1 Manipulating resource allocation in plants

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20 ABSTRACT

21	The distribution of nutrients and assimilates in different organs and tissues is in a
22	constant state of flux throughout the growth and development of a plant. At key
23	stages during the life cycle profound changes occur and perhaps one of the most
24	critical of these is during seed filling. By restricting the competition for reserves in
25	Arabidopsis plants the ability to manipulate seed size, seed weight, or seed content
26	has been explored. Removal of secondary inflorescences and lateral branches
27	resulted in a stimulation of elongation of the primary inflorescence and an increase
28	in the distance between siliques. The pruning treatment also led to the
29	development of longer and larger siliques that contained fewer, bigger seeds. This
30	seems to be a consequence of a reduction in the number of ovules that develop
31	and an increase in the fatty acid content of the seeds that mature. The data show
32	that shoot architecture could have a substantial impact on the partitioning of
33	reserves between vegetative and reproductive tissues and could be an important
34	trait for selection in rapid phenotyping screens to optimise crop performance.
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40 **INTRODUCTION**

41	Plants accumulate and redistribute macro and micronutrients throughout their life
42	cycle. Ultimately the plant is seeking to optimise the way it responds to
43	environmental cues and to integrate these with the genetically determined
44	elements of development in order to maximise its reproductive potential and
45	ensure the production and survival of the next generation. Plants, like other living
46	organisms, can be divided into those that have an r- or a K- reproductive strategy
47	(MacArthur and Wilson, 1967; Pianka, 1970). Most weed species adopt an r-
48	strategy and produce many thousands of small seeds with minimum investment of
49	resource per seed; in contrast, many trees invest in relatively few reproductive
50	units per individual but each unit is associated with a substantial quantity of
51	resource to help protect and nurture the subsequent generation (Pianka, 1970). For
52	example, the reproductive propagule of an Arabis weed species is a seed weighing
53	less than 1mg and sunflower seeds are about 100mg each, whereas coconuts
54	typically weigh over 1kg.

55 In order to optimise reproductive potential plants need to (re)mobilise nutrients from sources to sinks in a highly orchestrated way. This process is spatially and 56 temporally dependent on the stage of the plant's life cycle and the reproductive 57 strategy of the plant under consideration. For instance, an annual plant will 58 ultimately die after seed filling is complete; in this case the seeds are the means by 59 which the genetic information of an individual over-winters and survives until 60 growing conditions become favourable once again. In some species seed dormancy 61 can last months or even years and provides a highly effective means of spreading 62

63	the timing of germination and potentially avoiding adverse environmental
64	challenges (Barton, 1961). Work at the Millennium Seed Bank at the Royal Botanic
65	Gardens, Kew revealed that 200 year old seeds, collected by the Dutch explorer Jan
66	Teerlink during a voyage to the Cape of Good Hope in 1803, were still viable. The
67	species included a Leucospermum from the family Proteaceae, and two species
68	from the family Fabaceae including an Acacia and a legume called Liparia villosa,
69	indicating a great range of species for which long-term dormancy is a viable option.
70	For annual plants such as Arabidopsis, under optimal growth conditions, the
71	process of resource allocation ensures that ultimately nearly all the nitrogen
72	resources produced during the photosynthetic period will be remobilised from the
73	vegetative tissues into the developing seeds. In contrast, perennial plants commit a
74	more limited proportion of resources to the reproductive phase of their life cycle as
75	it is advantageous to retain some photosynthetic capacity for as long as the
76	environmental conditions allow. For this reason, it has been speculated that the
77	anthocyanins produced in autumnal leaves provide photoprotection to enable
78	maximal re-absorption during times when the leaves are vulnerable to the effects
79	of high light accompanied by low temperatures (Archetti et al., 2009). For a
80	perennial plant it is less essential to ensure reproductive success every year due to
81	their longer life spans, and tolerance to environmental stress is often achieved by
82	slower growth in addition to niche adaptations such as an evergreen habit. Bulbous
83	plants provide an example where temperature is key to regulating source-sink
84	balance; lower temperatures result in delayed leaf senescence and larger bulbs due
85	to an improved equilibrium between carbon fixation capacity and sink strength
86	(Gandin et al., 2011). Having lived for over 4800 years the bistlecone pine,

