

Consequences of Didymo Blooms in the transnational Kootenay River basin

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Abstract

Stream habitat changes that affect primary consumers often indirectly impact secondary consumers such as fishes. Blooms of the benthic algae Didymosphenia geminata (Didymo) represent one such habitat change known to affect stream macroinvertebrates. However, the potential indirect trophic impacts on fish consumers via modifications to their diet are poorly understood. The overall goal of this project was to determine if Didymo blooms in streams of the Kootenay River basin of British Columbia and Montana affect the condition and growth of fishes. and to see whether trophic mechanisms were responsible for any observed changes. We therefore quantified the diet, condition, and growth rate of trout, charr, and sculpin in a paired, Didymo vs. reference study, during the summer of 2018 and across a gradient of Didymo abundance in 2019. In the 2018 study, trout diets were 81% similar despite obvious differences in the composition of macroinvertebrate assemblages between the Didymo and reference streams. Trout abundance was higher in the stream with Didymo, but the amount of invertebrates in the drift was higher in the stream without Didymo. Growth rate and energy demand by individual trout was similar between the two streams. In the 2019 study, across a gradient of coverage, Didymo abundance was correlated only with the percent of aquatic invertebrates in trout diets and did not affect diets of charr or sculpin. Variation in fish condition was low across study streams. Thus, Didymo blooms may impact trout diets to a small extent, but we found no evidence this impact translates to changes in condition or growth. The relationship of fish abundance to Didymo blooms bears further study, but we found no obvious trophic mechanisms that would explain any differences. We suggest future studies prioritize research on potential impacts during winter months and on species with limited mobility that may be most greatly impacted by Didymo.

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Introduction

Fish growth and production in coldwater systems is highly dependent on both allochthonous and autochthonous sources of energy (Horton 1961; Huryn 1996; Bellmore et al. 2013). In the interior Columbia River basin, a long history of logging, mineral extraction, and river impoundment has altered in-stream habitats and riparian areas (Hand et al. 2018), resulting in a lack of structure and nutrients that alters the availability of food resources to aquatic organisms (Meredith et al. 2014; Minshall et al. 2014). Habitat change can alter stream macroinvertebrate assemblages and impact consumers of both larval and adult life-stages of aquatic insects (Power et al. 1996; Nakano et al. 1999; Baxter et al. 2005; Malison and Baxter 2010). Such changes within the interior Columbia River basin have indeed led to shifts in fish assemblage structure (Frissell 1993). Understanding how specific habitat change alters the flow of in-stream energy sources to fish consumers can thus be of great importance to conservation and management efforts (Cross et al. 2011, 2013; Bellmore et al. 2012; Scholl et al. 2019).

Instream habitat components that alter primary and secondary production such as woody debris and stream substrates are major topics of research, but ephemeral habitat components such as macrophytes and algae are less often considered in restoration and management. In recent years, increasing reports of severe blooms of the diatomaceous algae *Didymosphenia geminata* (hereafter, Didymo) have led to significant concern about its causes and consequences for freshwater organisms (Bickel and Closs 2008; Gillis and Chalifour 2010; James et al. 2010; Anderson et al. 2014; James and Chipps 2016; Jellyman and Harding 2016). Overgrowths (colloquially, blooms) of this North American-native are characterized by the production of a long polysaccharide stalk from individual diatoms, which can lead to large areas of the substrate becoming covered. However, the precise causes of Didymo blooms remain a current topic of investigation (Taylor and Bothwell 2014).

At high Didymo bloom coverage, stream invertebrate assemblages originally dominated by Ephemeroptera, Plecoptera, and Trichoptera (EPT taxa), typically shift towards dominance by Chironomidae, Oligochaeta, Nematoda, or Cladocera, taxa generally associated with reduced habitat quality in trout streams (Kilroy et al. 2009; Gillis and Chalifour 2010; James et al. 2010; Byle 2014; Larned and Kilroy 2014; Jellyman and Harding 2016). There has been widespread concern about the consequences of blooms for trout (Gillis and Chalifour 2010; James et al. 2010; Jellyman and Harding 2016) because EPT taxa are often a primary food source for salmonid species (Behnke 2010). However, to date, it is unclear if Didymo blooms have any significant negative or positive impacts on trout species. Jellyman and Harding (2016) found that blooms in several New Zealand rivers were correlated with lower trout abundances, dietary percent EPT, and stomach fullness. In contrast, production of Atlantic Salmon (Salmo salar) in Icelandic and Norwegian rivers has remained high despite the presence of severe Didymo blooms (Jonsson et al. 2008; Lindstrøm and Skulberg 2008), and spawner abundance and escapement of Pacific salmon and steelhead (Oncorhynchus spp.) in Vancouver Island streams either increased or did not change in relation to blooms (Bothwell et al. 2008). In four South Dakota streams the condition and feeding of large Brown Trout (Salmo trutta) was not correlated with Didymo blooms, while body condition in juveniles was higher (James and Chipps 2010). However, the study was also affected by drought, making causal inference difficult. As such, no individual study has

successfully examined the mechanistic links between Didymo blooms, macroinvertebrates, and fishes necessary to make causal inference. Further, no studies have addressed the potential effects of blooms on inland native trout populations or on nongame species such as members of the family Cottidae.

To better understand the trophic consequences of Didymo blooms, we assessed the relationship between blooms, fish diet, condition, and growth over two summers in a Columbia River subbasin, the mountainous Kootenay (Kootenai in the U.S.) basin of British Columbia, Idaho, and Montana (Fig. 1), much of which falls within the globally-rare, inland temperate rainforest biome (Dellasala et al. 2011). We employed a multi-faceted research approach in which we examined potential Didymo bloom impacts on fish: 1) temporally - in a reference-impact study of two streams during one summer, and 2) spatially – in a survey of fishes across Kootenay basin streams representing a gradient of bloom severity.



Fig. 1. Location of study streams (red dots) within the Kootenay River basin (left) and the upper Libby Creek subbasin (right). Inset A shows the location of the Kootenay basin within the larger Columbia River watershed.

Study Location

To determine the potential effects of Didymo blooms on fishes, we combined a high frequency sampling approach with a high spatial resolution approach. Twice-monthly through the summer of 2018, we sampled two streams located in the Cabinet Mountains of northwestern Montana, Bear Creek and nearby Ramsey Creek (Fig. 1). Both creeks have similar physical characteristics (Table 1), but Bear Creek contains obvious Didymo blooms while Ramsey Creek does not. The two streams thus offer an opportunity to examine potential effects of blooms on biotic communities in a paired, reference-impact framework.

During both the summer of 2018 and 2019, we examined 131 locations on 103 individual streams for the presence of Didymo blooms in the Kootenay River basin (Appendix A). In 2019, we surveyed fishes in 28 of those streams (Fig. 1) representing large differences in bloom coverage: 0 - 80% (Table 2). Ten of those streams were located in British Columbia provincial parks (Fig. 2).



Fig. 2. British Columbia portion of the Kootenay River basin. Sample sites (red dots) are shown in provincial parks (dark grey).

Table 1. Bear and Ramsey Creek habitat measurements - 2018.

