

1 **Digestive enzyme activities in the guts of bonnethead sharks (*Sphyrna tiburo*)**
2 **provide insight into their digestive strategy and evidence for microbial**
3 **digestion in their hindguts**

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27 **Abstract**

28 Few investigations have studied digestive enzyme activities in the alimentary tracts of sharks to
29 gain insight into how these organisms digest their meals. In this study, we examined the activity
30 levels of proteases, carbohydrases, and lipase in the pancreas, and along the anterior intestine,
31 spiral intestine, and colon of the bonnethead shark, *Sphyrna tiburo*. We then interpreted our data
32 in the context of a rate-yield continuum to discern this shark's digestive strategy. Our data show
33 anticipated decreasing patterns in the activities of pancreatic enzymes moving posteriorly along
34 the gut, but also show mid-spiral intestine peaks in aminopeptidase and lipase activities, which
35 support the spiral intestine as the main site of absorption in bonnetheads. Interestingly, we
36 observed spikes in the activity levels of N-acetyl- β -D-glucosaminidase and β -glucosidase in the
37 bonnethead colon, and these chitin and cellulose, respectively, degrading enzymes are likely of
38 microbial origin in this distal gut region. Taken in the context of intake and relatively long
39 transit times of food through the gut, the colonic spikes in N-acetyl- β -D-glucosaminidase and β -
40 glucosidase activities support the contention that bonnetheads take a yield-maximizing strategy
41 to the digestive process, with some reliance on microbial digestion in their hindguts. This is one
42 of the first studies to examine digestive enzyme activities along the gut of any shark, and
43 importantly, the data match with previous observations that sharks take an extended time to
44 digest their meals (consistent with a yield-maximizing digestive strategy), and that the spiral
45 intestine is the primary site of absorption in sharks.

46 Key words: elasmobranch, trypsin, lipase, β -glucosidase, maltase, chitin, spiral intestine,
47 pancreas

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49 **1. Introduction**

50 Generally, the digestive strategies of animals fit within a spectrum of physiological
51 parameters called a rate-yield continuum (Fig. 1). On one end, yield-maximizers consume
52 relatively large meals less frequently, hold the digesta in their digestive tract for long periods of
53 time, and have relatively high digestive efficiency of total organic matter. Rate-maximizers, on
54 the other hand, tend to consume large amounts of low-quality food at a high frequency, pass food
55 through the gut quickly, and have relatively low digestibility of total organic matter. Rate-
56 maximizers tend to readily digest the more soluble fractions of their diet (German 2009; German
57 and Bittong, 2009; Karasov and Douglas, 2013; German et al., 2015), whereas yield maximizers
58 can also digest the more structural elements (e.g., chitin) of their food, either endogenously, or
59 with the aid of an enteric microbial community (Crossman et al., 2005; Skea et al., 2005;
60 Karasov and Martinez del Rio 2007; Karasov and Douglas, 2013). Hence, in addition to diet
61 itself, an animal's digestive strategy affects its ecological role, and thus, it is important to move
62 beyond diet analysis in studies of trophic ecology and also investigate an animal's nutritional
63 physiology (Crossman et al., 2005; Skea et al., 2005, 2007; Karasov and Martinez del Rio 2007;
64 Karasov and Douglas, 2013; German et al., 2015).

65 Most fishes, including sharks, do not masticate their food before ingesting it into the
66 digestive tract. Thus, the chemical means of digestion (i.e., hydrochloric acid, digestive
67 enzymes) are crucial in nutrient acquisition in fishes (Papastamatiou and Lowe 2004, 2005;
68 Clements and Raubenheimer, 2006; German 2011). In fact, the activity levels of digestive
69 enzymes are often used to infer digestive function in fishes. Generally, carbohydrase activities
70 (e.g., amylase) are elevated in herbivores, whereas some proteases (e.g., aminopeptidase) can be
71 elevated in carnivores (Hidalgo et al., 1999; German et al., 2004; German et al., 2015).

72 Moreover, the patterns of enzymatic activity along a fish's gut can provide insight into the
73 digestive strategy taken by a fish to digest a given diet (Fig. 2A; Skea et al., 2005, 2007; Day et
74 al., 2011; German, 2009; German and Bittong, 2009; German et al., 2015). Key to these studies
75 is a spike in the activities of microbially-derived digestive enzymes—especially enzymes that
76 degrade insoluble, structural compounds like cellulose, carrageenan, and chitin—in the hindguts
77 of fishes adopting a yield-maximizing strategy, and a lack of such an activity spike in rate-
78 maximizing fishes. The reason for the distal intestine spike in microbial enzymatic activity in
79 fishes taking a yield maximizing strategy is that yield maximizers tend to have rich microbial
80 communities in their hindguts (i.e., foregut microbial digestion, as in ruminant mammals, is
81 unknown in fishes; Moran et al., 2005; Clements and Raubenheimer, 2006; Clements et al.,
82 2014). Thus, patterns of digestive enzymatic activity along a fish's gut are useful in
83 understanding their digestive strategy and trophic ecology.

84 As of late, there has been an increase in the interest in shark trophic ecology, and yet,
85 investigations of digestive strategies in sharks are limited and have primarily focused on the
86 stomach (e.g., Papastamatiou and Lowe, 2004, 2005; Papastamatiou 2007). Thus, in this study,
87 we investigated the bonnethead shark, *Sphyrna tiburo*, which is a small, coastal hammerhead
88 species and is one of the most abundant elasmobranch taxa in coastal Florida waters.
89 Bonnethead sharks generally consume crustaceans, cephalopods, and mollusks (Cortés et al.,
90 1996), although some young-of-the-year bonnetheads consume copious amounts of seagrass
91 (Bethea et al., 2007). Hence, while having a typical carnivorous diet, bonnetheads may also need
92 to digest chitin (in crustacean exoskeletons) and plant structural polysaccharides (e.g., cellulose
93 in seagrass).

