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# <sup>Q1</sup>A 10-year evaluation of the functional basis for regeneration habitat preference of trees in an African evergreen forest

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### ABSTRACT

The spatial distribution of tree juveniles in relation to light environments may reflect species differences in growth, survival, and functional traits and will shape the nature of forest regeneration. Long-term field experiments are important to evaluate this issue because of the potentially very long juvenile period in trees. Here, we combine a 10-year seedling survival-growth data with the results of community ordination and multivariate analyses of functional traits to ask how observed juvenile light guilds are related to species functional traits and seedling performance. We transplanted seedlings at a standardized height of 11 cm into the shaded understory and quantified their growth and survival for 10-year. Using the community-wide stem distribution data, we categorized 33 species including the focal 11 species to understory vs. gap/edge guilds. Then, we determined differences between the two guilds in seedling survival, growth, as well as seed size, adult height, and a series of leaf traits, including toughness and chemical traits (fiber, protein, phenolics, tannins, alkaloids, saponins). Among the 11 nonpioneer species whose seedlings were planted into the understory, there was no significant difference in 10-year survival between light guilds, but species in gap/edge guild tended to achieve greater height than species in the understory guild. The leaf chemical traits of 33 species did not differ between the two juvenile light guilds, but gap/edge species had smaller seeds, taller adults, and tougher leaves than understory species. We used logistic regression as a complementary approach to assess the extent to which plant traits varied between light guilds and the most parsimonious model based on AIC<sub>c</sub> ranking included only leaf toughness and had an Akaike weight of 0.52. In addition, across the 11 species planted as seedlings, these traits were not significantly related to survivorship or growth over 10 years. A Principle Components Analysis illustrated associations among traits. We conclude that light guilds in terms of juvenile stem distribution could not be explained by long-term field performance of postestablishment seedlings alone. Earlier seedling stage or later sapling stage may be more important in differentiation of light guilds. For the species examined difference in growth rates could be linked to seed size and adult stature, but not to the adult leaf chemical traits considered. These results suggest the importance of examining ontogenetic shifts and relationships among functional traits for a better understanding of regeneration strategies of tropical trees.

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### 1112 **1. Introduction**

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Canopy gaps create environmental heterogeneities, which in turn contribute to the maintenance of tree species diversity (Denslow, 1987; Brokaw and Busing, 2000; Wright, 2002; Obiri

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and Lawes, 2004). Historically, forest ecologists have recognized 16 species differences in gap dependency based on observation of the 17 relative abundance of stems in gaps vs. shaded understory (Shirley, 18 1943). While tree species are often classified into a few of juvenile 19 light guilds, this is a continuum (Grubb, 1996; Poorter et al., 2005; 20 Gilbert et al., 2006). At one end are pioneers that specialize on gaps 21 for regeneration, growth, and reproduction, while at the other 22 extreme are highly shade-tolerant species whose saplings are 23 preferentially distributed in the shaded understory (Hubbell and 24 Foster, 1986; Swaine and Whitmore, 1988; Dalling and Hubbell, 25

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26 2002). However, most tree species have intermediate light 27 requirements for regeneration (Augspurger, 1984; Wright et al., 28 2003). Furthermore, dispersal limitation and stochasticity may 29 play an important role in determining which species arrive and 30 persist in a given gap (Hubbell et al., 1999; Harms et al., 2000; 31 Brokaw and Busing, 2000). Still, contrasting these groups is useful 32 for determining suites of traits that vary in concert and for 33 providing a mechanistic understanding of the apparent sorting of 34 tree species along light gradients (Westoby and Wright, 2006). The 35 next logical challenge to ecologists is to identify the functional 36 traits that contribute to observed differences in light guilds of 37 juveniles and adults among coexisting tree species (Poorter, 2007; 38 Poorter et al., in review).

39 Spatial distributions of saplings along light gradients are shaped 40 as a cumulative consequence of three life stages: seeds, young 41 seedlings, and older established seedlings. A species' ability to 42 colonize gaps is thought to be achieved by producing a large 43 number of small seeds or maintaining these seeds in the soil seed 44 bank (Dalling et al., 2002). Light requirements for germination and 45 early seedling survival strongly constrain the spatial distribution of 46 seedlings during the first stages of life and have lasting effects on 47 sapling distributions (Augspurger, 1984; Pearson et al., 2002). 48 Survival and growth of established seedlings further modify the 49 distribution patterns of saplings relative to treefall gaps. Unfortu-50 nately, this last stage is the least well understood, as most existing 51 comparative or community-wide studies of growth and survival 52 are relatively short-term and limited to early seedling (Augspur-53 ger, 1984; Kitajima, 1994; Poorter, 1999; Gilbert et al., 2006) or 54 sapling stages (e.g., with stems >2 m tall and 1 cm DBH, Hubbell 55 and Foster, 1986; Poorter et al., 2003). There are few data on the 56 critical transitional stage from seedlings to saplings. The short 57 duration of many comparative studies is unfortunate, because 58 recruitment of seedlings to saplings can take many years especially 59 in the shaded understory where growth rates are extremely slow 600 2 (Connell and Green, 2000; Osada and Takeda, 2003).