2018 Habitat Measurements							
	Bear Ck. (Didymo)	Ramsey Ck. (No Didymo)					
Temp. (°C) ±SD	9.79 ± 2.32	9.79 ± 2.40					
Mesohabitat							
Cascade	76%	83%					
Riffle	16%	10%					
Pool	8%	7%					
Substrate Size	26.7 cm	23.2 cm					
Wetted Width	7.24 m	7.17 m					
Nutrients (µg/L) ±SD							
SRP	1.995 ±0.368	1.530 ±0.409					
Bromide	below detection	below detection					
Fluoride	below detection	below detection					
Nitrate	74.5	25					
Phosphate	below detection	below detection					
Sulfate	1235	930					

	Streams Surveyed in 2019								
Stream	Subbasin (State/Prov.)	% Didymo Coverage	% Overstory Cover	Wetted Width (m)	Dominant Vegetation	Rosgen Channel Type	Large Woody Debris #	Aug. Stream Temp. (°C)	Fish Spp. Observed
	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	DIDYMO	COV	VVVV	VEG	CHAN	LWD	TEMP	
Blacktail Cr.	Yaak R. (MT)	1.7	82.6	3.4	Pine	В	12	10.1	RB
Boulder Cr.	Koocanusa (MT)	33.3	80.2	4.7	Cedar	A	6	12.5	WCT
Bear Cr.	Kootenai R. (MT)	30.6	88.7	8.3	Cedar	В	19	11.9	BULL, RB
Big Cherry Cr.	Kootenai R. (MT)	29.0	74.5	5.9	Cedar	A	75	12.5	BULL, RB, RBxCT, WCT, SLCOT
Burnt Cr.	Yaak R. (MT)	31.6	64.5	7.3	Cedar	Α	12	14.4	EB, LN DC, MWF, RB, SLCOT
Coffee Cr.	Kootenay L. (BC)	2.0		14.3	Cedar	А		12.9	BULL, WCT
Davis Cr.	Kootenay L. (BC)	20.5	60.6	9.8	Cedar	А	5	11.8	BULL, MWF, RB, SLCOT
E. Fork Pipe Cr.	Kootenai R. (MT)	24.6	83.7	3.2	Alder	В	12	10.9	EB, RB, RBxCT, SL COT
E. Fork Yaak R.	Yaak R. (MT)	52.6	31.5	7.6	Pine	В	4	11.4	RB
Granite Cr.	Kootenai R. (MT)	21.3	68.1	10.3	Cedar	В	7	11.6	BULL, EB, RB, SLCOT
Hope Cr.	Lardeau R. (BC)	16.6	67.3	4.8	Cedar	А	16	12.7	BULL, MWF, RB
Kokanee Cr.	Kootenay L. (BC)	0.0	32.0	15.9	Alder	А		13.9	RB
Lake Cr.	Kootenai R. (MT)	27.0	6.3	19.0	Alder	В	0		RB, TCOT
Leigh Cr.	Kootenai R. (MT)	3.2	88.0	4.6	Cedar	А	20	9.9	EB, RBxCT
Lizard Cr.	Elk R. (BC)	15.4	26.0	6.8	Pine	В	5	11.5	EB, RBxCT
Lockhart Cr.	Kootenay L. (BC)	40.6	80.4	5.0	Cedar	А		11.5	BULL, LNDC, MWF, RB
Mobbs Cr.	Lardeau R. (BC)	48.3	28.1	3.2	Alder	Side Channel	8	10.6	BULL, RB, SLCOT
Lost Ledge Cr.	Kootenay L. (BC)	5.7	69.2	4.9	Cedar	А	9	12.8	RB
N. Fork 17 Mile Cr.	Yaak R. (MT)	17.7	66.0	3.8	Cedar	А	15	11.9	EB, RB, SLCOT, WCT
Outlet Cr.	White R. (BC)	80.5	64.0	6.3	Pine	В	8	18.2	RB
Parmenter Cr.	Kootenai R. (MT)	0.9	83.2	7.7	Cedar	В	40	10.7	EB, RB, SLCOT
Pinkham Cr.	Koocanusa (MT)	39.4	53.4	4.5	Alder	А		11.3	EB, RB
Solo Joe Cr.	Yaak R. (MT)	0.0	81.1	2.8	Pine	А	21	11.7	RB
Trail Cr.	Fisher R. (MT)	13.0	63.7	4.9	Pine	В	10	13.9	EB, TCOT, WCT
W. Fisher Cr.	Fisher R. (MT)	1.3	26.9	5.6	Pine	В	2	11.1	BULL, TCOT, WCT
Wolf Cr.	Fisher R. (MT)	0.0	32.0	9.8	Alder	В	3	20.2	LNDC, LSSU, MWF, RSSH, TCOT, WCT
Woodbury Cr.	Kootenay L. (BC)	0.2	37.9	10.1	Cedar	В		8.6	BULL, WCT
Weasel Cr.	Wigwam R. (MT)	52.5	58.0	3.5	Pine	В	5	15.3	RBxCT
	Species Codes:		Brook Trout (E	B), Bull Trout	(BULL), Longn	ose Dace (LNDC), Largescale Such	(er (LSSU), Moun	tain Whitefish (MWF), Rainbow Trout (RB),

Brook Trout (EB), Bull Trout (BULL), Longnose Dace (LNDC), Largescale Sucker (LSSU), Mountain Whitefish (MWF), Rainbow Trout (RB), Rainbow-Cutthroat Hybrid (RBxCT), Redside Shiner (RSSH), Slimy Sculpin (SLCOT), Torrent Sculpin (TCOT), and Westslope Cutthroat Trout (WCT)

Table 2. List of streams surveyed in 2019.

Methods

Didymo vs. Control Stream Study - 2018

We selected a three-hundred meter long reach for study in both Bear and Ramsey Creeks. The fish assemblages of both were predominantly composed of Columbia River Redband Trout (*O. mykiss gairdneri*) and Bull Trout (*Salvelninus confluentus*). Ramsey Creek also contained a small number of Columbia Slimy Sculpin (*Uranidea cognata* syn. *Cottus cognatus*). We measured five habitat variables to ensure Bear and Ramsey Creeks were suitable for comparison: mean substrate size (*sensu* Wolman 1954), channel width, mesohabitat composition (percent cascade, riffle, & pool), water temperature (30-minute recording interval, Onset HOBO© data loggers), and water chemistry (Lachat 8500 QuikChem FIA and IC). Every two weeks, we systematically estimated percent-of-substrate covered by blooming Didymo using a five-gallon bucket with a clear bottom, making five evenly-spaced estimates along lateral transects, each twenty paces apart from reach-top to bottom. We then combined, twice-monthly estimates to form monthly Didymo bloom coverage estimates.

Food-web structure was determined by macroinvertebrate and fish sampling concurrent with Didymo coverage estimation. In conjunction with Didymo bloom measurements, we collected drifting macroinvertebrates by placing two separate 10 x 18 inch drift nets in the stream for 30 minutes and pooling the combined samples in 70% ethanol. Samples were always taken between the hours of 10:00 a.m. and 5:00 p.m. The day following each Didymo and macroinvertebrate sampling event, we collected fishes through single-pass backpack electroshocking (LR-24

Backpack Shocker Smith-Root©, Vancouver, WA). We completed multiple passes during the final sampling event (September) to maximize summer-long recapture. Each fish was anesthetized with clove oil, weighed, measured, and marked by clipping a small section of the caudal fin. We gastrically lavaged individuals larger than 100 mm to collect diets and, if captured during June or July, implanted a uniquely-coded, 12 mm passive integrated transponder (PIT) tag (Model HDX12, Biomark©, Boise, ID). Gut evacuation was assumed to be minimal due to cold temperatures and processing generally less than an hour after capture. Using the average percent growth between individuals measured in June and July, we back-calculated June weights for individuals tagged in July. This represented 57% of Redband Trout in Bear Creek and 68% in Ramsey Creek. For PIT-tagged Redband and Bull Trout captured in September, we also compared summer growth to the total number of times that fish had been captured to test for handling effects.