94 We measured digestive enzyme activities in the guts of bonnethead sharks to examine
95 what compounds these fish could digest, and to infer their digestive strategy based on the
96 digestive enzyme activity patterns along their guts (Table 1; Fig. 2A). We, therefore, tested the
97 hypothesis that bonnethead sharks are “yield-maximizers” and consume relatively large meals
98 relatively infrequently (Fig. 1; German et al., 2015). Thus, sharks, including bonnetheads, would
99 be expected to have enzymatic patterns consistent with their yield-maximizing strategy that
100 would include some amount of microbial digestion in the hindgut (Fig. 2A; German et al., 2015).
101 Sharks are known to have long transit times (i.e., >20 hours) of food in the gut (Wetherbee et al.
102 1987), which is another indication of a yield-maximizing strategy towards digestion. However,
103 elasmobranchs (including sharks) have a relatively short intestine coupled to a “spiral valve”
104 (Fig. 2B), which is a convoluted region of the intestine that resembles a spiral staircase or a
105 rolled scroll of paper in cross section (Holmgren and Nilsson, 1999; Chatchavalvanich et al.,
106 2006; Theodosiou et al., 2007). Although it is accepted that the spiral valve (heretofore called
107 the spiral intestine) increases the absorptive surface area in the elasmobranch gut (Holmgren and
108 Nilsson, 1999; Chatchavalvanich et al., 2006; Wilson and Castro, 2011), patterns of digestive
109 enzyme activities in the anterior intestine vs. the spiral intestine, or how enzyme activities
110 change along the spiral intestine, are largely unknown (Kuz'mina 1990; Holmgren and Nilsson,
111 1999). Activities of the intestinal enzymes maltase, sucrase, trehalase, and alkaline phosphatase
112 have been measured in membrane vesicle preparations of dogfish spiral intestine tissue (Crane et
113 al., 1979), but it was not clear from where in the spiral intestine these vesicle preparations were
114 taken, or how activities change moving along the spiral intestine. Therefore, our investigation
115 also provides insight into the role of the spiral intestine as a site of digestion and/or absorption in
116 sharks.

117 We measured the activity levels of six digestive enzymes in the guts of bonnethead
118 sharks that reflected this species ability to digest carbohydrates (maltase, N-acetyl- β -D-
119 glucosaminidase, and β -glucosidase), protein (trypsin and aminopeptidase), and lipids (lipase;
120 Table 1). We measured the activities of these enzymes in the pancreas, anterior intestine, along
121 the spiral intestine, and in the colon (Fig. 2B), and used the activity patterns to infer whether
122 these sharks are rate- or yield-maximizers, predicting the latter. We did not measure digestive
123 enzyme activities (e.g., pepsin, chitinase) in the stomachs of the sharks, as that is the focus of a
124 different study, although a cursory examination of the stomach contents of the sharks used in this
125 study confirmed their carnivorous diet (Cortés et al., 1996; Bethea et al., 2007).

126 **2. Materials and Methods**

127 *2.1 Shark capture and tissue preparation*

128 Six bonnethead sharks were collected in gill nets off the coast of Cedar Key (29.115° N,
129 83.034° W) and Cumberland Sound (30.795° N, 81.492° W), Florida, USA. Sharks were
130 incidental mortalities from monthly surveys of shark nursery habitat within Florida coastal
131 waters (e.g., Bethea et al., 2011). Immediately following collection, freshly dead sharks were
132 measured (stretch total length \pm 0.5 cm) and then dissected on a cutting board kept on ice (4°C).
133 The sharks were 98 ± 9.4 cm (mean \pm SEM) in length, and were small adults or large juveniles.
134 Each digestive system was removed by cutting just anterior to the stomach and at the anus. The
135 pancreas was excised and frozen individually on dry ice in a 50 mL centrifuge vial. The guts
136 were gently uncoiled, measured, and the stomachs excised. The stomachs were placed in
137 individual bags and frozen on dry ice for later use in gut content analyses. The remaining
138 digestive tract was divided into the following sections: anterior intestine, spiral intestine (SI), and
139 colon (Fig. 2B). The SI was further subdivided into three sections of equal length: the proximal,

140 mid, and distal SI (Fig. 2B). Each section was emptied of their contents by pushing with the
141 blunt side of a razorblade, and the tissue rinsed with shark Ringer's solution; the contents and
142 intestinal tissues were then placed in separate 50 mL centrifuge vials and frozen on dry ice
143 (German and Bittong, 2009). Contents were recovered from each gut region of each shark.
144 Frozen samples were then shipped on dry ice to UC Irvine where they were stored at -80°C until
145 analyzed (within six months).

146 Gut tissues or contents from each gut region from individual sharks were weighed
147 (regional gut or content mass \pm 0.001 g) and homogenized following German and Bittong
148 (2009). Intestinal contents were homogenized in 25 mM Tris-HCl, pH 7.5, whereas intestinal
149 tissues were homogenized in 350 mM mannitol with 1 mM Hepes, pH 7.5. Colon tissue and
150 contents were homogenized in sodium acetate pH 5.5, based on the acidic conditions
151 documented in this gut region of the bamboo shark (*Chiloscyllium plagiosum*; Anderson et al.
152 2010). The supernatants of homogenates were collected and stored in small aliquots (100-200
153 μ l) at -80°C until just before use in spectrophotometric or fluorometric assays of digestive
154 enzyme activities. The protein content of the homogenates was measured using bicinchoninic
155 acid (German and Bittong, 2009; Smith et al., 1985). Stomach contents were cursorily examined
156 in all specimens confirming the carnivorous diet of the bonnetheads.