61 In this paper we report growth and survival of experimentally 62 planted seedlings in the understory over a 10-year period along 63 with other selected functional traits for tree species in a moist-64 evergreen forest at Kibale National Park, Uganda. Our two main 65 objectives are to explore (1) how species contrasting in stem 66 distribution between understory vs. gap/edge differ in growth and 67 survival in the understory during the seedling-sapling transition, 68 and (2) what additional functional traits are associated with 69 preferred regeneration habitats among tree species in Kibale. For 70 functional traits, we examine seed size, maximum adult height, 71 and adult leaf traits. These are traits often examined in recent 72 analyses of functional traits encompassing multiple sites and 73 species in the Neotropics (Wright et al., 2007; Poorter et al., in 74 review). Our report is the first such analysis that we are aware of 75 for a wet African forest to explore functional traits relationships 76 with juvenile distribution, growth, and survival.

### 77 2. Materials and methods

### 78 2.1. Study site

79 This study was conducted in Kibale National Park in western 80 Uganda (795 km<sup>2</sup>; 0° 13′-0° 41′N and 30° 19′-30° 32′E; Chapman 81 et al., 1997). Kibale is a mid-altitude moist-evergreen forest with a 82 relatively species poor flora (68 tree species were identified in 83 4.8 ha of vegetation sampling; Chapman et al., 1997). The study 84 was conducted at Kanyawara (~1500 m elevation), which receives 85 approximately 1719 mm of rainfall annually (1990-2006) that 86 peaks during two rainy seasons and is well disperse throughout the 87 year, falling on average 166 days per year. Because of this well dispersed rain, the forest floor rarely dries and wilting of 88

understory herbs is infrequent. The annual mean daily maximum temperature of 23.1 °C and minimum temperature of 15.1 °C. The soils of the region are dark gray to red sandy loams or sandy clays and the fertility of the soils range from fair to good (for details on soils see Struhsaker, 1997).

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The forest is classified as transitional between lowland rain and montane forest with a canopy height averaging 25–30 m, and natural disturbances leading to treefall gaps are common (Howard, 1991). The forest is notable for its lack of aggressive colonizers (e.g., *Musanga* spp., *Cercropia* spp.) typical of other tropical regions, but other early successional species are present (e.g., *Albizia grandibracteata, Polyscias fulva, Trema orientalis*; Zanne and Chapman, 2005).

### 2.2. Species and assessment of habitat association type

The determination of light guild is difficult without long-term 103 monitoring of growth and mortality of marked individuals. Hence, 104 statistically significant bias of stem distribution to gaps vs. shaded 105 understory (Welden et al., 1991), as well as expert opinions based on 106 field observations (Mostacedo and Fredericksen, 1999), are still 107 commonly used methods for assessing light guilds. Poorter et al. 108 (2005) developed a method to assess juvenile light demand as 109 continuous variable derived from logistic regression of crown-110 exposure index against individual height. The relationships of 111 functional traits and juvenile vital rates achieve using this index is 112 similar to their relationships with light guild classes based on expert 113 opinions (Poorter and Kitajima, 2007). In this paper, we used two 114 means of assessing species light guilds, one based on spatial bias of 115 stem distribution to forest understory vs. gap/edge (Zanne and 116 Chapman, 2005), and another based on expert opinion of how 117 common seedlings and saplings were in the shaded understory. 118