We identified and measured drift and diet macroinvertebrates to family and used published length-to-mass conversions to estimate biomass (Benke et al. 1999; Sabo et al. 2002; Baumgärtner and Rothhaupt 2003; Gruner 2007; Miyasaka et al. 2008) and caloric content (Montana Fish, Wildlife & Parks, unpublished data). Conversions are provided in Appendix B.

Because a shift to a macroinvertebrate assemblage of smaller and more abundant individuals may favor juvenile fishes (James and Chipps 2016), we identified large and small sizeclasses of Redband and Bull Trout using length-frequency histograms (Appendix C). We then calculated size-specific abundances using Lincoln-Petersen mark-recapture estimation in which the final sampling date was the recapture event and all previous sampling events a single marking event (Lincoln 1930). We determined this approach to be a reasonable because movement of PIT-tagged fishes between the abutting upper and lower halves of Bear Creek was negligible and thus assumed the closed population assumption of Lincoln-Petersen estimation was satisfied (Appendix D).

We compared taxon-specific proportions of drifting macroinvertebrates in Bear and Ramsey Creeks by calculating the monthly percent similarity (Schoener 1970):

(1)

Percent Similarity =
$$100 - 0.5\left(\sum_{i=1}^{n} |B_i - R_i|\right)$$

where B_i is the percent of invertebrates of taxa *i* in Bear Creek and R_i is the percent of invertebrates of taxa *i* in Ramsey Creek. Using the same equation, we compared trout diets to the availability of invertebrates in the drift as a measure of selection. Then, we also compared trout diets between the two streams using percent energetic content for each diet taxa. To evaluate how likely observed differences between groups were (drift vs. drift, diet vs. drift, and diet vs. diet), we used Pearson's chi-squared tests. We further report monthly and summer-long gut fullness and relative number and energetic content of invertebrates in the drift between the two streams.

By pairing individual caloric demand with trout diet composition, we created energy-flow food webs. We used a novel modification of the Benke-Wallace trophic-basis of production method that accounts for thermal preferences to calculate energetic demand (Benke and Wallace 1980). The Benke-Wallace method was originally developed for use with benthic macroinvertebrates and does not account for differential allocation of energy by organism size and water temperature, factors known to strongly influence fish growth (Brown et al. 2004). Thus we used two different numbers for the proportion of total assimilated energy allocated to growth (net production efficiency or NPE) in large vs. small fishes as suggested by Bellmore et al. (2013). We then modified this proportion by observed stream temperatures as compared to species' thermal optimums such that a fish's consumption in kilocalories was calculated

(2)

$$Consumption = \sum_{i=1}^{n} \frac{(DietProportion_{i} \times Growth \times EnergyDensity)}{(TempFactor \times TissueAllocation \times (Digestible_{i} - 0.2Digestible_{i}))}$$

where *DietProportion*_i is the average proportion by kilocalories of food type *i* in the diet,

Growth is the summer growth (Jun.-Sept.) in grams of the average fish,

EnergyDensity is the energy density (kcal/gram) of the fish,

TissueAllocation is the theoretical maximum proportion of assimilated energy allocated to fish tissue growth (net production efficiency), which was set as 0.22 for large size-class trout and 0.5 for small size-class trout and Slimy Sculpin

Digestible; is the estimated digestible proportion of food type i, and

TempFactor is the temperature correction factor calculated according to the equation

(3)

 $TempFactor = e^{-((0.2 \times (StreamTemp - OptimTemp))^4)}$

where *StreamTemp* is the average stream temperature for the measurement interval over which growth was recorded and *OptimTemp* is the thermal optimum for the given species of fish. This equation is an approximation of a fish's thermal optimum curve that asymptotes at an energy-allocation-to-tissue value of zero (Appendix E). We derived thermal optimum values from previous field and laboratory studies: 13.1°C for Redband Trout (Bear et al. 2007), 12.0°C for Bull Trout (Dunham et al. 2004), and 12.1°C for Slimy Sculpin (Wehrly et al. 2004).

We used estimated digestible proportions (*Digestible*_i) for each food type from Hanson et al. (1997) and subtracted a value of $0.2Digestible_i$ to account for specific dynamic action (Hanson et al. 1997). Thus *Digestible*_i – $0.2Digestible_i$ is the assimilation efficiency of food type *i*.

To derive total estimated consumption by each species, we multiplied estimated summer Benke-Wallace consumption values by calculated fish abundances in each stream. Then, we multiplied the proportion of energy of each prey item in the average diet of each fish species by the reach-level consumption estimates. Thus, we obtained estimates of total energy flow from all prey to fish predators and compared results for Bear and Ramsey Creeks (Appendix F).

Multi-Stream Didymo Survey – 2019

In a representative 30.5 meter (100 ft.) reach of each selected stream, we estimated Didymo coverage using the same method as in 2018. We also recorded, six other habitat variables: wetted-width (n=5), canopy density (n=5 using a densitometer [Strickler 1959]), dominant vegetation type, substrate type (Cummins 1962), Rosgen channel type (Rosgen 1994), number of large woody debris items (*sensu* Kershner et al. 2004), and stream temperature. From reach top-to-bottom, we measured wetted width and canopy density, while we qualitatively assessed vegetation, substrate, and channel type. We estimated mean August temperatures by adding the time-specific difference of each stream's temperature to a reference temperature logger (Bear Creek for streams flowing into the Kootenay River below the Fisher River confluence, Outlet Creek for those above the Fisher confluence, and Trail Creek for Fisher River tributaries).

In the same reach, we collected fishes through two-pass (one upstream, one downstream) backpack electroshocking. We anesthetized, weighed and measured all fishes and then released leuciscids and catostomids. Using an in-field assessment in which we gastrically-lavaged fish, we assessed the diets of salmonids and cottids by spreading the diet contents in a 30 x 15 cm white pan and recording the number of individuals of each invertebrate taxa. We identified insects to order except for Simuliidae and Chironomidae, which we identified to family. Other invertebrates we identified to Class or Phylum, and vertebrates to the lowest practical taxonomic level (usually species).

We generated two response metrics of fish condition (Fulton's K [Heincke 1908; Ricker 1975] and residual analysis of observed vs. predicted weights [Fechhelm et al. 1995]) and four metrics of diet composition (%Diptera, %EPT, %Aquatics, and gut fullness [# Diet Items/Fish Length]) for each fish. Using weighted, univariate logistic (%Diptera, %EPT, %Aquatics) and linear regressions (gut fullness and fish condition) in which fish sample size was the relative weight of each stream in the regression, we analyzed each response metric compared to Didymo and the other six habitat variables. We removed four streams (Kokanee, Coffee, Mobbs & Solo Joe Creeks) from regressions due to low sample size or substantially different substrate type. We grouped fish by genus due to otherwise small sample size if compared only within species (charr *Salvelinus* and sculpin *Uranidea*) or significant hybridization in the basin (trout *Oncorhynchus*), which made some field ID's difficult. For each comparison of a habitat variable to a diet metric, we calculated an R² (or Nagelkerke's pseudo-R² for logistic regression [Nagelkerke 1991]) and p-value, and considered variables with an R² greater than 0.2 and a p-value less than 0.2 to be a non-spurious correlation.