Methuselah, is a prime example of a plant's ability to endure abiotic stresses. Grime 87 (1977) proposed that resource-allocation strategies in plants could be divided into 88 89 three categories: Competitive, Stress-tolerant and Ruderal Strategies (denoted C-, 90 S- and R-strategies respectively). Grime summarised the consequences of C-, S- and R-strategies on resource allocation for plant growing in three broad habitat 91 92 categories (Table 1) and it is interesting to review these in the light of considering 93 crop plants and the habitats under which they are grown in a modern agricultural environment. 94

95 **RESOURCE ALLOCATION STRATEGIES OF EXISTING CROP PLANTS**

96 The major cereal crops have undergone many centuries of domestication and, in 97 the developed world, are typically grown in environments where water and nutrients are plentiful. Selection by 'man' has promoted C-strategy plants that 98 99 compete well in a monoculture and make large changes in root/shoot biomass as a consequence of stress, meaning that when these crops are grown in environments 100 101 experiencing prolonged stress they survive less well (see Table 1). This may be one 102 explanation why the major grain crops of the world do not yield well in countries 103 which habitually experience high levels of abiotic (or biotic) stresses, unlike plants 104 native toarid environments which can respond quickly to resource pulses. For 105 instance, high temperatures and low rainfall across southern Australia in 1982, 1994 and 2004 restricted the total wheat yield to less than 10 million tonnes, 106 107 whereas the favourable environment in the 1983/4 season that followed the major drought in 1982 produced in excess of 22 million tonnes (Sutton, 2009). In contrast, 108 109 more recently domesticated crop plants, such as *Brassica* species, retain the weedy

110	characteristics of their wild progenitors and continue to adopt an R-strategy. In
111	consequence, these plants produce seed very rapidly, especially when experiencing
112	abiotic stress (Sinniah et al., 1998), and ensure that the genepool is carried
113	forwards to the next generation. In these situations it would appear that some seed
114	can be produced with such speed that very little resource reallocation is required;
115	this is perhaps not surprising since the seed in a Brassicaceous plant is itself
116	photosynthetic in the initial stages of its development, but it is surrounded by a
117	photosynthetic pod that can supply photosynthates at a highly localised site (Hua at
118	al., 2011). However, if <i>Brassica</i> crops are grown under conditions of minimal abiotic
119	stress they undergo a normal pattern of leaf and pod senescence that suggests
120	resource reallocation does occur to enhance seed number and quality.
121	Observations of Brassica rapa and Brassica napus inflorescences indicates that
122	seed development is frequently terminated following development of the first-
123	formed pods and resumes at the end of flowering, leading to regions of the
124	inflorescence without mature siliques (Figure 1). McGregor (1981) also reported
125	high levels of pod abortion in <i>B. campestris</i> and <i>B. napus</i> which was attributed to
126	over production of flowers and pod formation sites, so that a plant retains spare
127	pods which can fully develop should any become damaged, indicating that pod
128	abortion and the development of excess pod formation sites is normal and not just
129	a response to injury or biotic/abiotic stress. The ability to selectively abort pods
130	during periods of high abiotic stress, and resume once the stress has diminished,
131	would enable plants to become S-strategists. Bosac et al. (1994) and Stewart et al.
132	(1996) both found that exposing the racemes of <i>B. napus</i> or <i>B. campestris</i> to ozone
133	caused pod abortion, pod abscission or fewer seeds per pod. Black et al. (2000)

made the same observation and raised the possibility that for a wild plant this was
potentially disadvantageous; if seed size is too low then plants may lose their
competitive advantage against those growing nearby.

