.79	17.45	17.28	7.94	6.98
18:4 n 3	1.47	1.28	3.12	2.12	0.89	1.22	3.67	3.60	1.56	0.93
20:5 n 3	4.00	10.55	16.39	13.85	16.88	26.29	17.31	15.93	11.07	12.54
20:3 n 3	0.06	0.06	0.20	0.18	0.37	0.30	0.23	0.23	0.98	0.48
22:6 n 3	1.92	0.15	0.00	0.32	0.30	0.00	0.33	0.13	27.08	29.50
n3	14.86	20.70	39.70	35.57	26.51	36.60	39.00	37.18	48.63	50.41
18:2 n 6	5.03	6.66	4.73	7.06	4.34	3.03	3.20	4.46	4.44	3.62
18:3 n 6	0.15	0.72	0.38	0.22	0.19	0.17	0.46	0.57	0.07	0.07
20:3 n 6	0.03	0.00	0.00	0.07	0.02	0.00	0.11	0.58	0.31	0.20
20:4 n 6	1.48	1.92	0.81	1.23	3.89	2.88	2.53	4.49	2.54	2.75
22:2 n 6	2.42	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.13	0.00
nб	9.49	9.93	5.92	8.57	8.44	6.09	6.30	10.10	7.50	6.64
n3:n6	1.57	2.08	6.70	4.14	3.11	6.31	6.49	4.93	7.10	7.84
PUFA	25.02	31.09	45.62	44.42	35.25	42.94	45.30	47.30	56.22	57.11
608		22.07		=						

612		A nove1 1	Diamaga	
		Annual Biomass Input (g/m ² /year)		
613			•	
	Fatty acid	Fresh	Fresh	
614		Alder	hemlock	
615	14:0	10.5	3.75	
015	15:0	1.33	0.66	
(1)	16:0	90.58	20.1	
616	17:0	1.96	0.85	
	18:0	24.08	11.98	
617	20:0	12.92	4.08	
	22:0	22.86	2.75	
618	23:0	1.51	3.02	
018	24:0	13.06	3.97	
	SAFA	178.82	51.17	
619	14:1	1.23	0.32	
	15:1	0	0.81	
620	16:1	5.18	2.38	
020	17:1	0.46	0.84	
	18:1n9	44.38	14.76	
621	20:1	1.05	0	
	22:1n9	0	0.26	
622	24:1	0	0	
022	MUFA	51.21	19.37	
623	18:3 n 3	66.43	8.79	
023	18:4 n 3	6.23	0.74	
	20:5 n 3	2.73	0	
624	20:3 n 3	0.77	0.29	
	22:6 n 3	0.95	5.38	
625	n3	77.11	15.20	
020	18:2 n 6	38.15	11.20	
	18:3 n 6	0.32	0.32	
626	20:3 n 6	0	0	
	20:4 n 6	1.54	0	
627	22:2 n 6	0	0	
	n6	40.01	11.52	
629	n3:n6	7.32	1.52	
628	PUFA	119.70	28.92	
	SAFA:	11,0	20.72	
629	(MUFA+PUFA)	3.68	1.06	

611 Appendix 2. Annual biomass inputs of fatty acids from senesced vegetation.