Results

Didymo vs. Reference Stream Study - 2018

Differences in all four habitat variables were small between Bear (Didymo) and Ramsey Creeks (No Didymo), giving us confidence the two were suitable for comparison (Table 1). Didymo bloom severity in Bear Creek increased from 10.9% coverage in June to 22.6% coverage in

August before falling to 18.9% in September (Fig. 3). The June to August Didymo growth was significant (p<0.01), but the decline from August to September was not (p = 0.21).



Didymo Coverage & Aquatic Invertebrate Drift

Fig. 3. Monthly, percent of stream substrate covered by Didymo in Bear Creek, 2018 (top). Pie charts show proportions of major aquatic invertebrate taxa in the drift in Bear and Ramsey Creeks.

Percent composition of drifting invertebrates between the two streams generally became less similar as Didymo coverage increased (June-September: 84.2%, 63.1%, 68.5% and 66.6% similar; Fig. 3, Appendix G). Percent EPT in the drift was initially 12.1% higher in Ramsey Creek but by September was 20.3% higher in Bear Creek. However, Ephemeroptera larvae were proportionally more abundant in Ramsey Creek during all months (June-September: 10.5%, 0.7%, 11.8%, and 8.5% higher; Fig. 3). Percent of larval and pupal Diptera in Bear Creek was 17.8% higher than Ramsey Creek in June, 32.1% higher in July, 30.6% higher in August, and 12.0% lower in September. Both total drifting invertebrates and total energy of drifting invertebrates similarly diverged later in the summer with the streams having similar numbers in June, Ramsey Creek having higher numbers in July and August, and Bear Creek having higher numbers in September. The summer-long amount of total energy of drifting invertebrates was 2.2 times higher in Ramsey Creek.

Reach abundance estimates for Redband and Bull Trout were higher in Bear Creek (Table 2). Slimy Sculpin (n=20) were only in Ramsey Creek. Relative growth of Redband Trout varied by size-class. Summer relative growth of small trout (<105 mm) was estimated to be 0.0292 g/g/d in Bear Creek but only 0.0033 g/g/d in Ramsey Creek, but this difference was likely driven by a very small sample size of small Redband Trout during June (3 in Bear Creek and 1 in Ramsey Creek). Relative growth of large size-class Redband Trout (>105 mm) was similar between the two streams: 0.0027±0.0004 g/g/d in Bear Creek and 0.0029±0.0007 g/g/d in Ramsey Creek (mean \pm standard error; Table 3). Growth for similar size class trout was likewise similar between the two

streams (Appendix H). We observed no negative impact of even frequent capture on growth of PIT-tagged fish (Appendix I).

	Stream	Pop Abundance	Individual Growth	Indiv Summer Consumption	Pop Consumption
Redband Trou	t		(g/g/d)	(kcal)	(kcal)
Small	Bear	132	0.0292	20.9	2755.8
	Ramsey	91	0.0033	18.8	1712.5
Large	Bear	196	0.0027	75.3	14724.5
	Ramsey	81	0.0029	54.2	4398.1
Bull Trout					
Small	Bear	60	0.0136	21.3	1277.0
	Ramsey	2	NA	NA	NA
Large	Bear	45	0.0011	23.9	1080.5
	Ramsey	3	NA	NA	NA
Slimy Sculpin					
	Ramsey	20	0.0030	8.8	176.8

Table 3. Population (reach) abundance, growth, and consumption estimates for each fish species and size class in Bear and Ramsey Creeks. Bioenergetics consumption estimates are shown for comparison to Benke-Wallace estimates though population-level estimates used the Benke-Wallace method.

Redband Trout diets were 40.7% similar to the drift in Bear Creek (χ^2 test: p < 0.01) and 40.1% similar to the drift in Ramsey Creek (χ^2 test: p < 0.01; Fig. 4). By energetic content, Redband diets were 81.2% similar between Bear and Ramsey Creeks for the whole summer (χ^2 test: p = 0.84): 55.6% similar in June, 77.5% similar in July, 99.7% similar in August, and 75.0% similar in September (Appendix G). Gut fullness was not significantly different between the two streams in any month. Diets of small individual Redband Trout in Bear Creek had more EPT (78.6%±8.4) than large individuals (46.4%±3.0), while gut fullness and %Diptera were similar.

Benke-Wallace consumption estimates for large, individual Redbands were 39% higher in Bear Creek, while small size-class estimates were similar between the two streams (Table 3). Reach-level energetic demand by all Redband Trout were estimated at 17,500 kcal in Bear Creek and 6,111 kcal in Ramsey Creek (Table 3). The primary sources of energy (>5% of demand) for Redband Trout in Bear Creek were Ephemeroptera (38.0% of energy intake), Hymenoptera (15.1%), Trichoptera (14.4%), Plecoptera (9.5%) and Diptera (7.6%); (Fig. 5). Primary energy sources for Ramsey Creek Redbands were Ephemeroptera (45.8%), Hymenoptera (15.7%), Diptera (9.8%), Trichoptera (9.0%), and Plecoptera (6.3%); (Fig. 5). Primary sources of energy for Bull Trout in Bear Creek were Ephemeroptera (48.0%), Trichoptera (13.1%), Nematoda (7.2%), Plecoptera (6.3%), and Hymenoptera (5.1%); (Fig. 5). We collected only 4 Bull Trout 3 Slimy Sculpin diets in Ramsey Creek, and we did not consider this sufficient to draw conclusions as to average diet compositions.



Fig. 4. Invertebrate taxa in Bear and Ramsey Creek drift (top) and proportion in Redband Trout diets (bottom) by month.

Fig. 5. Energy-flow food web for fishes in Bear and Ramsey Creeks. Line thickness represents proportion of total energy demand by the given fish species met by each invertebrate taxa. Only taxa representing at least 5% of energy demand are shown.



Multi-Stream Didymo Survey – 2019

Between-site variation in Fulton's K was too low to assess possible explanatory variables (coefficients of variation [CV] \leq 0.1; Appendix J). Between-site variation in fish relative condition, calculated as a fish's observed weight compared to its predicted weight, was similarly low for trout and sculpin (CV of 0.12 and 0.04, respectively) and moderately low for charr (CV = 0.28). Despite slightly more variation in charr relative condition between sites, there was no relationship between condition and Didymo coverage (R² = 0.03, p = 0.46).

For all diet metrics across all three fish taxa, percent Didymo cover was only correlated with percent of aquatic invertebrates in *Oncorhynchus* diets (Fig. 6). Canopy cover, LWD, riparian vegetation type, and stream temperature were also correlated with percent aquatic invertebrates in *Oncorhynchus* diets, with LWD having the highest pseudo-R² (Appendix J). In fact, few fish diet metrics were correlated with any habitat variable (Appendix K). However, percent of aquatic invertebrates in vertebrates in trout diets was positively associated with pine vegetation types (Fig. 7).



Fig. 6. Correlations of Didymo coverage to each fish taxa's diet and condition from 2019. Each dot is the average value for fish in a single stream. ρ^2 is Nagelkerke's pseudo- R^2 value. *** indicates a p-value ≤ 0.05 .



Fig. 7. Violin plot of percent aquatic invertebrates in trout diets showing the spread across different riparian vegetation types.

Discussion

During the summers of 2018 and 2019, we examined the response of trout, charr, and sculpin to Didymo blooms over space and time. While Didymo appeared to impact the macroinvertebrate assemblage of Bear Creek, the macroinvertebrate food sources and subsequent growth rates of trout did not appear to be affected. Across a gradient of Didymo bloom coverages in 2019, Didymo was weakly correlated with percent of aquatic invertebrates in trout diets but we observed little variation in condition of trout, charr, & sculpin.