137 Crop breeding programmes still largely focus on increasing yield, particularly in the 138 light of the goal to feed an ever-increasing global population. Most crops originate 139 from R-strategy plants, yet breeding programmes are frequently orientated 140 towards C-strategy plants. In those countries where land use for crops is marginal 141 and abiotic stress levels are high it might be more desirable to select for S-strategy plants. Although S-strategy plants can be overgrown by competitors in a wild 142 habitat, in farmed land this is manageable through weed control or less dense seed 143 144 planting rates. A crop ideotype would be an S-strategy plant with the ability of an Rstrategy plant to reproduce rapidly when the environmental conditions become 145 harsh, thus ensuring at least a minimum yield each year. 146

147 MANIPULATING SOURCE: SINK RATIOS

Using the R-strategy plant Arabidopsis as a model to determine how the 148 149 manipulation of resource allocation can impact on seed quality and yield it has been 150 possible to explore how a crop ideotype might be optimised and developed in a breeding programme. Plants were grown under near optimal conditions in order to 151 maximise seed number. The objective was to ascertain how much plasticity in 152 153 resource allocation was retained by a plant and whether there was potential to 154 increase seed size, yield and quality to make an R-strategy plant invest additional resources into seeds. If the mechanism of resource allocation can be better 155 156 understood then it can be manipulated in crop plants that still harbour weedy traits

157	from their ancestors in order to produce improved varieties. Such varieties might
158	be 'tailored' to suit differing environmental conditions. Selective 'pruning' of parts
159	of the inflorescence was carried out to alter source-sink relationships across the
160	plant. Control plants were compared to those that had either secondary
161	inflorescences removed, or all secondary inflorescences and lateral branches from
162	the main inflorescence removed, thus representing a decrease in sink strength as
163	the pruning treatments became more severe (see Fig. 2A).
164	Analyses of the growth morphology of the plants revealed that the main
165	inflorescence became significantly elongated as pruning became more severe, but
166	that the number of pods on the main stem remained the same as a consequence of
167	larger spaces between the pod formation sites (Fig. 2B, C). The rationale for
168	stimulating inflorescence elongation is unclear, as any competition for light would
169	have been reduced not enhanced by the pruning treatment. Whether this change
170	resulted from an alteration in cell number or size has yet to be determined,
171	however Dale (1959) also observed stem elongation in disbudded cotton plants.
172	Apical dominance was mechanically reinforced by the selective removal of lateral
173	branches and secondary stems and this seems to have resulted in the main
174	inflorescence becoming even more dominant, but the advantage of this is not clear.
175	An alternative explanation is that without the drain of lateral branches on
176	photosynthate reserves the pruned plants were able to implement a more extreme
177	shade avoidance response and elongate more than their highly branched
178	competitors in order to capture the available light more effectively. Rosette leaf
179	number and biomass increased significantly with the pruning of lateral and

secondary branches (sinks), but the rate of senescence of individual leaves was the 180 same as the control plants. This finding is in agreement with that of Nooden and 181 182 Penney (2001) who found that stem removal increased the longevity of the plant 183 but not of individual rosette leaves, enabling the rosette to remain a source of photosynthates and other metabolites for much longer (Bennett et al., 2011; Fig. 184 185 2D). In common with findings in soybean (Seddigh and Jolliff, 1986), it would appear 186 that the strength of the sink is the dominant factor in determining resource reallocation from rosette leaves into the pods. Without the pull from multiple 187 188 branches containing developing seeds, the rosette continues to develop without 189 overall senescence in the absence of other environmental cues such as day length 190 or temperature change. Other experiments in soybean have shown that in the 191 absence of sufficient sink strength the stomata close, leading to a reduction in 192 photosynthesis, although the consequence of this on the distribution of existing 193 photoassimilates and the timing of senescence was not established (Setter and 194 Brun, 1980). However, some authors contest the theory that reproductive 195 development is intrinsically linked to leaf senescence in Arabidopsis. Hensel et al. 196 (1993) reported that the reproductive organs had no influence on leaf senescence 197 and rosette leaf age was the only driver of senescence; Nooden and Penny (2001) 198 took the idea a step further and argue that loss of meristem activity represents the 199 primary signal for whole plant senescence. Early research on source-sink 200 relationships in plants developed the 'self-destruct hypothesis' (Sinclair and De Wit, 201 1976) which assumed that increased nutrient remobilisation out of the leaves was 202 the driver for early senescence. Later researchers in the field found that the rate of 203 photosynthesis in leaves was reduced when sink strength was reduced