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158 *2.2 Assays of digestive enzyme activity*

159 All assays were carried out at 22°C in duplicate or triplicate using a BioTek Synergy H1
160 Hybrid spectrophotometer/fluorometer equipped with a monochromator (BioTek, Winooski,
161 VT). All assay protocols generally followed methods detailed in German and Bittong (2009),
162 unless otherwise noted. All pH values listed for buffers were measured at room temperature

163 (22°C), and all reagents were purchased from Sigma-Aldrich Chemical (St. Louis). All reactions
164 were run at saturating substrate concentrations as determined for each enzyme with gut tissues
165 from bonnethead sharks. Each enzyme activity was measured in each gut region of each
166 individual shark, and blanks consisting of substrate only and homogenate only (in buffer) were
167 conducted simultaneously to account for endogenous substrate and/or product in the tissue
168 homogenates and substrate solutions.

169 Maltase and Sucrase activities were measured following Dahlqvist (1968), as described
170 by German and Bittong (2009). We used 112 mM maltose (or 100 mM sucrose) dissolved in
171 200 mM phosphate buffer, pH 7.5 (sodium acetate pH 5.5 for the colon tissue and contents). The
172 maltase and sucrase activity was determined from a glucose standard curve and expressed in U
173 (μmol glucose liberated per minute) per gram wet weight of gut tissue.

174 β -glucosidase and N-acetyl- β -D-glucosaminidase activities were measured following
175 German et al. (2015), using 200 μM solutions of the substrates 4-methylumbelliferyl- β -D-
176 glucoside and 4-methylumbelliferyl-N-acetyl- β -D-glucosaminide, respectively, dissolved in 25
177 mM Tris-HCl (pH 7.5; sodium acetate pH 5.5 for the colon tissue and contents). Briefly, 90 μL
178 of substrate were combined with 10 μL of homogenate in a black microplate and incubated for
179 30 minutes. Following incubation, 2.5 μL of 1 M NaOH was added to each microplate well, and
180 the fluorescence read immediately at 365 nm excitation and 450 nm emission. Each plate
181 included a standard curve of the product (4-methylumbelliferone), substrate controls, and
182 homogenate controls, and enzymatic activity (μmol product released per minute per gram wet
183 weight tissue) was calculated from the MUB standard curve (German et al., 2011).

184 Trypsin activity was assayed using a modified version of the method designed by
185 Erlanger et al. (1961). The substrate, 2 mM N α -benzoyl-L-arginine-p-nitroanilide hydrochloride

186 (BAPNA), was dissolved in 100 mM Tris-HCl buffer (pH 7.5; sodium acetate pH 5.5 for the
187 colon tissue and contents). Trypsin activity was determined with a p-nitroaniline standard curve,
188 and expressed in U (μmol p-nitroaniline liberated per minute) per gram wet weight of gut tissue.

189 Aminopeptidase activity was measured using 2.04 mM L-alanine-p-nitroanilide HCl
190 dissolved in 200 mM sodium phosphate buffer (pH 7.5; sodium acetate pH 5.5 for the colon
191 tissue and contents). Aminopeptidase activity was determined with a p-nitroaniline standard
192 curve, and activity was expressed in U (μmol p-nitroaniline liberated per minute) per gram wet
193 weight of gut tissue.

194 Lipase (nonspecific bile-salt activated) activity was assayed using 0.55 mM p-nitrophenyl
195 myristate (in ethanol) in the presence of 5.2 mM sodium cholate dissolved in 25 mM Tris-HCl
196 (pH 7.5; sodium acetate pH 5.5 for the colon tissue and contents). Lipase activity was
197 determined with a p-nitrophenol standard curve, and expressed in U (μmol p-nitrophenol
198 liberated per minute) per gram wet weight of gut tissue.

199 The activity of each enzyme was regressed against the protein content of the
200 homogenates to confirm that there were no significant correlations between the two variables.
201 Because no significant correlations were observed, the data are not reported as U per mg protein.

202

203 *2.3 Statistical analyses*

204 Prior to all significance tests, a Levene's test for equal variance was performed and
205 residual versus fits plots were examined to ensure the appropriateness of the data for parametric
206 analyses. Where necessary, data were log-transformed prior to analysis. All tests were run using
207 SPSS statistical software (version 20). Comparisons of mass-specific enzymatic activities were

208 made among gut regions with ANOVA followed by a Tukey's HSD with a family error rate of P
209 = 0.05.

210

211 **3. Results**

212 The digestive enzyme activities of the bonnetheads generally followed the patterns
213 predicted in Table 1 and Fig. 2A. Trypsin showed a strong decreasing pattern moving distally
214 along the gut, with significantly ($P < 0.001$) higher activities in the pancreas than other gut
215 regions, and activities in the tissues tended to be of the same magnitude as those in the contents
216 (Fig. 3). Lipase activities showed a similar pattern moving along the guts as trypsin did
217 ($P = 0.003$), but there were elevated activities of this enzyme in the mid SI samples (Fig. 3).
218 Aminopeptidase activities showed a significant spike ($P < 0.001$) in the mid SI samples, and
219 activities were generally elevated in the tissues in comparison to the contents (Fig. 4). N-acetyl-
220 β -D-glucosaminidase activities were more elevated in the SI tissues than the intestine ($P < 0.001$),
221 but there were spikes in the activity of this enzyme in the tissue and contents ($P = 0.003$) of the
222 colon (Fig. 4), suggesting potential microbial production of this enzyme in the distal most
223 regions of the sharks' digestive tract. Maltase activities generally decreased ($P = 0.057$) moving
224 distally along the digestive tract, whereas β -glucosidase activities clearly spiked ($P = 0.062$) in the
225 colon of the sharks (Fig. 5). This latter result again suggests a microbial source for this enzyme,
226 although β -glucosidase activity was not reliably detectable in the gut contents of any gut region.