As Didymo bloom coverage in Bear Creek increased to its maximum in August 2018, the proportion of the invertebrate drift made up by larval Diptera (primarily Simuliidae and Chironomidae) diverged between the two streams, remaining relatively high in Bear Creek while decreasing in Ramsey Creek. Numerous other studies have similarly found high proportions of Diptera, especially Chironomid larvae, where Didymo is in bloom (Marshall 2007; Kilroy et al. 2009; Gillis and Chalifour 2010; Anderson et al. 2014; Ladrera et al. 2015; Sanmiguel et al. 2016). Yet despite their relative abundance in Bear Creek, Diptera comprised a disproportionately small percent of Redband Trout diets in both streams, indicating strong negative selection. Ephemeroptera, Hymenoptera and Nematoda were strongly selected for by Redband Trout in both streams. Overall, Redband Trout diets were highly similar between the Didymo and reference streams in 2018 (81.2% similar) despite differences in the availability of certain prev taxa. In fact, diets were most similar in August (99.7% similar), when Didymo coverage was at its peak. Correspondingly, major energy sources and growth rates of trout did not differ greatly between Bear and Ramsey Creeks. It is however possible that Didymo coverage in Bear Creek was not severe enough to cause the proportional shifts in macroinvertebrate composition such that trout would have been impacted by food limitation. While a no-Didymo comparison was not available for Bull Trout since so few were captured in Ramsey Creek, Bull Trout in Bear Creek also did not utilize larval Diptera as a major energy source, which may be consequential only at very high Didymo coverage. Further, sexually mature Bull Trout in these systems were likely allocating energy to pre-spawn gamete production, which may have affected overall growth.

Stream resident trout are considered generalist invertivores (Behnke 1992), but strong selection by Redband Trout in both Bear and Ramsey Creeks in 2018 for the same taxa indicates this subspecies may show strong preferences for mayflies (Ephemeroptera). However, given interior (non-steelhead) Redband Trout occupy only 42% of their historic range across the West and only 2% of historic range in Montana (Muhlfeld et al. 2015), it is important to carefully evaluate land management actions such as timber harvest or road construction that may impact sensitive mayfly species.

In our 2019 survey of 28 streams with varying levels of coverage, Didymo bloom severity was not correlated with most measures of fish diet and was only a weak predictor of aquatic invertebrates in trout diets. In conjunction with the 81.2% similarity of diets between Bear and Ramsey Creeks in 2018, this suggests Didymo may alter the composition of trout dietary macroinvertebrates to a small extent, but that shift does not alter condition or growth rates of trout. This disconnect may indicate trout in Kootenay River headwaters are not food limited during summer months, or that much greater diet perturbations are necessary to affect trout growth. Alternatively, the lack of variability in fish condition across streams may suggest fishes in these populations conform to the theory of ideal-free distribution (Fretwell 1969; Sutherland et al. 1988) such that fish condition between streams is relatively homogenous but abundances vary based on where forage is most available. As such, distribution of fish condition in Kootenay basin headwaters may be relatively stable - i.e. exist in a state of equilibrium (*sensu* Nash 1951).

Although not the impetus of our study, we observed interesting differences in percent of aquatic invertebrates in trout diets in streams with differing riparian vegetation (Fig. 7). Allan et al. (2003) found riparian communities dominated by alder in Alaska coastal temperate rainforests, provided more terrestrial invertebrates to juvenile salmon (*Oncorhynchus kisutch*) than did those dominated by a mix of hemlock and spruce (*Picea spp.*). Similarly, we observed higher proportions of riparian invertebrates in trout diets in alder-dominated streams than in pine-dominated streams (largely lodgepole pine *Pinus contorta*). In contrast to the finding of Allan et al. (2003), trout in our inland temperate rainforest streams with riparian communities dominated by cedar and hemlock, had similar aquatic-terrestrial ratios to alder-dominated streams (Fig. 7).

Our study examined the impacts of Didymo blooms only into early Fall during both years, a time when terrestrial invertebrate inputs, and trout reliance upon them, are high (Nakano and Murakami 2001). It is possible terrestrial inputs act as a buffer to shifts in aquatic invertebrate composition caused by blooms and some negative or positive consequence of Didymo can only be observed by studying fishes across seasons. In fact, some studies have reported severe Didymo blooms during winter months (e.g. Kolmakov et al. 2008), and we observed severe blooms in the Lardeau River during April of 2018 when snowpack was still high. Trout growth in headwater streams is higher in summer months but foraging (Thurow 1997) and growth (Al-Chokhachy et al. 2019) still occur over winter. We therefore suggest potential impacts of Didymo on fishes be examined during winter. Further, due to the multitude of studies indicating impacts

to macroinvertebrate assemblages, the relationship of Didymo to imperiled invertebrates, especially sedentary taxa that may not be able to avoid Didymo blooms such as freshwater mussels, bears further study.

Implications for Management

Authors of previous studies have suggested nutrient amendments (James et al. 2015; Coyle 2016) and dam releases (Cullis et al. 2015) as viable means to manage nuisance Didymo blooms. Indeed, both methods show promise for reduction of blooms at local scales. The impetus for this bloom reduction may be independent of concern for fishes, including aesthetics, fouling of infrastructure, or to prevent hypoxia. However, we did not observe any major impacts of Didymo blooms on the diet, condition, or growth of trout in Kootenay basin headwaters. This overall result is similar to those for Brown Trout in a South Dakota stream (James and Chipps 2016). Therefore, it is not clear efforts to control Didymo blooms in headwater streams will benefit fish condition.

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Appendix 1. All 131 unique stream locations examined for presence of Didymo blooms in the Kootenai basin. Streams which we quantitatively assessed coverage are listed as 'Y'. We visually estimated covered for streams listed as 'N'.