204	(Wittenbach, 1983; Crafts-Brander, 1984) and Seddigh and Jolliff (1986) proposed
205	that increasing sink strength would drive increased production of carbon and
206	nitrogen by the leaves, thus switching the emphasis back in line with our current
207	thinking that sink strength 'pulls' resources from the rosette (Bennett et al., 2011).
208	Wild type Arabidopsis plants have been shown to have sub-maximal levels of
209	nitrogen remobilisation from the leaves into developing seeds, but if over-
210	expression of cytosolic pyruvate, orthophosphate dikinase (PPDK) is engineered
211	then nitrogen is exported more rapidly from senescing leaves and seeds on the
212	transgenic lines were larger than wild type (Taylor et al., 2010). This work
213	demonstrates that rosette nitrogen is present in excess of that demanded by the
214	sink (pods) but it is only metabolised in proportion to the pull from the developing
215	pods. The PPDK over-expressing transgenics provide evidence that it is possible to
216	alter the dynamics of remobilisation and source-sink relationships and the capacity
217	for additional resource accumulation is present in the seeds, even in a species such
218	as Arabidopsis where lipid is the primary form of storage metabolite.

219 **RESOURCE ALLOCATION TO THE PODS CAN BE ALTERED**

Reduction in sink strength from pruning lateral and secondary branches altered pod morphology compared to the control. Resultant pods were longer and with a bigger area (Fig. 2E), however, the total number of seeds within each pod was significantly reduced, yet the total weight of seeds in the most extreme manipulation was significantly heavier (Fig. 2F). These observations indicate that when the number of reproductive sites was reduced the number of seeds that even start to develop within a pod is less than the total that could be theoretically sustained within a pod,

as opposed to lots of seeds being formed and then some aborted. These data 227 suggest that the sink strength is determined by the number of pods rather than the 228 229 number of seeds. For a weedy species such as Arabidopsis there is a logic to the 230 argument that the minimum unit that a plant 'measures' is a pod rather than a seed. Since pod shatter will release all the seeds within that pod simultaneously it 231 232 makes ecological sense that the rate of development of all seeds within a single pod 233 is highly coordinated. If the number of pods is too small to pull sufficient resources from the rosette then the co-ordination sequence could be reprogrammed to 234 235 enable a few large, viable seeds to develop rather than lots of small seeds with 236 impaired viability. The mechanism by which a pod is able to 'measure' the amount 237 of resource remains unexplained but our observations indicate that if sink strength 238 is low then not all the potential sites of attachment in a pod are used to develop 239 seeds. As a consequence low numbers of seeds develop per pod to guarantee that all the seeds formed will meet a minimum threshold of resources required for 240 241 viability. One hypothesis is that R-Strategy annual plants operate a 'minimum' 242 viability threshold for reproductive success' and ensure that some seeds will be 243 produced with the best chance of survival. If conditions remain favourable and 244 further resources are available then the plant will invest in more units of 245 reproduction (pods), thus increasing sink strength and mobilising resources more 246 fully from the rosette. This is evident when pods from the main inflorescence of un-247 pruned plants are compared to those from the lateral branches. The pods on the 248 lower section of the main inflorescence are the first formed and contain the 249 heaviest seeds, whereas the later formed pods on lateral branches contain lighter 250 and significantly fewer seeds (Fig. 3). Seeds from pods on lateral branches have high

viability, but further detailed experiments are necessary to establish if there is a
fitness cost if these smaller seeds are selected over multiple generations. Other
researchers have raised the possibility that an R-strategy plant such as Arabidopsis
makes a late, or even continuous, decision about the volume of seed set based on
the available nutrients during the reproductive stage, rather than it being
predetermined by growth and development during the vegetative phase (Schulze et
al., 1994).