227

228 **4. Discussion**

229 Two key observations support the contention that bonnethead sharks adopt a yield-
230 maximizing strategy to digestion: the activity levels of N-acetyl- β -D-glucosaminidase and β -

231 glucosidase were elevated in the sharks' colons. These distal intestine enzyme activity spikes are
232 consistent with other fish species known to have a yield-maximizing strategy in the digestive
233 process (Skea et al., 2005; German et al., 2015), and suggest that bonnetheads have an active
234 microbial population in their hindguts that may aid in digestion. Overall, the patterns of
235 digestive enzyme activities in the bonnethead guts largely matched our predictions (Table 1; Fig.
236 2A), with pancreatic enzyme activities largely decreasing moving down the gut (Fig. 3), and
237 some brushborder enzyme activities (e.g., aminopeptidase) peaking in the mid spiral intestine
238 (Fig. 4). This latter result suggests that the spiral intestine is likely the most active, absorptive
239 section of the shark intestine, consistent with increased epithelial surface area in this gut region
240 (Holmgren and Nilsson, 1999; Chatchavalvanich et al., 2006; Wilson and Castro, 2011).
241 Measurements of nutrient transport rates along the elasmobranch epithelium are lacking, but we
242 hypothesize that these rates would be highest in the spiral intestine.

243 Sharks are known for consuming large meals, and holding those meals for extended
244 periods of time in the stomach (Wetherbee et al., 1987; Holmgren and Nilsson, 1999;
245 Papastamatiou 2007) before releasing chyme into the anterior intestine (Meyer and Holland,
246 2012). What happens to digesta after passage into the anterior intestine is largely unstudied in
247 many elasmobranchs, but flow likely follows the typical “plug-flow” model (Penry and Jumars,
248 1987) through the anterior intestine until the digesta reaches the spiral intestine, where transit
249 may be slowed (Holmgren and Nilsson, 1999). Slowed flow anywhere in the intestine is also
250 consistent with a yield-maximizing strategy. In one of the most detailed analyses of the
251 intestinal epithelium in any elasmobranch, Chatchavalvanich et al. (2006) showed that the spiral
252 intestine of the white-edge freshwater ray (*Himantura signifer*) had more complex folding
253 patterns (i.e., more absorptive surface area) than the anterior intestine in that species.

254 Observations in other elasmobranchs support this contention (Holmgren and Nilsson, 1999;
255 Wilson and Castro, 2011). Given that we observed that aminopeptidase and lipase activities
256 peaked in the mid spiral intestine, similar to mid-intestine spikes in activities of these enzymes in
257 other fish species that lack a spiral intestine (Chakrabarti et al., 1995; Harpaz and Uni, 1999;
258 Smoot and Findlay, 2000; German 2009; German et al., 2015), this portion of the intestine seems
259 to be the primary site of amino acid and fatty acid absorption in bonnetheads. Thus, the spiral
260 intestine essentially encompasses what is called the “intestine” in most other fishes, with
261 regionality of function changing from proximal to distal ends (German 2009; German et al.
262 2015), unlike some earlier cursory analyses that claimed little regionality in digestive enzyme
263 activity in the elasmobranch gut (Kuz'mina 1990).

264 The elevated trypsin and aminopeptidase activities in the bonnethead pancreatic and
265 spiral intestinal tissues, respectively, make sense, as the pancreas is the site of synthesis for
266 trypsin, and enterocytes the site of synthesis for aminopeptidase (Karasov and Martinez del Rio,
267 2007). However, the bonnethead trypsin and aminopeptidase activity levels are about 10X
268 higher than activities in the pancreatic or intestinal tissues of carnivorous teleost fishes measured
269 using the same methods and equations for calculations as this study (German, 2009; German et
270 al., 2015). Thus, bonnetheads appear to be efficient at digesting protein, which is probably an
271 important nutrient for these carnivorous animals. Lipase activity in the bonnetheads is not
272 exceptionally elevated in comparison to other fishes, but the broad distribution of lipolytic
273 activity along the gut (Fig. 3) suggests that bonnetheads likely readily digest lipids with great
274 efficiency. To our knowledge, there aren't any studies of protein and/or lipid digestibility in
275 sharks consuming their natural prey items, but Wetherbee and Gruber (1993) measured apparent
276 digestive efficiencies of 62-83% and 76-88% for energy and organic matter, respectively, in

277 lemon sharks (*Negaprion brevirostris*) consuming different sized meals of a fish diet. Given that
278 fish would be primarily protein and lipid (Horn, 1989) it follows that the high organic matter
279 digestibility by lemon sharks would primarily be of protein and lipid. We attempted to measure
280 amylolytic activity in the bonnetheads, but we were not able to reliably detect enzymatic activity
281 against starch, suggesting that bonnetheads may be poor at digesting soluble carbohydrates, like
282 starch. This is corroborated by the relatively low maltase activities in the bonnethead intestine
283 (Fig. 4), although using membrane vesicle preparations may improve detection of maltase (Crane
284 et al., 1979).