The statistical assessment of stem distribution described in detail by Zanne and Chapman (2005) is briefly summarized here. Trees and shrubs of all size classes were enumerated in 24 small plots  $(5 \text{ m} \times 5 \text{ m})$  in each of four habitats (closed canopy forest, treefall gaps, forest/grassland edge, and grassland; *N* = 96 plots). For gap plots, treefall gaps were located throughout the forest in which the crown of the fallen trees had no fine branches and leaves remaining, vet little decay of the bole had occurred. Gap size averaged  $300 \text{ m}^2$ (range: 39–636 m<sup>2</sup>). Since few large trees and shrubs >2 m tall occurred in small plots (5 m  $\times$  5 m), trees and shrubs of larger sizes were also sampled in large plots (10 m  $\times$  50 m) randomly located in forest and grassland habitats (not gap and edge habitats). To determine confidence intervals of stem counts in different habitat types, the data were randomized with a custom-written program in R 1.6.2 (R Development Core Team, 2002). The natural structure of the data within the plots was maintained by randomly reassigning plots to habitats without replacement for 10,000 iterations. Significant species associations with a habitat were reported if observed stem counts were more extreme than the 95% confidence limits. For 10 species not found in these habitat plots or found only rarely (Table 2), habitat associations were based on reports in Eggeling and Dale (1952), Polhill (1952), Hamilton (1991), Katende et al. (1995), and Lwanga (1996). For the analysis reported here, gap and edge associated species were combined as "gap/edge" species in contrast to "understory" species for more balanced sample size between groups.

Our second method of assessing juvenile light guilds is based on 145 146 the abundance of seedlings and saplings in shaded understories (common, frequent, or rare in the understory). This relied on data 147 of small individuals (height <0.5 m) in the plot data of Zanne and 148 Chapman (2005) as well as published information (Eggeling and 149 Dale, 1952; Polhill, 1952; Hamilton, 1991; Katende et al., 1995; 150 Lwanga, 1996), and an independent assessment made by Peter 151 Grubb based on his experiences with seedlings and sapling in 152

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#### Table 1

The 10-year survival and mean height growth of seedlings transplanted at a common size of 11 cm along understory transects

	Stem distribution bias	Seedlings and saplings in shade	# Planted	Survival (%)	Growth (cm)
Balanites wilsoniana	Understory	Common	54	6	18
Chrysophyllum gorungosanum	Understory	Common	59	29	36
Uvariopsis congensis	Understory	Common	64	63	43
Symphonia globulifera	Understory	Common	11	27	55
Mimusops bagshawei	Understory	Common	57	30	57
Cordia millenii	Gap/edge	Common	13	40	55
Monodora myristica	Gap/edge	Often	22	50	37
Tabernaemontana sp.	Gap/edge	Often	9	33	71
Warbugia stuhlmanni	Gap/edge	Rare	16	50	66
Ficus exasperata	Gap/edge	Rare	27	7	144
Pseudospondias microcarpa	Gap/edge	Rare	64	55	178
Understory	Summary		49	31	42
Gap/edge	Summary		25	39	92
Common	Summary		43	33	44
Often	Summary		16	42	54
Rare	Summary		36	37	129

Q6 Species are grouped according to their preferred light guilds based on stem distributions and seedling and sapling abundance in shade (see Section 2). Bold letters for stem distribution bias indicate the species reported in Zanne and Chapman (2005).

153 Kibale (P. Grubb, personal communication). All 15 "understory" 154 species are common as seedling and saplings in shade, whereas 9 of 155 18 "gap/edge" species are classified to be "common" in shade as 156 seedlings (Table 2). This latter group corresponds to "partial shade-157 tolerant" species that can establish in shade, but strongly require 158 gaps to reach maturity (Poorter and Kitajima, 2007). The remaining 159 "gap/edge" species are observed only "often" or "rare" in shaded 160 understories as seedlings and saplings, corresponding to pioneers 161 Q3 or open swamp species (Grubb, 2006; P. Grubb personal 162 communication). While the analysis using the three guild groups 163 yielded similar trends, the small number of species in each group 164 limits the statistical power. Thus, we report only the results from 165 comparing two guilds. Approximately the same number of species 166 from each guild was used for seedling transplant experiment 167 (Table 1). For examination of the relationships of seed mass, adult 168 height, and leaf traits with light guilds, we included an additional 22 common tree species in Kibale, plus the 11 species used in the 169 170 seedling experiment (Table 2).