258 The observation that seed size within a pod can be manipulated by source/sink 259 manipulation raises the question whether this is the consequence of altered 260 partitioning of resource allocation into each seed or pod. Seeds accumulate a 261 mixture of lipids, proteins and carbohydrates as they develop. In Arabidopsis the 262 protein and carbohydrate are mostly in the embryo, whereas the endosperm contains the majority of the lipid component. The experiments reported here have 263 revealed that enhancing the source strength through selective stem removal causes 264 265 a trend towards an increase in the fatty acid content of the seeds, although protein content did not alter (Fig. 4A, B). This suggests that the developing embryo is of a 266 fixed size, but the endosperm component can increase beyond a minimum 267 268 threshold level. Partitioning of individual fatty acids did not change, indicating that 269 the ratio is genetically determined and therefore fixed. In Arabidopsis the dominant fatty acids are linoleic acid, eicosenoic acid and linolenic acid which are omega 6, 270 omega 9 and omega 3 unsaturated fatty acids respectively. These were found in 271 272 similar proportions to that established by other workers (Katavic et al., 1995; 273 Penfield et al., 2004), although less oleic acid (omega 9) was detected than both of

these groups who found comparable levels in both the embryo and endosperm ofArabidopsis seeds.

276 IMPLICATIONS FOR CROP PRODUCTION

An analysis of all the different parameters we have measured in relation to 277 resource allocation in Arabidopsis has revealed that seed and pod number per plant 278 is more closely related to protein and lipid content than seed mass or pod area (Fig. 279 280 5). This analysis suggests that if the target is to increase the number of viable seeds per pod the consequence should be further mobilisation of lipid and protein 281 282 content into each individual pod which will result in a redistribution of these resources amongst a greater number of seeds within those pods. However, the 283 overall mass of seeds per pod is unlikely to be increased so the usefulness of 284 increasing seed number is limited to production scenarios where the aim is to 285 286 increase the number of seeds that can give rise to viable plants e.g. seed production for the leafy vegetable market. If the target is to increase seed size and nutrient 287 288 content *per se* then it is necessary to alter the source-sink relationship within the 289 plant and to start to shift the architecture away from that of an obligate R-strategy 290 plant and towards that of an S- or C-strategy plant.

There are many reasons why the typical architecture observed in an R-strategy plant that has been adapted for crop production is less than optimal. Most ruderal plants either have a highly branched habit, if they are dicotyledonous, with minimal coordination of pod development and shattering between branches or, in the case of monocotyledonous plants, the natural architecture is to produce numerous tillers with a consequential separation of heading dates and maturation rates on

297 different tillers. In a few examples (e.g. wheat, barley, rice) long-term selection of domesticated crops has enabled much closer development of seed heads present 298 299 on different tillers, but in years when the environment is less favourable, for 300 example when late season rain affects a wheat crop, new tillers are produced that 301 mature several weeks after the main inflorescence. In crops that have received far 302 less attention from plant breeders, such as the numerous small grain crops grown in Africa and Asia, the coordination between flowering heads is extremely poor, 303 304 leading to a considerable reduction in the proportion of potential yield from each 305 plant that is actually harvested. For these reasons it would be desirable to replicate the architecture enforced by our selective stem removal experiments, but it is 306 307 important to appreciate the potential consequences of the change for seed yield 308 and quality.

It is well known that adjusting the seed rate (planting density) for numerous crops 309 310 can have a large influence on plant morphology. Plants grown closely together will 311 reduce the number of branches they produce and adopt a shade avoidance strategy of elongating the main stem to capture the maximum amount of available light 312 (Robson et al., 1996). Whilst excessive stem height can lead to problems with 313 314 lodging there are ways of preventing this through the introgression of genes that lead to dwarfing, such as *Rht* in wheat, and the resultant plants are then able to 315 concentrate their resource allocation into the seed head or pods on the primary 316 inflorescence. Since non-branched plants are grown in a much more densely 317 planted stand the actual yield per m² may not alter significantly, but the 318 319 coordination will be greatly improved by achieving a monoculture of primary

inflorescences from any crop, meaning more of the seeds formed will actually be atthe optimal harvest stage simultaneously.