	All I	ocations Exami	ned for Did	ymo Blooms		
Date	Stream Name	State/ Province	Latitude	Longitude	% Didymo	Measured (Y) or Estimated (N)
2019-Aug	Alexander Creek	MT	48.39266	-115.32864	0%	N
2019-Aug	Barron Creek	MI	48.51344	-115.30235	0%	N
2018-Aug	Bear Creek	MT	40.17997	-115.573162	10.9%	ř V
2010-Aug 2019-Aug	Bear Creek	MT	48 16886	-115 58853	30.6%	Y
2018-Aug	Big Cherry Creek	MT	48.32744	-115.529159	0.070	Ň
2018-Aug	Big Cherry Creek	MT	48.35288	-115.525966	Õ	N
2019-Aug	Big Cherry Creek	MT	48.26734	-115.53004	0%	Ν
2019-Aug	Big Creek	MT	48.82052	-115.35209	0%	Ν
2019-Aug	Big Foot Creek	MT	48.58376	-115.70378	0%	N
2019-Jun	Big Sand Creek	BC	49.37202	-115.240065	0%	N
2019-Aug	Blacktail Creek	MT	48.95124	-115.54154	5-10%	N
2019-Aug	Boulder Creek	MT	48.82052	-115.29097	33.3%	Ý
2018-Aug	Bramlet Creek Bristow Creek		48.03754	-115.49219	0%	N N
2019-Aug 2019-Aug	Bunker Hill Creek	MT	40.04427	-115.29245	0%	N
2019-Aug	Burnt Creek	MT	48 72936	-115 87002	31.6%	Y
2018-Jun	Cable Creek	MT	48.16965	-115.585025	0%	Ň
2019-Aug	Callahan Creek	MT	48.45636	-115.89153	0%	Ν
2019-Jun	Canyon Creek	MT	48.43051	-115.283832	0%	Ν
2019-Aug	Cascade Creek	BC	50.39822	-117.09391	<5%	Ν
2018-Aug	Cedar Creek	MT	48.4093 <mark>1</mark>	-115.665338	0	N
2018-Aug	Cedar Creek	MT	48.43052	-115.628991	<5%	N
2019-Aug	Cedar Creek	MT	48.4096	-115.66522	<5%	N
2019-Aug	Cedar Creek (Kootenay L.)	BC	49.74661	-116.91241	0%	N
2019-Aug	Coffee Creek	BC	49.69663	-116.91/81	2%	Y
2019-Aug	Cooper Creek	BC	50.19901	-116.96092	<5%	N
2019-Jun 2019 Aug	Davis Crook	INT I	40.47742	-115.20000	20.5%	N V
2019-Aug	Deen Creek	MT	48 26923	-115 53397	20.5%	N
2019-Aug	Deer Creek	ID	48.81707	-116.11524	0%	N
2019-Aug	Duhamel Creek	BC	49.57474	-117.22723	0%	N
2019-Aug	Duncan River	BC	50.19634	-116.95234	0%	Ν
2019-Jun	Dunn Creek	MT	48.3851	-115.313085	0%	Ν
2019-Aug	East Fork Pipe Creek	MT	48.61675	-115.61885	24.6%	Y
2019-Aug	East Fork Yaak River	MT	48.94885	-115.53378	52.6%	Y
2018-Aug	Elliot Creek	MT	48.23637	-115.46749	0%	N
2018-Aug	Fisher River	MI	48.06978	-115.374032	5%	N
2019-Jun	Five Mile Creek		48.53573	-115.210572	0%	N
2019-Aug	Flattall Creek		40.03313	-115.70071	0%	N
2018-Aug	Flower Creek	MT	48 34452	-115 606768	0	N
2018-Aug	Flower Creek	MT	48 39004	-115 560911	0%	N
2018-Aug	Granite Creek	MT	48.30597	-115.543096	0	N
2019-Aug	Granite Creek	MT	48.30613	-115.54307	<5%	Ν
2019-Aug	Granite Creek	MT	48.29544	-115.62011	21.3%	Y
2019-Aug	Hammill Creek	BC	50.20204	-116.94646	5-10%	N
2019-Aug	Hellroaring Creek	MT	48.78759	-115.91856	0%	N
2019-Aug	Hope Creek	BC	50.45751	-117.19081	16.6%	Y
2018-Aug	Horse Creek	MT	48.30163	-115.598662	0	N
2019-Aug 2019-Aug	Hudson Creek	MT	40.30100	-115.09000	0%	N
2019-Aug	Jackson Creek	MT	48 46479	-115 31589	0% 0%	N
2019-Aug	Kaslo River	BC	49,9085	-116.90636	0%	N
2019-Aua	Kokanee Creek	BC	49.60506	-117.12635	0%	Y
2018-Aua	Lake Creek (Fisher R.)	MT	48.03812	-115.482052	0	Ň
2018-Aug	Lake Creek (Fisher R.)	MT	48.03766	-115.490971	0	Ν
2018-Aug	Lake Creek (Fisher R.)	MT	48.03915	-115.471111	0%	Ν
2019- Aug	Lake Creek (Kootenai R.)	MT	48.44915	-115.879329	27%	Y
2019-Aug	Leigh Creek	MT	48.22127	-115.60603	3.2%	Y
2018-Aug	Libby Creek	MT	48.31436	-115.537251	0	N
2018-Aug	Libby Creek	MT	48.22552	-115.477984	0	N

Appendix 1 (cont.)

2018-Aug	Libby Creek	MT	48.22456	-115.477006	0	Ν
2018-Aug	Libby Creek	MT	48.3 1 436	-115.504728	0%	Ν
2019-Jun	Little Sand Creek	BC	49.37789	-115.29201	5-10%	N
2019-Sep	Lizard Creek	BC	49.48972	-115.104813	15.4%	Y
2019-Aug	Lockhart Creek	BC	49.5084	-116.785994	40.6%	Y
2019-Aug	Lost Ledge Creek	BC	49.90834	-116.90733	5.68%	Y
2019-Aug	McKillop Creek	MT	48.148	-115.27532	0%	N
2019-Aug	Meadow Creek	ID	48.81943	-116.1462	0%	N
2019-Aug	Meadow Creek (Lardeau R.)	BC	50.22831	-116.98587	0%	N
2019-Aug	Meadow Creek (Yaak R.)	MI	48.78425	-115.92321	20-30%	N
2019-Aug	Movie Diver	BC	00.00073	-117.27104	48.3%	ř N
2019-Aug	North Fork 17 Mile Crook		40.02030	-110.1402	30% 17.7%	N
2019-Aug	North Fork Parsnin Creek	MT	48.00022	115 3/280	0%	N
2019-Aug	North Fork Vaak River (Vaak R.)	MT	40.07.094	-115 61128	0%	N
2019-Sen	Outlet Creek	BC	50 16812	-115 464053	80.5%	Y
2018-Aug	Parmenter Creek	MT	48 39906	-115 57565	0.00	N
2018-Aug	Parmenter Creek	MT	48.37799	-115.628871	0	N
2018-Aug	Parmenter Creek	MT	48.39272	-115.579392	0%	N
2019-Aug	Parmenter Creek	MT	48.37814	-115.62908	0.9%	Ŷ
2019-Jun	Pinkham Creek	MT	48.82799	-115.24295	39.4%	Y
2019-Aug	Pinkham Creek	MT	48.82799	-115.24295	39.4%	Y
2019-Aug	Pipe Creek	MT	48.4278	-115.59636	<5%	Ν
2019-Aug	Pipe Creek	MT	48.61675	-115.61885	0%	Ν
2019-Aug	Placer Creek	ID	48.82678	-116.13982	0%	Ν
2019-Aug	Pleasant Valley Fisher River	MT	48.04044	-115.29337	5-10%	N
2019-Aug	Poplar Creek	BC	50.4 1 548	-117.12218	5-10%	N
2018-Aug	Prospect Creek	MT	48.31877	-115.575627	0%	N
2019-Aug	Quartz Creek	MT	48.60519	-115.6947	20-30%	N
2018-Jun	Ramsey Creek	MT	48.13918	-115.536531	0%	Y
2019-Aug	Rapid Creek	BC	50.43936	-117.15664	<5%	N
2019-Aug	Raven Creek	MI	48.04519	-115.28/15	0%	N
2019-Aug	Red Top Creek	MI	48.76113	-115.91/94	0%	N
2019-Aug	Redfish Creek	BC	49.65643	-117.04791	0%	N
2019-Aug	ROSS Creek	BC MT	49.00043	-110.93313	0%	IN N
2010-Aug	Shoop Crook		40.30300	-115.594735	0%	IN N
2019-Jun 2019-Aug	Sitkum Creek	BC	40.02200	-117 17/27	0%	N
2019-Aug 2019-Aug	Skin Creek		49.39300	-116 12033	0%	N
2019-Aug	Snell Creek	MT	48 19548	-115 27532	0%	N
2019-Aug	Snowshoe Creek	MT	48 2138	-115 59783	0%	N
2019-Aug	Solo Joe Creek	MT	48.92425	-115.53963	0%	Y
2018-Aug	South Fork Flower Creek	MT	48.34277	-115.602578	0%	N
2019-Aug	South Fork Miller Creek	MT	48.03825	-115.4603	<5%	Ν
2019-Aug	Spread Creek	MT	48.82335	-115.85119	5-10%	Ν
2019-Jun	Staples Creek	BC	49.917	-115.648183	0%	Ν
2019-Aug	Sullivan Creek	MT	48.87102	-115.25779	0%	Ν
2019-Jun	Sutton Creek	MT	48.76062	-115.280701	0%	N
2018-Aug	Swamp Creek	MT	48.17664	-115.448145	0%	N
2019-Jun	Ten Mile Creek	MT	48.595	-115.206589	0%	N
2019-Aug	Tenderfoot Creek	BC	50.48269	-117.21991	<5%	N
2018-Aug	Trail Creek	MT	48.03856	-115.460284	0	N
2019-Aug	Trail Creek	MT	48.03825	-115.4603	13%	Y
2019-Aug	Unnamed Tributary	MI	48.24024	-115.56314	0%	N
2019-Aug	Unnamed Tributary (Quartz Cr.)	MI	48.57516	-115.69742	0%	N
2019-Aug			48.20577	-115.59134	29%	Y
2019-Aug	Vinal Creek		48.86076	-115.04443	0%	IN N
2019-Juli 2010 Son	Wahahu Creek		40.00001	-113.211244	52 5%	N V
2019-Sep	West Eisher Creek	MT	48.94904	-115/28004	JZ.J /0	N
2018-Aug	West Fisher Creek	MT	48 04227	-115 472095	~570	N
2018-Aug	West Fisher Creek	MT	48 05628	-115 40030	<5%	N
2019-Aug	West Fisher Creek	MT	48.04253	-115 47351	1.3%	Y
2019-Aug	West Fork Creek	MT	48.42768	-115.60329	0%	N
2019-Aua	West Fork Yaak River	MT	48,934	-115.67286	0%	N
2019-Aug	Wolf Creek	MT	48.23393	-115.28529	0%	Y
2019-Aug	Woodbury Creek	BC	49.80625	-117.028353	0.16%	Y
2019-Aug	Yaak River	MT	48.78713	-115.91965	20-30%	Ν
2019-Aug	Young Creek	MT	48.87102	-115.21744	0%	Ν