### 171 2.3. Seedlings transplanted to forest understory

172 Seedlings were grown from seeds in a nursery under a semitransparent roof. Seeds germinated on wet cloth towels were 173 174 planted into individual pots (250 ml) in topsoil collected from the 175 forest and seedlings were grown in the nursery (see Chapman and 176 Chapman, 1996 for details). When seedlings were approximately 177 11 cm tall (2-3 months after germination), they were transplanted 178 at 5 m intervals along 100-200 m transects (10 in total) that were 179 separated from each other by approximately 100 m. The species 180 available for planting on a specific date were randomly assigned to 181 transect location, avoiding locations within 20 m of any canopy 182 gaps on a transect. Seedlings were transported to the field in the 183 pots they grew in and then bare-rooted seedlings were planted into 184 holes dug in the soil and were watered on the day of planting only. 185 Seedlings that died in the first month were replaced. Transplanting 186 took place between October 1990 and May 1992 as nursery-grown 187 seedlings became available, with most (70%) being transplanted 188 during 1990–1991. As our objective was to study the survival and 189 growth of established seedlings rather than very young seedlings, 190 this experimental protocol circumvented the vulnerable emer-191 ging-seedling stage and allowed comparison of growth and 192 survival across species. Plants were revisited opportunistically 193 during the 10-year experiment to monitor survival, causes of 194 death, and growth. Here, growth was assessed by height, and 195 survival was assessed by %survival at the end of the study.

### 2.4. Measurement of seed size, adult leaf traits, and adult stature 196

Seed size was estimated at the mean length of approximately 30 197 seeds per species, typically collected from five fruiting trees. The 198 chemical and physical leaf characteristics of adult leaves were 199 determined using leaves collected typically from three locations in 200 the middle of adult crowns from one adult tree that was found in a 201 habitat that was thought to represent the species preferred habitat 202 (an analysis of interspecific variation in the chemical properties of 203 leaves at this site can be found in Chapman et al., 2003). Although 204 205 changes in light environment and ontogenetic shifts can cause changes in leaf traits, adult leaf traits and juvenile functional traits 206 207 may be linked via life history correlations (Poorter, 2007; Wright et al., 2007). Leaves were dried, stored in sealed plastic bags, 208 transported to North America, and ground in a Wiley mill to pass 209 through a 1-mm mesh screen. Protein (estimated from nitrogen 210 content), fiber (ADF), saponins, and presence/absence of alkaloids, 211 and cyanogenic glycosides were determined following the proce-212 dures outlined in Chapman and Chapman (2002). Nitrogen was 213 evaluated because of its importance to plants and because many 214 herbivores select food bases on protein content. Fiber represents an 215 important structural element of leaves and often serves to deter 216 herbivore foraging. The secondary compounds that were selected 217 are a small proportion of the great diversity of compounds plants use 218 to defend their leaves. The selected compounds were those that have 219 documented impact of herbivore foraging and that we were 220 equipped to analyze (Chapman and Chapman, 2002). None of the 221 study species had cyanogenic glycosides. Leaf fracture toughness, 222 tannins, and total phenolic estimates were obtained from Dominy 223 (2001). Maximum adult height data were derived from the literature 224 (Eggeling and Dale, 1952; Polhill, 1952; Hamilton, 1991; Katende 225 et al., 1995; Lwanga, 1996). 226

### 2.5. Statistical analysis

Our main objective was to compare growth, survival, and228functional traits of species between light guilds. Ten-year survival229and growth of seedlings, as well as functional traits, were230compared between the two guilds with a two-tailed *t*-test with231the degrees of freedom appropriately adjusted if the variances of232growth were correlated with functional traits.233234

Furthermore, to avoid type 1 error associated with multiple *t*-235 tests of the same hypothesis (i.e., difference between two light guilds); presence/absence in shade was examined as a function of 237

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Table 2

Functional traits (seed size, maximum adult height, and adult leaf traits) of 33 common tree species in Kibale National Park, Uganda, including 11 species used in the seedling transplanting experiment (marked with (\*))