285 What is intriguing is the presence of elevated N-acetyl- β -D-glucosaminidase activity in
286 the colons of the bonnetheads. Bonnetheads clearly consume chitin with their diet rich in
287 crustaceans (Bethea et al., 2007), and like other fishes that consume chitin (Goodrich and Morita,
288 1977; Gutowska et al. 2004; German et al., 2010; German et al., 2015), this may be an important
289 source of carbon and nitrogen for these sharks. Indeed, N-acetyl- β -D-glucosaminidase activities
290 in the bonnethead intestine are also at least 5X higher than carnivorous, omnivorous, and
291 detritivorous teleost fishes measured using the same methods and equations for calculations as
292 this study (German and Bittong, 2009; German et al., 2015). The source of the N-acetyl- β -D-
293 glucosaminidase is likely endogenous along the gut walls of the anterior and spiral intestine, but
294 the spike in N-acetyl- β -D-glucosaminidase activity in the colon (including the colon contents)
295 strongly suggests a microbial origin of these activities in the hindgut (German and Bittong, 2009;
296 German et al., 2015).

297 Interestingly, the colons of sharks and skates are known to be acidic (pH 5.5-6.4,
298 depending on species), which is more acidic than their intestines (which tend to be pH 7.0-7.5;
299 Anderson et al., 2010), and is more similar to the pH of a typical vertebrate colon, which is a site

300 of microbial fermentation (Karasov and Martinez del Rio, 2007; Karasov and Douglas, 2013).
301 The colonic environment is anaerobic, which allows enteric microbes to use fermentative
302 pathways to produce short chain fatty acids (SCFA); these SCFA (e.g., acetate, propionate) are
303 the reason for the lower pH of the colon in most vertebrates and the SCFA can be absorbed by
304 the host and used for ATP production (Bergman, 1990; Stevens and Hume, 1998; Karasov and
305 Martinez del Rio, 2007; Karasov and Douglas, 2013). Although we didn't measure SCFA
306 production in the bonnetheads, SCFA production is well known in the hindguts of fishes, and is
307 usually higher in herbivores than in carnivores (Clements et al. 1995; Clements et al. 2014;
308 German et al. 2015), although some carnivores (e.g., largemouth bass, *Micropterus salmoides*)
309 do show seasonally high levels of SCFA production in their hindguts (Smith et al., 1996). The
310 omnivorous *Phytichthys chirus*, which also consumes a crustacean-rich diet, has elevated levels
311 of N-acetyl- β -D-glucosaminidase activities in its hindgut (German et al., 2015), further
312 suggesting that hindgut microbial digestion of chitin may be wide-spread in carnivorous fishes
313 with chitin-rich diets, as we see in the bonnetheads. The slower transit time of food through a
314 carnivorous gut is amenable to a yield-maximizing strategy, and the activities of microbially-
315 produced enzymes being elevated in the hindgut also support that carnivores can be yield-
316 maximizers with some reliance on microbial symbionts in the digestive process.

317 Along these lines, we were surprised to observe a spike in β -glucosidase activity in the
318 bonnethead colon, as this enzyme digests the breakdown products of cellulose and other β -
319 glucosides, like laminarin (German and Bittong, 2009). Up to 62% of young-of-the-year
320 bonnethead diet (by mass) can be composed of seagrass, which appears degraded by the time it
321 reaches the hindguts of the sharks (Bethea et al., 2007). Certainly, if bonnetheads have a
322 microbial community in their hindguts that are capable of degrading chitin, they may also be able

323 to degrade other β -glucosides. The activity levels of β -glucosidase in the bonnethead colon are
324 about 2X higher than those observed in the hindguts of *Cebidichthys violaceus*, an herbivorous,
325 teleost fish that digests algal material in its hindgut with the aid of an enteric microbial
326 community (German et al., 2015). Clearly, feeding trials to examine the digestibility of seagrass
327 by bonnetheads are necessary to confirm this supposition, but it does appear possible that
328 bonnetheads have the enzymatic machinery to degrade components of seagrass. Indeed, the
329 main carbohydrate in seagrass is cellulose, and sucrose is the photosynthate (Kuiper-Linley et al.,
330 2007). We did not readily detect sucrase activities in the bonnethead intestine, but attempting
331 sucrase assays on membrane vesicle preparations (as we suggest for maltase) may produce more
332 consistent results for this enzyme (Crane et al. 1979). Using stable isotope analysis ($\delta^{13}\text{C}$ and
333 $\delta^{15}\text{N}$ signatures), Bethea et al. (2011) showed that the bonnetheads occupied a different trophic
334 space (in particular, a lower trophic level from the perspective of $\delta^{15}\text{N}$) than their congener, the
335 scalloped hammerhead shark (*Sphyrna lewini*), which is more piscivorous. Although
336 invertebrate vs fish diets would be enough to result in niche segregation from the perspective of
337 stable isotope analysis, it is possible that the digestion of seagrass, and its epibionts, may
338 contribute to the lower $\delta^{15}\text{N}$ and enriched $\delta^{13}\text{C}$ signatures observed in bonnethead tissues relative
339 to scalloped hammerhead tissues (Bethea et al. 2011), but this needs to be explored in more
340 detail.