Bear Creek Redband Trout Length-Frequency

Ramsey Creek Redband Trout Length-Frequency



Length (mm)

		Prey	/ Mass & I	Energy Conversions		
		Lengt	h to Mass		kilooolorioo nor	Based
Taxa	Stage	Coe	fficients	Source	kilocalories per	Proportion
	3-	h	3		gram dry mass	Indigestible
Incosto	Lonyoo	2 700	0.0064	Banka at al. 1990	E 11E210E22	0.1
Distora	Larvae	2.700	0.0004	Benke et al. 1999	J. 115510555	0.1
Athorioidao	Larvae	2.092	0.0025	Benke et al. 1999	4.045075520	0.1
Riophariaidaa	Larvae	2.000	0.004	Benke et al. 1999	4.270001500	0.1
Corotonogonidoo	Larvae	2.460	0.0007	Benke et al. 1999	4.270001300	0.1
Ceratopogonidae	Larvae	2.409	0.0025	Benke et al. 1999	4.742017400	0.1
Simuliidaa	Larvae	2.017	0.0018	Benke et al. 1999	3.427029020	0.1
Simulidae	Larvae	0.011	0.002	Benke et al. 1999	4.290047555	0.1
Hemintere	Larvae	2.001	0.0029	Benke et al. 1999	4.270001300	0.1
Cerividee	Larvae	2.7.34	0.0100	Benke et al. 1999	5.457649904	0.1
Corridae	Larvae	2.904	0.0031	Benke et al. 1999	4.020227029	0.1
Veliidee	Larvae	2.090	0.015	Benke et al. 1999	5.457649904	0.1
Vellidae	Larvae	2.719	0.0126	Benke et al. 1999	3.437649904	0.1
Conniideo	Larvae	2.704	0.0094	Benke et al. 1999	4.020227029	0.1
Capilidae	Larvae	2.002	0.0049	Benke et al. 1999	4.020227029	0.1
Nemeuridee	Larvae	2.724	0.0065	Benke et al. 1999	4.020227029	0.1
Nemoundae	Larvae	2.702	0.0056	Benke et al. 1999	5.457649904	0.1
Perildae	Larvae	2.0/9	0.0099	Benke et al. 1999	4.626227629	0.1
Periodidae	Larvae	2.742	0.0196	Benke et al. 1999	4.828227629	0.1
Pteronarcyldae	Larvae	2.573	0.0324	Benke et al. 1999	4.828227829	0.1
Odonata	Larvae	2.792	0.0078	Benke et al. 1999	4.828227629	0.1
Epnemeroptera	Larvae	2.832	0.0071	Benke et al. 1999	5.472659943	0.1
Ameletidae	Larvae	2.588	0.0077	Benke et al. 2000	5.472659943	0.1
Baetidae	Larvae	2.8/5	0.0053	Benke et al. 1999	5.472659943	0.1
Baetiscidae	Larvae	2.905	0.0116	Benke et al. 1999	5.472659943	0.1
Caenidae	Larvae	2.172	0.0054	Benke et al. 1999	5.472659943	0.1
Epnemereilidae	Larvae	2.676	0.0103	Benke et al. 1999	5.472659943	0.1
Heptageniidae	Larvae	2.754	0.0108	Benke et al. 1999	5.589738241	0.1
Siphionuridae	Larvae	3.446	0.0027	Benke et al. 1999	5.472659943	0.1
Trichoptera	Larvae	2.839	0.0056	Benke et al. 1999	5.002345411	0.1
Brachycentridae	Larvae	2.818	0.0083	Benke et al. 1999	5.002345411	0.1
Glossosomatidae	Larvae	2.958	0.0082	Benke et al. 1999	5.002345411	0.1
Hydropsychidae	Larvae	2.926	0.0046	Benke et al. 1999	5.389604398	0.1
Lepidostomatidae	Larvae	2.649	0.0079	Benke et al. 1999	5.002345411	0.1
Leptoceridae	Larvae	3.212	0.0034	Benke et al. 1999	5.002345411	0.1
Limnephilidae	Larvae	2.933	0.004	Benke et al. 1999	4.015080424	0.1
Polycentropodidae	Larvae	2.705	0.0047	Benke et al. 1999	5.002345411	0.1
Psychomylidae	Larvae	2.873	0.0039	Benke et al. 1999	5.002345411	0.1
Rhyacophilidae	Larvae	2.48	0.0099	Benke et al. 1999	5.002345411	0.1
Coleoptera	Larvae	2.91	0.0077	Benke et al. 1999	5.5597 18164	0.1
Eimidae	Larvae	2.8/9	0.0074	Benke et al. 1999	4.828227829	0.1
Amphipoda	Larvae	3.015	0.0058	Benke et al. 1999	4.004678203	0.1
Decapoda	Larvae	3.620	0.0147	Benke et al. 1999	3.940039300	0.1
Hydrachnidia	Larvae	1.66	0.1327	Baumgartner & Rothhaupt 2003	4.828228967	0.1
Negociaeta		1.000	0.000		4.003450122	0.1
Nematoda Enhemerontera	Adult	2/0	0.00983607	Clancy, unpublished data Sabo et al. 2002	4.716154015	0.1
	Adult	2.49	0.014	Sabo et al. 2002	5.457649904	0.1
Plecontera	Adult	1 60	0.012	Sabo et al. 2002	5 457649904	0.1
Hemintera	Adult	3 3 3	0.20	Sabo et al. 2002	5 457649904	0.1
Trichontera	Adult	20	0.005	Sabo et al. 2003	5 457649904	0.1
Coleoptera	Adult	2.0	0.01	Sabo et al. 2002	5 550718164	0.1
Hymenontera	Adult	1.56	0.64	Sabo et al. 2002	4 632007801	0.1
Orthontera	Adult	2.55	0.00	Sabo et al. 2002	5 08030761	0.1
Aranoao	Adult	2.55	0.05	Sabo et al. 2002	4 828228067	0.1
Dintera		2.14	0.05	Sabo et al. 2002 Sabo et al. 2002	5 786870076	0.1
Culicidae	Adult	2.20	0.04 0.030	Sabo et al. 2002	5 786870076	0.1
Formicidae	Adult	2.000	0.032	Sabo et al. 2002 Sabo et al. 2002	4 552011264	0.1
Parlidaa	Adult	2.000	0.027	Sabo at al. 2002	5 457640004	0.1
Stanhylinidae		2.019	0.000	Sabo et al. 2002 Sabo et al. 2002	5.457049904	0.1
Veenidee	Adult	3 700	0.001	Sabo at al. 2002	1 622007201	0.1
Lenidontera	Adult	J.120 2 210	0.001	Gruper 2002	4.032097001 5 15765	0.1
Collembola	Adult	2.010	0.0179	Gruner 2003 Gruner 2003	5 457640004	0.1
Insecta		2.009	0.0000	Calculated Average	5 361210400	0.1
A stinger to to the	Adun	2.000	0.0000000		5.501210409	0.1
Actinopterygii		2.935	0.00000307	Gancy, unpublished data	5.089403633	0.033