Species	Stem	Seedlings and	Seed	Adult	Fiber	Protein	Saponins	Alkaloids	Phenolics	Tannins	Toughness
	distribution	saplings in shade	Size (cm)	height (m)	(%)	(mg/g)	(mm)	P/A	(%)	(%)	(J/m²)
Symphonia globulifera <sup>*</sup>	Understory	Common	7.05	40	30.6	11.2	14		0.5	1.7	580
Newtonia buchananii	Understory	Common	5.8	50	38.5	17.7	5	0	1.87	7.2	438
Balanites wilsoniana <sup>*</sup>	Understory	Common	4.77	35					0.67	0	892
Lovoa swynnertonii	Understory	Common	4.5	40	52.9	14.2	5	0	0.7	1	521
Strombosia scheffleri	Understory	Common	3.66	30	31.4	20.1	19	0	0.13	0	881
Rothmannia urcelliformis	Understory	Common	3.54	10	23.7	17.2	11	0	0.17	0	461
Cola gigantea	Understory	Common	2.85	50							
Chrysophyllum gorungosanum	Understory	Common	2.45	45	38.8	17.1	14	0	2.08	5.7	1022
Mimusops bagshawei <sup>*</sup>	Understory	Common	1.54	40	41	10.6	3	0	2.89	6	1024
Pouteria altissima	Understory	Common	1.19	50	41.5	16.8	12	0	3.5	3.8	1165
Uvariopsis congensis <sup>*</sup>	Understory	Common	1.12	15	33.9	20.6	0.5	1	0.13	0	272
Blighia unijugata	Understory	Common	1.01	20	35.6	15	5	0	0.28	0	1720
Trilepisium madagascariense	Understory	Common	0.74	30	27.8	19	14	0	1.41	3.5	1508
Euadenia eminens	Understory	Common	0.67	20					0.07	0	267
Diospyros abyssinica	Understory	Common	0.44	30	27	21.1	2	1	2.26	0	489
Cordia millenii <sup>*</sup>	Gap/edge	Common	4.12	45	46.5	13	2	0			
Lepisanthes senegalensis	Gap/edge	Common	1.5	15	44.8	12.6	5	0	1.72	3.6	596
Antiaris toxicaria	Gap/edge	Common	0.73	45	38.3	27.2	10	0	1.87	3	726
Dovyalis macrocalyx	Gap/edge	Common	0.44	6	42.3	10.2	14	0			
Teclea nobilis	Gap/edge	Common	0.44	25	26.8	20.4	5	1	0.61	0	282
Coffea eugenoides	Gap/edge	Common	0.36	4					1.37	0	602
Celtis africana	Gap/edge	Common	0.24	30	14.8	23.7	5	0	0.41	0.5	292
Fagaropsis angolensis	Gap/edge	Common	0.21	30	11.5	24.8	10	1			
Pancovia sp. near turbinata	Gap/edge	Common		20	48.1	18.1	0	0	1.65	1.1	361
Monodora myristica*	Gap/edge	Often	2.62	30	37	13.9	16	0	0.33	0	1110
Tabernaemontana sp.*	Gap/edge	Often	0.75	15	33.2	24.7	15	0			
Cordia abyssinica	Gap/edge	Rare	1.92	15					0.33	0	462
Pseudospondias microcarpa <sup>*</sup>	Gap/edge	Rare	1.41	30	36.1	12.9	12	0	1.67	5.6	300
Warbugia stuhlmanni <sup>*</sup>	Gap/edge	Rare	0.75	40	37.9	12.1	15	1			
Trema orientalis	Gap/edge	Rare	0.1	15					0.8	0	245
Ficus exasperata <sup>*</sup>	Gap/edge	Rare	0.01	25	22.2	26.1	12	0	0.39	0.2	515
Neoboutonia macrocalyx	Gap/edge	Rare	0.68	15	23.8	20.1	8	0			
Macaranga schweinfurthii	Gap/edge	Rare	0.31	15	25.2	13.5	16	0	5.06	35.6	673
Understory	Summary		2.76	33.67	35.23	16.72	8.71	0.18	1.19	2.06	802.86
Gap/edge	Summary		0.98	23.33	32.57	18.22	9.67	0.20	1.35	4.13	513.67
Common	Summary		2.15	30.21	34.79	17.53	7.78	0.21	1.21	1.86	704.95
Often	Summary		1.69	22.50	35.10	19.30	15.50	0.00	0.33	0.00	1110.00
Rare	Summary		0.37	22.00	27.28	17.95	12.75	0.25	2.08	11.93	477.67

Species are sorted by stem distribution to understory vs. gap/edge (bold: reported in Zanne and Chapman, 2005), expert opinion on seedling and sapling abundance in shade, and seed size (large to small). Adult leaf chemistry is on a dry mass basis (fiber, protein), relative assay of a unit dry mass (mm of foam rise for saponin, % of gallic acid standard for phenolics, and % of quebracho tannin standard for tannin), or presence/absence (alkaloids; 1: present, 0: absent). Values are missing when not tested. (\*\*) open swamp species.

the various leaf traits using logistic regression. We used sevencandidate models of leaf characters (starting with all variables and

progressively removing the least significant variable) and ranked them according to Akaike's Information Criterion (AIC<sub>c</sub>) adjusted for small sample size (Anderson et al., 2000). Akaike weights are standardized to a scale of 0–1 and give the approximate probability that a model "i" is the best model in the set of models considered. The 20 species in which all leaf traits were quantified were included in the models.