341 In conclusion, we measured digestive enzyme activities along the guts of bonnethead
342 sharks in an effort to understand their digestive strategy and discern what compounds they might
343 be able to digest. The patterns of enzymatic activity along their guts suggest that bonnetheads
344 take a yield-maximizing strategy to the digestive process, and that these sharks likely harbor an
345 enteric microbial community in their colons that may aid in digestion of complex carbohydrates

346 (e.g., chitin, cellulose). We also elucidated that the spiral intestine is likely the primary site of
347 digestion and absorption in the bonnethead gut, and future studies should focus on the spiral
348 intestine to discern the digestive capabilities of elasmobranchs, but also determine the role of the
349 anterior intestine, which is currently unclear. Indeed, there is broad interest in sharks, and to
350 better understand their ecological roles, we need to move beyond feeding observations and truly
351 grasp what they are eating, digesting, and excreting back into their environments in order to
352 make better predictions of how sharks will thrive in a changing world.

353

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360 we did not require any approval of an institutional animal care and use committee (i.e., the
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362

363 **References**

364 Anderson, W.G., Dasiewicz, P.J., Liban, S., Ryan, C., Taylor, J.R., Grosell, M.,
365 Weihrach, D., 2010. Gastro-intestinal handling of water and solutes in three species of
366 elasmobranch fish, the white-spotted bamboo shark, *Chiloscyllium plagiosum*, little skate,
367 *Leucoraja erinacea* and the clear nose skate *Raja eglanteria*. *Comp Biochem Physiol A*
368 155, 493-502.

369 Bergman, E., 1990. Energy contributions of volatile fatty acids from the gastrointestinal
370 tract in various species. *Physiol Rev* 70, 567-590.

371 Bethea, D.M., Carlson, J.K., Hollensead, L.D., Papastamatiou, Y.P., Graham, B.S.,
372 2011. A comparison of the foraging ecology and bioenergetics of the early life-stages of
373 two sympatric hammerhead sharks. *Bull Mar Sci* 87, 873-899.

374 Bethea, D.M., Hale, L., Carlson, J.K., Cortés, E., Manire, C.A., Gelsleichter, J., 2007.
375 Geographic and ontogenetic variation in the diet and daily ration of the bonnethead shark,
376 *Sphyrna tiburo*, from the eastern Gulf of Mexico. *Mar Biol* 152, 1009-1020.

377 Chakrabarti, I., Gani, M.A., Chaki, K.K., Sur, R., Misra, K.K., 1995. Digestive enzymes
378 in 11 freshwater teleost fish species in relation to food habit and niche segregation. *Comp*
379 *Biochem Physiology* 112A, 167-177.

380 Chatchavalvanich, K., Marcos, R., Poonpirom, J., Thongpan, A., Rocha, E., 2006.
381 Histology of the digestive tract of the freshwater stingray *Himantura signifer* Compagno
382 and Roberts, 1982 (Elasmobranchii, Dasyatidae). *Anat Embryol* 211, 507-518.

383 Clements, K.D., Angert, E.R., Montgomery, W.L., Choat, J.H., 2014. Intestinal
384 microbiota in fishes: what's known and what's not. *Mol Ecol* 23, 1891-1898.

385 Clements, K.D., Choat, J.H., 1995. Fermentation in tropical marine herbivorous fishes.
386 *Physiol Biochem Zool* 68, 355-378.

387 Clements, K.D., Raubenheimer, D., 2006. Feeding and nutrition, in: D.H. Evans (Ed.),
388 *The physiology of fishes*. CRC Press, Boca Raton, FL, 47-82.

389 Cortés, E., Manire, C.A., Hueter, R.E., 1996. Diet, feeding habits, and diel feeding
390 chronology of the bonnethead shark, *Sphyrna tiburo*, in southwest Florida. *Bull Mar Sci*
391 58, 353-367.

392 Crane, R.K., Boge, G., Rigal, A., 1979. Isolation of brushborder membranes in vesicular
393 form from the intestinal spiral valve of the small dogfish (*Scyliorhinus canicula*).
394 *Biochim Biophys Acta* 554, 264-267.

395 Crossman, D.J., Choat, J.H., Clements, K.D., 2005. Nutritional ecology of nominally
396 herbivorous fishes on coral reefs. *Mar Ecol Progr Ser* 296, 129-142.

397 Dahlqvist, A., 1968. Assay of intestinal disaccharidases. *Analyt Biochem* 22, 99-
398 107.

399 Day, R.D., German, D.P., Manjakasy, J.M., Farr, I., Hansen, J., Tibbetts, I.R., 2011.
400 Enzymatic digestion in stomachless fishes: how a simple gut accommodates both
401 herbivory and carnivory. *J Comp Physiol B* 181, 603-613.

402 Erlanger, B.F., Kokowsky, N., Cohen, W., 1961. The preparation and properties of two
403 new chromogenic substrates of trypsin. *Arch Biochem Biophys* 95, 271-278.

404 German, D.P., 2009. Do herbivorous minnows have "plug-flow reactor" guts? Evidence
405 from digestive enzyme activities, gastrointestinal fermentation, and luminal nutrient
406 concentrations. *J Comp Physiol B* 179, 759-771.

407 German, D.P., 2011. Digestive Efficiency, in: A.P. Farrel (Ed.), *Encyclopedia of Fish*
408 *Physiology: From Genome to Environment*. Academic Press, San Diego, 1596-1607.

409 German, D.P., Bittong, R.A., 2009. Digestive enzyme activities and gastrointestinal
410 fermentation in wood-eating catfishes. *J Comp Physiol B* 179, 1025-1042.

411 German, D.P., Horn, M.H., Gawlicka, A., 2004. Digestive enzyme activities in
412 herbivorous and carnivorous prickleback fishes (Teleostei: Stichaeidae): ontogenetic,
413 dietary, and phylogenetic effects. *Physiol Biochem Zool* 77, 789-804.