322 What has become clear from the above findings and from existing literature is that increasing the nutrient supply to the plants during the vegetative stage does not 323 324 guarantee an increase in yield and/or seed quality. Masclaux-Daubresse and 325 Chardon (2011) showed that nitrogen limitation negatively affected plant biomass, 326 yield, and harvest index but did not change the dry weight of individual seeds. 327 There is evidence that the leaves produce photosynthates to an excess of those 328 required by the developing pods and seeds and that there is a positive feedback 329 loop through which the pods (in the case of a Brassicaceous plant) pull the 330 resources they need from the rosette. Whilst supplying adequate nutrition to the leaves and roots will avoid morphological constraints imposed by a starvation 331 stress, more nutrition is not necessarily a guarantee of greater yield (Allen & 332 Morgan, 1972) and a restricted nutrient supply does not necessarily mean that 333 334 senescence will occur more rapidly; Abdallah et al. (2011) demonstrated that short or long-term sulphur limitation delays senescence and allows the plant to 335 remobilise much more sulphur and nitrogen out of its leaves. Agronomic 336 337 consideration should be given to treatments that would make plants increase the 338 rate of resource remobilisation from the source to the sink, for example increasing the expression of PPDK (Taylor et al., 2010) or Arabidopsis thaliana NITRATE 339 TRANSPORTER2.1 (ATNRT2.1; Chopin et al., 2007), or ways to adjust the feedback 340 341 mechanism between source and sink and how it changes through different stages of development. Since these are rather subtle changes it is likely to fall to plant 342

breeders to target suitable genes that will change developmental patterns of
resource allocation and tip the delicate balance of source sink relationships in the
appropriate direction at different stages of crop growth.

346 **PLANT BREEDING TARGETS FOR ALTERED ARCHITECTURE AND RESOURCE**

347 ALLOCATION

348 The control of branch formation in plants has become much better characterised 349 over recent years. Shoot and root branching is in part mediated by the MAX family 350 of genes that function through regulating the strigolactone pathway. Recessive 351 mutants of MAX2 exhibit excessive branching (Stirnberg et al., 2002) and MAX2 encodes an F-Box leucine rich repeat protein that modulates strigolactone 352 signalling, reportedly by targeting the protein for ubiquitination and degradation by 353 the 26S proteasome. Strigolactones appear to act downstream of auxin, and the 354 355 two plant growth regulators interact so that auxin is able to mediate apical 356 dominance by inducing the expression of strigolactone biosynthesis genes (Brewer 357 et al., 2009). MAX2 was originally isolated as ORE9, a gene involved in promoting 358 leaf senescence and thereby potentially increasing the source of metabolites 359 available for seed production (Woo et al., 2001). It is interesting to note that 360 max2/ore9 mutants exhibit increased branching but show delayed plant 361 senescence. In contrast, silencing of the BRANCHED1b gene resulted in increased 362 branching in tomato, but no notable change in fruit maturation or seed 363 development (Martin-Trillo et al., 2001). BRC1b, along with FINE CULM1 are believed to be downstream of strigolactone synthesis and presumably downstream 364 365 of resource allocation during senescence, since fruit development and ripening are

not altered in the plants. Clearly the consequences of this genetic manipulation 366 differs from that brought about by pruning treatments and it would be interesting 367 368 to determine if the mutations above resulted in the production of more or heavier 369 seeds. Work by Abreu and Munné-Bosch (2009) with the NahG transgenics and sid2 370 mutations in Arabidopsis, both of which reduce salicylic acid synthesis, showed that 371 seed production was increased by 4.4-fold and 3.5-fold respectively, compared to 372 the wild type. Salicylic acid deficiency also resulted in higher seed nitrogen content 373 and increased pro-vitamin A and vitamin E than in wild type plants. NahG and sid2 374 both exhibited increased branching per plant, along with reduced seed weight per 100 seeds, increased seed number per fruit and per plant. These results support 375 376 those we found with manipulation of architecture by pruning; in both cases the 377 least branched phenotype has the highest seed weight and the fewest seeds per 378 pod.