247 Multivariate trait associations were examined using Principal 248 Components Analyses (PCA) with varimax rotation and all variables 249 were log transformed. As we did not have data for all variables for all species, a compromise between species number and variable 250 inclusion was made; an analysis containing eight variables and 251 252 19 species was the best compromise allowing the most species and 253 variables. Statistical analyses were done with SPSS 14.0 (SPSS Inc., Chicago, IL) and JMP v 3.0 (SAS Institute, Cary, NC). 254

### 255 3. Results

### 256 3.1. Seedling survival and growth in the understory

Over the 10 years, survivorship in the forest understory was
extremely variable among regeneration strategies and species
(range 6–63%), averaging 35.5% (S.D. = 18.5%; Table 1). There were

no differences in 10-year survival between understory vs. gap/edge 260 species (t = -0.712, d.f. = 9, p = 0.494), or between species that 261 were common vs. rare as seedlings in shade (t = -0.561, d.f. = 9, 262 p = 0.589; Table 1). However, there was a tendency for gap/edge 263 species to grow faster than understory species (t = -2.094, 264 d.f. = 5.9, p = 0.082), as well as a trend for species whose seedlings 265 were rare in shade to grow faster than those whose seedlings were 266 commonly observed in shade (t = -2.138, d.f. = 3.3, p = 0.114). This 267 latter result is particularly interesting, given that growth was 268 measured for 10 years in the shaded understory, without any 269 conspicuous gap-opening over these seedlings. 270

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### 3.2. Seed size, adult stature, and leaf traits

Species classified as belonging to the edge/gap guild had significantly smaller seeds (t = -3.00, d.f. = 20.4, p = 0.007) and smaller maximum adult height (t = 2.357, d.f. = 31, p = 0.025) than understory species. In contrast, there was no difference between species common and those rare as juveniles in shade (seed size t = 1.164, d.f. = 30, p = 0.253; adult height t = 0.729, d.f. = 31, p = 0.471; Table 2).

Height growth of 11 species in the understory was only weakly and negatively correlated with seed size (r = -0.50, p = 0.11) and was not correlated with maximum adult height. Five species tested positive for alkaloids; three were gap/edge species, two were

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#### Table 3

Eigenvalues and factor loadings for seed size, adult height, and leaf traits for the first four axes using Principal Components Analyses for 19 species for which all data were available

	Axis 1	Axis 2	Axis 3	Axis 4
Eigenvalue	1.98	1.98	1.54	1.22
Cumulative % variance	24.7	49.4	68.7	83.9
Seed size (cm)	0.85	-0.22	0.13	0.14
Protein (mg g <sup>-1</sup> )	-0.80	-0.27	-0.18	0.38
Fiber (%)	0.75	0.19	-0.03	0.40
Phenolics (%)	-0.05	0.95	0.06	0.18
Tannins (%)	0.13	0.93	0.18	-0.01
Saponins (mm)	0.04	0.16	0.90	-0.03
Toughness (J m <sup>-2</sup> )	0.13	0.05	0.79	0.20
Maximum adult height (m)	0.11	0.14	0.17	0.90

Numbers in bold denote strong factor loadings (>0.50).

283 understory species, and four and one were common and rare as 284 juveniles in shade, respectively. Seed size was weakly positively 285 correlated with fiber (r = 0.38, p = 0.059) and negatively correlated 286 with protein contents of adult leaves (r = -0.39, p = 0.049), which 287 were negatively correlated with each other (r = -0.55, p = 0.003). 288 However, no significant differences were found between light 289 guilds in any of the chemical leaf characteristics (saponins, 290 phenolics, tannins, p > 0.4 in all cases). Understory species though 291 had tougher leaves than gap/edge species (t = -2.41, p = 0.028), as 292 did species whose juveniles are common in shade relative to those 293 whose juveniles are rare in shade (t = -2.06, p = 0.052).

294 We used logistic regression as a complementary approach to 295 these t-tests to assess the extent to which plant traits varied 296 between light guilds. This revealed a significant difference for 297 toughness, but not for any other the other leaf characteristics (all 298 chemical traits p > 0.4). The most parsimonious model based on 299 AIC<sub>c</sub> ranking included only leaf toughness and had an Akaike 300 weight of 0.52. No significant correlations were found between the 301 leaf chemical traits and seedling survival or growth over 10 years 302 in the understory.

303 In a PCA of plant traits, seed size and fiber were positively and 304 protein was negatively related to Axis 1 and phenolics and tannins 305 were positively related to Axis 2 (Table 3). Axis 1 and Axis 2 explain 306 an equal proportion (24.7%) of total variance, while Axis 3 explains 307 an additional 19.8%. Saponins and leaf toughness were positively 308 related to Axis 3 and adult height was positively related to Axis 4. 309 None of the axes were correlated to seedling growth or survival, 310 which could be due to limited statistical power from low sample 311 sizes.