414 German, D.P., Nagle, B.C., Villeda, J.M., Ruiz, A.M., Thomson, A.W., Contreras-

415 Balderas, S., Evans, D.H., 2010. Evolution of herbivory in a carnivorous clade of
416 minnows (Teleostei: Cyprinidae): effects on gut size and digestive physiology. *Physiol*
417 *Biochem Zool* 83, 1-18.

418 German, D.P., Sung, A., Jhaveri, P.K., Agnihotri, A., 2015. More than one way to be an
419 herbivore: convergent evolution of herbivory using different digestive strategies in
420 prickleback fishes (family Stichaeidae). *Zool* (In press).

421 German, D.P., Weintraub, M.N., Grandy, A.S., Lauber, C.L., Rinkes, Z.L., Allison, S.D.,
422 2011. Optimization of hydrolytic and oxidative enzyme methods for ecosystem studies.
423 *Soil Biol Biochem* 43, 1387-1397.

424 Goodrich, T.D., Morita, R.Y., 1977. Incidence and estimation of chitinase activity
425 associated with marine fish and other estuarine samples. *Mar Biol* 41, 349-353.

426 Gutowska, M., Drazen, J., Robison, B., 2004. Digestive chitinolytic activity in marine
427 fishes of Monterey Bay, California. *Comp Biochem Physiol A* 139, 351-358.

428 Harpaz, S., Uni, Z., 1999. Activity of intestinal mucosal brush border membrane
429 enzymes in relation to the feeding habits of three aquaculture fish species. *Comp*
430 *Biochem Physiol Part A* 124, 155-160.

431 Hidalgo, M.C., Urea, E., Sanz, A., 1999. Comparative study of digestive enzymes in fish
432 with different nutritional habits. Proteolytic and amylase activities. *Aquacult* 170, 267-
433 283.

434 Holmgren, S., Nilsson, S., 1999. Digestive system, in: W.C. Hamlett (Ed.), *Sharks,*
435 *skates, and rays: the biology of elasmobranch fishes.* The Johns Hopkins University
436 Press, Baltimore, MD, 144-173.

437 Horn, M.H., 1989. *Biology of Marine Herbivorous Fishes.* *Oceanogr Mar Biol*

438 Ann Rev 27, 167-272.

439 Karasov, W.H., Douglas, A.E., 2013. Comparative Digestive Physiology. *Comprehens*
440 *Physiol* 3, 741-783.

441 Karasov, W.H., Martínez del Rio, C., 2007. *Physiological ecology: how animals process*
442 *energy, nutrients, and toxins*. Princeton University Press, Princeton, NJ USA.

443 Kuiper-Linley, M., Johnson, C.R., Lanyon, J.M., 2007. Effects of simulated green turtle
444 *regrazing on seagrass abundance, growth and nutritional status in Moreton Bay, south-*
445 *east Queensland, Australia*. *Mar Freshwat Res* 58, 492-503.

446 Kuz'mina, V.V., 1990. Characteristics of enzymes involved in membrane digestion in
447 *elasmobranch fishes*. *Zhur Evolyut Biokhim Fiziolog* 26, 161-166.

448 Meyer, C.G., Holland, K.N., 2012. Autonomous measurement of ingestion and digestion
449 *processes in free-swimming sharks*. *J Exp Biol* 215, 3681-3684.

450 Moran, D., Turner, S., Clements, K.D., 2005. Ontogenetic development of the gastrointestinal
451 *microbiota in the marine herbivorous fish *Kyphosus sydneyanus**. *Microb Ecol* 49, 590-
452 597.

453 Papastamatiou, Y.P., 2007. The potential influence of gastric acid secretion during
454 *fasting on digestion time in leopard sharks (*Triakis semifasciata*)*. *Comp Biochem*
455 *Physiol A* 147, 37-42.

456 Papastamatiou, Y.P., Lowe, C.G., 2004. Postprandial response of gastric pH in leopard
457 *sharks (*Triakis semifasciata*) and its use to study foraging ecology*. *J Exp Biol* 207, 225-
458 232.

459 Papastamatiou, Y.P., Lowe, C.G., 2005. Variations in gastric acid secretion during
460 *periods of fasting between two species of shark*. *Comp Biochem Physiol A* 141, 201-214.

461 Penry, D.L., Jumars, P.A., 1987. Modeling animal guts as chemical reactors. The
462 Am Nat 129, 69-96.

463 Skea, G., Mountfort, D., Clements, K.D., 2005. Gut carbohydrases from the New
464 Zealand marine herbivorous fishes *Kyphosus sydneyanus* (Kyphosidae), *Aplodactylus*
465 *arctidens* (Aplodactylidae), and *Odax pullus* (Labridae). Comp Biochem Physiol B 140,
466 259-269.

467 Skea, G., Mountfort, D., Clements, K.D., 2007. Contrasting digestive strategies in four
468 New Zealand herbivorous fishes as reflected by carbohydrase activity profiles. Comp
469 Biochem Physiol Part B 146, 63-70.

470 Smith, T., Wahl, D., Mackie, R., 1996. Volatile fatty acids and anaerobic fermentation in
471 temperate piscivorous and omnivorous freshwater fish. J Fish Biol 48, 829-841.

472 Smoot, J.C., Findlay, R.H., 2000. Digestive enzyme and gut surfactant activity of
473 detritivorous gizzard shad (*Dorosoma cepedianum*). Can J Fish AquatSci 57, 1113-1119.