379 Other mutants that have already been characterised have the potential to be 380 utilised in the development of crops with altered resource allocation, many of which are also involved in plant growth regulator signalling (Quirino et al., 2000). 381 Improved resource allocation, and consequentially yield enhancement, was 382 383 achieved in rice by reducing the expression of the ethylene receptor ETR2, which 384 caused a 4% increase in thousand grain weight by adjusting starch acclimation and increasing sugar translocation into the grains (Wuriyanghan et al., 2009). It is well 385 known that several mutants in ethylene signalling pathways result in delayed 386 387 senescence of the leaves in dicotyledonous plants (Grbic and Bleecker, 1995) and 388 this may have the potential to alter seed filling at a later stage in development. Ma

and Wang (2003) identified the ETR1 homologue in wheat and initial experiments 389 suggest that it is involved in integrating signals from other hormones, such as 390 391 jasmonic acid, to regulate leaf senescence. Experiments that modulated cytokinin 392 signalling via a SAG12 feedback loop have shown that leaf senescence can be significantly delayed if cytokinin content in the leaves is kept high (Gan and 393 394 Amasino, 1995) and that this can enhance yield, although the impact of this 395 manipulation has been shown to be dependent on nutrient availability and under limited N supply the transgenics had significantly lower fruit dry weight (Wingler et 396 397 al., 2005). Using this strategy to specifically increase cytokinin content in the roots 398 of tomato plants has been reported to increase fruit yield by 30% (Ghanem et al., 399 2011). Cytokinins also appear important in modulating stress responses; when IPT 400 was placed under the regulation of a stress-responsive SARK promoter in rice 401 source/sink modifications led to improved drought tolerance and increased grain 402 yield under water-stress (Peleg et al., 2011). As a staple food crop, rice architecture 403 has been extensively researched and a recently identified point mutation in 404 OsSPL14 was reported to result in the "ideal" plant architecture by producing a 405 taller plant with fewer tillers, denser panicles and enhanced grain yield per panicle 406 (Jiao et al., 2010). The identification of such alleles, which not only affect rice 407 architecture but also yield, may provide targets for future breeding programs.

408 IMPLICATIONS FOR CROP DEVELOPMENT

The example given in the section above is a prime illustration of how below-ground
events influence the development of above-ground development and architecture.
Soil compaction can have serious consequences for the development of above-

ground biomass. Dry mass in barley was reduced by over 50% when soil bulk
density was increased from 1.1 to 1.7 g.cm⁻³ (Mulholland et al., 1996) with a
concomitant increase in xylem abscisic acid content in the compacted samples.
Some attempts have been made to relate soil compaction in different soils types to
yield in cereals (Hamza and Anderson, 2005), but without the ability to directly
observe below ground root architecture it will be difficult to assess how resource
allocation is related to soil structure across the whole plant.

419 The need for rapid phenotyping methods for plants growing *in vivo* has led to the 420 advent of techniques such as microscale X-ray Computed Tomography (microCT) 421 which enables root systems to be investigated in sufficient detail to provide 422 information about architecture as well as biomass (Lucas et al., 2011). Other phenomics centres are being established to investigate above-ground architecture 423 (Berger et al., 2010). Together these techniques will provide a better understanding 424 of how overall plant architecture is regulated, the relationship of architecture to 425 426 crop yield, an understanding of the variability that exists even between plants grown in a monoculture, and therefore how best precision agriculture can be 427 exploited to optimise plant growth and seed/fruit development. 428

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TABLE

Table 1. Morphogenetic responses to desiccation, shading, or mineral nutrient
stress of competitive, stress tolerant, and ruderal plants and their ecological
consequences of three types of habitat (Reproduced from Grime, 1977).