#### 312 4. Discussion

313 Overall, the two light guilds based on stem distribution bias to 314 understory vs. gap/edge differed only in three key functional traits. 315 Surprisingly, understory-survival of post-establishment seedlings 316 during 10 years showed no obvious relationship with light guilds 317 in shade. However, there was a tendency for gap/edge species to 318 grow faster than understory species in the shaded understory. 319 Similarly, 33 species did not show difference in relation to light 320 guilds with respect to leaf chemical traits (fiber, protein, phenolics, 321 tannins, alkaloids, saponins), except that gap/edge species had 322 smaller seeds, shorter adults, and less tough leaves compared to 323 understory species. However, these traits were not significantly 324 related to survivorship or growth of seedlings transplanted to the 325 understory.

Several aspects specific to this study may have contributed to some of these unpredicted results. First, these seedlings were transplanted at the post-establishment stage (2–3 months after germination) at a common height of 11 cm, and any seedlings that died within the first month were replaced. It is possible that mortality during and immediately after germination (Augspurger, 331 1984; Swaine and Whitmore, 1988), rather than mortality at the 332 post-establishment phase, is the important discriminating factor 333 for seedling establishment in shade vs. gap/edge. Also, classifica-334 tion of species into just two light guilds is too simple for 335 assessment of regeneration light requirements. Unfortunately, 336 the number of individuals and samples included in our analysis 337 restrict use of continuous index for light demands, such as the 338 average crown-exposure index at a given height (Poorter et al., 339 2005; Poorter, 2007) or one based on the growth and survival of 340 341 sufficient number of individuals (Gilbert et al., 2006).

342 The finding of greater height growth for seedlings of gap/edge species than understory species, especially those rarely found in 343 344 shade as juveniles, is in agreement with other results from studies 345 of the early seedling stage (Kitajima, 1994; Dalling et al., 1999; 346 Poorter, 1999; Walters and Reich, 1999). This suggests that species-specific traits that enhance growth rates, such as high 347 specific leaf area and biomass allocation to shoot extension, may be 348 349 the main determinants of height growth across light gradients (Kitajima, 1996; Walters and Reich, 1999; Poorter, 2001; Poorter 350 et al., 2003). Although some suggest that this tendency (i.e., slower 351 growth for more shade-tolerant species) disappear after early 352 seed-reserve dependent period (Sack and Grubb, 2001, 2003), we 353 found that cumulative height growth over 10 years was 354 continuously greater in shade for gap/edge species than unders-355 356 tory species.

Seed size is considered an important correlate of life history and 357 shade tolerance (Pearson et al., 2002; Rose and Poorter, 2003; 358 Moles and Westoby, 2004). In agreement with this, mean seed size 359 was greater for the understory guild than for in gap/edge guild in 360 Kibale. In a separate experiment addressing early seedling growth 361 and survival of 24 tree and shrub species in Kibale, Zanne (2003) 362 found a positive correlation between seed mass and survival, but 363 364 not between seed mass and growth rates for young seedlings, regardless of whether seedlings were planted in the understory, 365 366 gaps, edges, or grasslands. However, in our study seed size was not significantly correlated with post-establishment survival and 367 growth during the 10-year period. One likely explanation is that 368 seed size advantage disappears with time (Sack and Grubb, 2001, 369 370 2003; Moles and Westoby, 2004). Also, 11 species may insufficient to detect seed size relationship with survival, growth, and adult 371 height. Among four functional traits (seed mass, wood density, 372 specific leaf area, and maximum adult height), seed mass is the 373 374 best correlate of diameter growth rates and survival rates of stems >10 cm in a data set including over 200 species of neotropical tree 375 376 species from five sites (Poorter et al., in review). In the same data set, seed size is positively correlated with adult height consistently 377 across sites (Wright et al., 2007). The relationship of seed size with 378 adult height may be triangular, however; short species are 379 constrained from having large seeds, while tall species may have 380 large or small seeds (Grubb et al., 2005). In future research 381 consideration should be made of the fact that, unlike tropical 382 383 forests of many other regions, African forests still have abundant 384 large bodied seed dispersers, such as elephants (Loxodonta 385 africana) and chimpanzees (Pan troglodytes).