474 Stevens, C.E., Hume, I.D., 1998. Contributions of Microbes in Vertebrate
475 Gastrointestinal Tract to Production and Conservation of Nutrients. Physiol Rev 78, 393-
476 427.

477 Theodosiou, N., Hall, D.A., Jowdry, A.L., 2007. Comparison of acid mucin goblet cell
478 distribution and Hox13 expression patterns in the developing vertebrate digestive tract J
479 Exp Zool 308B, 442-453.

480 Wetherbee, B.M., Gruber, S.H., 1993. Absorption efficiency of the lemon shark *Negaprion*
481 *brevirostris* at varying rates of energy intake. Copeia 1993, 416-425.

482 Wetherbee, B.M., Gruber, S.H., Ramsey, A.L., 1987. X-radiographic observations of
483 food passage through digestive tracts of lemon sharks. Trans Am Fish Soc 116, 763-767.

484 Wilson, J.M., Castro, L.F.C., 2011. Morphological diversity of the gastrointestinal tract in
485 fishes, in: M. Grosell, A.P. Farrell, C.J. Brauner (Eds.), The multifunctional gut of fish.
486 Elsevier, San Diego, 1-55.

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491 **Figure 1.** Cumulative nutrient gained (solid black line) by a fish as a function of time spent
492 processing a meal (modified from German et al. 2015). The slope of the black line labeled “Max
493 Rate” is the maximum rate at which the nutrient can be absorbed from the meal. A rate-
494 maximizing strategy is characterized by a line tangential to the curve (red line “R”), with
495 defecation of gut contents occurring at time 1 (t1). A portion of the nutrient consumed is lost in
496 the feces (“Wastage”), but at t1 the animal can take a new meal. This is the Rate-maximizing
497 strategy with high-intake. Maximum yield (blue line “Y”) is attained by extending processing
498 time to time 2 (t2), however, this is done at the cost of reduced digestive rate. In animals with
499 lower intake, this strategy tends to involve longer retention times of food in the gut and can
500 include microbial fermentation in the hindgut (especially in herbivorous vertebrates).

501

502 **Figure 2. A.** Potential patterns of digestive enzyme activities along a shark gut. Pancreatic
503 enzymes are made in the acinar cells of the pancreas (not shown). Thus, other than the pancreas
504 itself, activities of pancreatic digestive enzymes would be expected to be highest in the anterior
505 intestine (where they are carried after traveling down the ductus choledochus). Brush border
506 enzymes tend to peak in the mid intestine of many fishes, which would be the spiral intestine in
507 sharks. However, microbially-produced enzymes peak in the distal intestines of fish utilizing a
508 Yield-maximizing strategy because microbes tend to be more concentrated in the distal intestines
509 of fishes (see Skea et al. 2005; German et al. 2015). Fish adopting a Rate-maximizing strategy
510 would not show a spike in microbial digestive enzymes in their distal intestines (see German
511 2009; German and Bittong 2009; German et al. 2015). **B.** *Sphyrna tiburo* with its digestive tract.
512 For this study, the stomach was excised, and the remaining digestive tract was divided into the
513 anterior intestine, proximal-, mid-, and distal-spiral intestine (PSI, MSI, and DSI, respectively),
514 and colon.

515

516 **Figure 3.** Trypsin (top) and lipase (bottom) activities in gut tissue (left column) or gut contents
517 (right column) in different gut regions of *Sphyrna tiburo*. See Fig. 1 for gut region definitions.
518 Activities are mean \pm SEM. Trypsin or lipase activities were compared among gut regions
519 independently for tissue or contents with ANOVA followed by Tukey’s HSD multiple
520 comparisons test. Regional enzymatic activity values for an enzyme and gut fraction (tissue or
521 contents) that share a letter are not significantly different from one another ($P>0.05$). There were
522 not enough intestinal contents in which to perform the lipase assay, and hence these values are
523 missing from the lipase gut content graph (right bottom). SI = spiral intestine.

524 **Figure 4.** Aminopeptidase (top) and N-acetyl- β -D-glucosaminidase (NAG; bottom) activities in
525 gut tissue (left column) or gut contents (right column) in different gut regions of *Sphyrna tiburo*.
526 See Fig. 1 for gut region definitions. Note the different scales for the y-axis of the
527 aminopeptidase graphs. Activities are mean \pm SEM. Aminopeptidase or NAG activities were
528 compared among gut regions independently for tissue or contents with ANOVA followed by
529 Tukey's HSD multiple comparisons test. Regional enzymatic activity values for an enzyme and
530 gut fraction (tissue or contents) that share a letter are not significantly different from one another
531 ($P>0.05$). There were not enough intestinal contents in which to perform the aminopeptidase
532 assay, and hence these values are missing from the aminopeptidase gut content graph (right top).
533 SI = spiral intestine.

534 **Figure 5.** Maltase (top) and β -glucosidase (bottom) activities in gut tissue (left column) or gut
535 contents (right column) in different gut regions of *Sphyrna tiburo*. See Fig. 1 for gut region
536 definitions. Activities are mean \pm SEM. Maltase or β -glucosidase activities were compared
537 among gut regions independently for tissue or contents with ANOVA followed by Tukey's HSD
538 multiple comparisons test. Regional enzymatic activity values for an enzyme and gut fraction
539 (tissue or contents) that share a letter are not significantly different from one another ($P>0.05$).
540 There was no detectable β -glucosidase activity in the intestine, and hence these values are
541 missing from the β -glucosidase gut tissue graph (left bottom). β -glucosidase was not repeatedly
542 detectable in gut contents, and hence, this graph is not shared. SI = spiral intestine.

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