Strategy	Response to stress	Habitat 1 ª	Habitat 2 ^b	Habitat 3 ^c
Competitive	Large and rapid changes in root:shoot ratio, leaf area and root surface area	Tendency to sustain high rates of uptake of water and mineral nutrients to maintain dry matter production under stress and to succeed in competition	Tendency to exhaust reserves of water and/or mineral nutrients both in rhizoshere and within the plant; etiolation in response to shade increases susceptibility to fungal attack	Failure rapidly to produce seeds reduces chance of rehabilitation after disturbance
Stress tolerant	Changes in morphology slow and often small in magnitude	Overgrown by competitors	Conservative utilisation of water, mineral nutrients and photosynthate allows survival over long periods in which little dry matter production is possible	
Ruderal plants	Rapid curtailment of vegetative growth and diversion of resources into seed production	-	Chronically low seed production fails to compensate for high rate of mortality	Rapid production of seeds ensures rehabilitation after disturbance

^a In the early stages of productive, undisturbed habitats (stresses mainly plant induced) and coinciding with competition.

^b In other continuously unproductive habitats (stresses more or less constant and due to unfavourable climate and/or soil) or in the late stages of succession to productive habitats.

^c In severely disturbed, potentially productive habitats (stresses either a prelude to disturbance e.g. moisture stress preceding plant fatalities or plant induced), between period of disturbance.

FIGURE LEGENDS

Figure 1. *Brassica napus* floral inflorescence showing regions of pod abortion.

Figure 2. Morphological influences on resource allocation in Arabidopsis. (A) The three selective stem removal treatments; from left to right: control, secondary branches removed, secondary and lateral branches removed (B) Height of the main inflorescence from each treatment (C) Distance between the pods on the primary inflorescence (D) Images of the rosettes at the end of flowering (E) Pod area and length for each treatment (F) Seed number and weight per pod. N=10 unless otherwise stated on graph. * indicates significant differences between the control and treatments (P <0.05; determined by ANOVA followed by Tukey's post-hoc analysis). Plants were grown under controlled conditions with a 16h light, 8h dark cycle. Temperature was maintained throughout at 20°C and 55% RH.

Figure 3. Comparison of thousand grain weight and the number of seeds per pod between the main inflorescence and lateral branches in un-manipulated plants. N=10. * indicates significant differences between the control and treatments (P <0.05; determined by ANOVA followed by Tukey's post-hoc analysis)

Figure 4. Resources allocated to the seeds. (A) Seed protein concentration from each treatment. Total protein quantification was performed using the Thermo Scientific Pierce BCA protein assay kit according to the manufacturer's protocol. The microtitre plate was read at an absorbance of 540nm. (B) Seed fatty acid profile to show amounts and proportional quantification of seeds from each treatment. N= 10 replicates per treatment. No significant differences were detected but the trend was for an increase in fatty acid content as the manipulations become more severe.

Seeds were freeze dried and the fatty acids measured via direct transmethylation (Smooker et al., 2011). Pentadecanoic acid and methyl heptadecanoate were used as internal standards, the former being present during the transmethylation process whilst the later was added just before the samples were run on the GC. 1µl aliquots of the heptane phase containing the fatty acid methyl esters (FAMEs) were analysed by gas chromatography with flame ionization detection (Agilent G1530a) using a CP-Sil 88 column (50 m length x 0.25 mm id., 0.25 µm film thickness; Chrompack). The GC conditions were: split mode injector (50:1), flame ionizer detector temperature 260°C, oven temperature 130°C for 3 min and increasing at 10°C/min for 6 min: total analysis time 20 min. FAMEs were identified by comparison to the Supelco FAME mix (Sigma-Aldrich).

Figure 5. Dendrogram of resource allocation variables generated by cluster analysis illustrating the relationship between plant architecture and the partitioning of resources.

Figure 1.









Figure 4.







□No Manipulation ■ Main Stem □ Primary Inflorescence