Because adult height should be positively correlated with adult 386 387 light environment (Thomas and Bazzaz, 1999), it is a surrogate for 388 adult light niche that can be compared to juvenile light niche (Poorter, 2007). In a Bolivian moist forest, average crown-exposure 389 390 index for 2 m tall samplings was greater for species with taller adults, suggesting that species more light demanding as adults 391 392 tend to also be more light demanding as juveniles (Poorter, 2007). In contrast, among 33 species in our study, mean adult height was 393 significantly higher for understory than gap/edge species. In the 394 Bolivian site and Kibale, there is a wide variation in adult height 395 within understory guild. However, the upper canopy layer is 396

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dominated by more pioneer species in the Bolivian site than in
Kibale, perhaps reflecting difference in historical disturbance
regime and seasonality. In the evergreen forest of Kibale, treefall
gaps tend to be small and filled by horizontal expansion of canopy
species rather than by pioneers (Zanne and Chapman, 2005).

402 Other studies have shown that adult leaf traits, such as specific 403 leaf area (=leaf area per unit leaf mass), nitrogen contents, and 404 photosynthetic rates, are significantly correlated not only with 405 adult light environment (Thomas and Bazzaz, 1999, in SE Asian 406 forest), but even more strongly so with juvenile light environment 407 (Poorter, 2007; in a seasonal Bolivian forest). Our study did not 408 examine specific leaf area, but protein content (i.e., nitrogen) 409 which should be positively correlated with photosynthetic rates, 410 was unrelated to adult height. In a closer examination, however, 411 the five highest protein content values, indicative of high 412 photosynthetic rates per unit leaf mass, all belong to species in the gap/edge guild (Table 2). 413

414 Poorter (2007) found that adult leaf traits were more strongly 415 constrained by juvenile light niche than by the adult light niche. In 416 contrast, the only adult leaf trait that differed between the two 417 light guilds in Kibale was fracture toughness. Indeed, this may 418 reflect selection for greater leaf toughness in seedlings of shade-419 tolerant species. Alvarez-Clare and Kitajima (2007) found that 420 seedling leaves have greater fracture toughness for shade-tolerant 421 species in a Panamanian forest. Zanne (2003) studied young 422 seedlings in Kibale and found that percent leaf herbivory was 423 negatively related to cotyledon toughness, which is related to leaf 424 turnover. Furthermore, understory species had lower leaf herbiv-425 ory than open-associated species across different planting 426 habitats, and seedlings growing in forest had lower leaf herbivory 427 than seedlings growing in gaps. In contrast to toughness, chemical 428 defense traits, including phenolics and tannins that are known to 429 be correlated with juvenile light demands elsewhere (Coley, 1983; 430 Kurokawa et al., 2004), were uncorrelated with juvenile light 431 demands or adult height in this study. Although we expected 432 possible relationships between adult leaf characteristics and 433 juvenile functional traits, such relationships may be weak if leaf 434 traits change with size, ontogeny, and light environments that the 435 species experience as adults (Osada et al., 2002; Poorter et al., 436 2005). However, for the Neotropical trees studied by Dominy et al. 437 (2003), adult canopy leaves and understory sapling leaves showed significant correlation for phenolics (r = 0.76, p < 0.0001, N = 28438 439 spp.), but only weak and non-significant correlations for tannins 440 (r = 0.53), proteins (r = 0.43), and fracture toughness of leaf lamina 441 (r = 0.43).

### 442 5. Conclusions

443 In conclusion, post-establishment seedlings of understory and 444 gap/edge guilds in Kibale showed only weak difference in height 445 growth, but not in survival in shade over 10 years. The equally high 446 survival of post-establishment seedlings of species in understory 447 and gap/edge guilds may indicate that their difference in stem 448 distributions along light gradients may be created at an earlier 449 regeneration stage. Alternatively, gaps may be the more forbidding 450 condition, i.e., light guilds may reflect species difference in growth 451 rate and survival in gaps and forest edges, rather than species 452 differences in survival and growth in shade. The tendency of gap/ 453 edge species to grow taller in shade may reflect strong selection for 454 rapid height growth to be competitive in gap/edge environments. 455 Of the functional traits we examined, seed size, adult height, and 456 leaf toughness differed between understory and gap/edge guilds in 457 the expected directions. Of these, only seed size was correlated 458 with seedling height growth in shade (negative correlation). Kibale 459 differed from Asian and Neotropical forests in terms of adult height 460 relationship with juvenile guilds and functional traits, perhaps reflecting differences in historical disturbance regimes and 461 regeneration strategy of dominant trees. Constraints of statistical 462 463 power limited our analysis to contrasts between two broad guilds based on overall stem distribution patterns. Regeneration strate-464 gies clearly need to be considered in a broader ecological context 465 (i.e., beyond understory and gap specialists) and the extent to 466 which ontogenetic shifts influence light niches and coexistence 467 needs to be closely evaluated. 468

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