Appendix 3. Biomass and energy conversions for fish prey items. Length (in millimeters)-tomass (grams_{DryMass}) conversions follow the equation: $Mass = a*Length^b$

Fish Movement Between Bear Creek Sections						
	Percent of Tagged Fish					
Month	Recaptured in Adjacent Section	Recaptured in Same Section				
July	3.4%	96.6%				
August	22.4%	77.6%				
September	8.3%	91.7%				

Appendix 4. Movement of trout between the abutting lower and upper halves of Bear Creek, MT.

Appendix 5. Example thermal adjustment curve for the modified Benke-Wallace method for a fish with a 13.1°C thermal optimum.



2018 Redband Trout Energy Sources					
	% Total Energ	gy Demand			
Prey Source	Bear Cr.	Ramsey Cr.			
Actinopterygii	0.2	0.0			
Arachnida	0.8	0.1			
Coleoptera	4.8	3.7			
Collembola	<0.1	<0.1			
Diptera Adult	3.0	3.0			
Diptera Larvae	4.6	6.8			
Ephemeroptera Adult	5.3	2.0			
Ephemeroptera Larvae	32.7	43.8			
Hemiptera Adult	0.6	0.1			
Hymenoptera	15.1	15.7			
Lepidoptera	2.1	1.2			
Nematoda	2.5	7.6			
Oligochaeta	1.7	0.6			
Plecoptera Adult	2.4	0.6			
Plecoptera Larvae	7.1	5.7			
Trichoptera Adult	0.4	0.2			
Trichoptera Larvae	14.0	8.8			
Other Insecta Adult	2.7	0.2			

Appendix 6. Average percent-of-energy derived from different prey sources by Redband Trout in Bear and Ramsey Creeks during the summer of 2018.

Results of Pearson's Chi-squared tests							
Bear Cr. Drift vs. Ramsey Cr. Drift							
	χ ²	df	p-value				
June	5.9	3	0.11				
July	6.4	3	0.09				
August	22.5	3	5.2x10⁻⁵				
September	3.2	3	0.37				
Full Summer	5.0	3	0.17				
Bear Cr. R	edband Diets	vs. Bear	Cr. Drift				
	χ ²	df	p-value				
Full Summer	79.1	19	2.7x10 ⁻⁹				
Ramsey Cr. D	rift vs. Ramsey	y Cr. Red	lband Diets				
	χ ²	df	p-value				
Full Summer	82.9	16	4.9x10 ⁻¹¹				
Bear Cr. Red	band Diets vs.	. Ramsey	/ Cr. Diets				
	X ²	df	p-value				
June	53.4	12	3.5x10 ⁻⁷				
July	16.9	15	0.32				
August	18.2	15	0.25				
September	26.5	13	0.01				
Full Summer	12.2	18	0.84				

Appendix 7. Pearson's Chi-squared test results comparing macroinvertebrate drift between Bear and Ramsey Creeks, Redband Trout diets to drift in each stream, and diets between the streams. χ^2 is the chi-squared test statistic and df is degrees of freedom.

Appendix 8. Redband Trout length (at first capture) compared to its summer long growth. Bear Creek (Didymo) is in red and Ramsey Creek (No Didymo) is in blue.



Appendix 9. Relationship of handling pressure and growth of trout during summer 2018 in Bear Creek.



Handling Pressure vs. Trout Summer Growth

Dispersion Statistics for Condition & Gut Fullness							
		Mean	Coefficient of Variation				
Trout	К	0.943	0.0612				
	Fullness	0.0878	0.471				
Charr	K	0.901	0.0764				
	Fullness	0.0756	0.447				
Sculpin	K	1.08	0.112				
	Fullness	0.0494	0.677				

Appendix 10. Statistics of spread for trout, charr, and sculpin condition (K) and gut fullness across the 24 streams included in analyses of 2019 data.

Appendix 11. Univariate linear regression results for the five continuous and two categorical habitat variables on trout, charr, and sculpin diet metrics. Categorical variables were assessed with an anova and post-hoc Tukey test. COV is canopy cover, WW is wetted width, LWD is large woody debris, TEMP is average August stream temperature, VEG is riparian vegetation type, and CHAN is Rosgen channel type.

Habitat Predictors of Fish Diet								
			Variable Diffe	rence?				
	DIDYMO	COV	WW	LWD	TEMP	VEG	CHAN	
Oncorhynchus	spp.							
<u>% Midges</u>	0.15	0.26***	0.08	0.53	0.00	No	No	
<u>% EPT</u>	0.00	0.00	0.03	0.51	0.05	No	No	
<u>% Aquatics</u>	0.45***	0.46***	0.00	0.53**	0.44***	Yes _{Pine-Cedar} ***	No	
<u>Fullness</u>	-0.01	0.00	0.01	0.27	0.02	No	No	
Salvelinus spp.								
<u>% Midges</u>	-0.02	0.04	0.01	0.38*	0.0	No	No	
<u>% EPT</u>	0.00	-0.01	0.28**	0.33	0.03	No	No	
<u>% Aquatics</u>	0.00	0.05	0.32*	0.36	0.02	No	No	
<u>Fullness</u>	0.02	0.02	0.10	0.15	0.00	No	No	
Uranidea spp.								
<u>% Midges</u>	0.22	0.00	0.03	0.00	0.02	No	No	
<u>% EPT</u>	0.13	0.02	0.00	0.00	0.00	No	No	
<u>% Aquatics</u>	-0.01	0.12	0.44	0.00	0.06	No	Yes*	
<u>Fullness</u>	-0.04	0.00	0.03	0.00	-0.02	No	No	

* indicates p≤0.2, ** indicates p≤0.1, *** indicates p≤0.05