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Glucosinolates, myrosinase hydrolysis products and flavonols found in rocket (*Eruca sativa* and *Diplotaxis tenuifolia*)

Luke Bell^{†*} & Carol Wagstaff[†]

[†]Department of Food & Nutritional Sciences and the Centre for Food Security, University of Reading, Whiteknights, Reading, Berkshire, UK. RG6 6AH

*Tel: +44(0) 7909447654

*Email: luke.bell@pgr.reading.ac.uk

Abstract

Rocket species have been shown to have very high concentrations of glucosinolates and flavonols, which have numerous positive health benefits with regular consumption. In this review we highlight how breeders and processors of rocket species can utilize genomic and phytochemical research to improve varieties and enhance the nutritive benefits to consumers. Plant breeders are increasingly looking to new technologies such as HPLC, UPLC, LC-MS and GC-MS to screen populations for their phytochemical content to inform plant selections. Here we collate the research that has been conducted to-date in rocket, and summarise all glucosinolate and flavonol compounds identified in the species. We emphasize the importance of the broad screening of populations for phytochemicals and myrosinase degradation products, as well as unique traits that may be found in underutilized gene bank resources. We also stress that collaboration with industrial partners is becoming essential for long-term plant breeding goals through research.

Key words: *Brassicaceae*, Isothiocyanates, Plant breeding, Indoles, Nitriles

26 **Introduction**

27 In recent years, several species of minor leafy-crops have risen to prominence as potentially
28 important commercial and edible species. One example is rocket, which has quickly gained
29 popularity in the Western diet. Originally found as an obscure crop in Mediterranean and
30 Middle-Eastern countries, rocket has become popular largely due to the pungent aromas and
31 tastes associated with it. Glucosinolates (GSLs)/isothiocyanates (ITCs) and flavonols derived
32 from many species ¹⁻⁴ have been shown to infer significant protection against cancer and
33 heart disease ⁴⁻¹¹. In Western countries, diets are generally lacking in fruits and vegetables.
34 Despite government initiatives (such as the “5-a-day” campaign in the UK and USA), these
35 diseases are increasingly leading to premature deaths ¹². Plant breeders aim to maximize
36 levels of such beneficial compounds, but with little genomic information about rocket species
37 presently available, this is a formidable task. This review will give an overview of research in
38 rocket, an outbreeding crop, and how breeders and processors can utilize it to enhance
39 beneficial compounds.

40 **Rocket species**

41 Rocket (also known as arugula, rucola and roquette) is a leafy vegetable crop that has gained
42 substantial popularity across the world, particularly over the last fifteen years ¹³⁻¹⁶. Two main
43 species are predominantly farmed as salad crops; these are *Eruca sativa* (‘salad’ or ‘cultivated’
44 rocket; sometimes referred to as *Eruca vesicaria* subsp. *sativa*) and *Diplotaxis tenuifolia* (‘wild’
45 rocket). Both species share a peppery taste and aroma that is very distinctive ¹⁷. They have
46 been reported to contain high levels of vitamin C, GSLs, flavonols and phenolics ¹⁸⁻²⁵. These
47 are all known to have both anti-oxidant and anti-cancer properties, and are also implicated in
48 lowering the risk of cardiovascular and cognitive disease. For excellent information on these
49 beneficial effects and their underlying causes, see Drewnowski & Gomez-Carneros ²⁶, Keum

et.al ²⁷, D'Antuono et.al ²⁸, Egea-Gilbert et.al ²⁹, Degl'Innoocenti et.al ³⁰, Bjorkman et.al ³¹ and Jeffery et.al ³².

Taxonomy and domestication

A distinction should be made that both *Eruca* and *Diplotaxis* species have overlapping characteristics, and that one can be easily mistaken for the other by the untrained eye, and/or before a certain level of maturity has been reached ²⁸. It is also arguable that *D. tenuifolia* is the least 'wild' of the two species even though the common name is 'wild rocket'. It is featured and favored in commercial products and breeding programs, and is likely to be more domesticated than *Eruca* species as a result. *Diplotaxis* varieties are generally uniform phenotypically, with *Eruca* varieties being more diverse in this respect ²³. No direct genomic evidence has been presented in the literature to suggest one species is any more or less genetically variable than the other. Variability in GSL data seems to support the hypothesis that *Diplotaxis* species are more 'wild' ³³, though it is not conclusive, as only a relatively small number of cultivars have been tested. This is a point that needs clarification through research and extensive breeding, as neither species can be considered fully domesticated ²⁹. For example, germination rates are variable, reproductive organs are typically small, seedpods shatter and disperse freely (rather than staying on the plant), and physical defenses such as leaf hairs are still present in many commercial varieties ³⁴.

Phytochemicals in *Eruca sativa* and *Diplotaxis tenuifolia*: types and structures

Glucosinolates

GSLs are β -thioglucoside *N*-hydrosulphates that are responsible for the sharp and bitter-tasting flavors found in cruciferous vegetables ^{35,36}. In combination with the enzyme myrosinase (thioglucoside glucohydrolase, EC 3.2.1.147), GSLs are hydrolyzed to create isothiocyanates, nitriles, thiocyanates, epithionitriles, indoles, oxazolidine-2-thiones,

75 cyanopithioalkanes, ascorbigens, goitrogens and epithioalkanes ³⁷⁻⁴⁹; see Figure 1. Many of
76 these hydrolysis products have antibacterial, antifungal and insect repellent effects ⁵⁰⁻⁵⁵. GSLs
77 and ITCs are being increasingly used as 'biofumigants' to suppress soil borne pathogens,
78 nematodes and weeds. Some of the volatile products have the opposite effect of attracting
79 species that can tolerate high GSL concentrations, such as types of ovipositing insect ^{56,57}.

80 The conditions under which hydrolysis of GSLs occurs will affect the respective
81 proportions of the chemicals produced; pH, iron ions, thiol ions, temperature and hydration
82 play a particularly prominent role in this process *in vivo* ⁵⁸. The separation of GSLs in
83 specialist 'S-cells' from myrosinase in myrosin cells means that the two components only
84 come into contact upon tissue disruption; for example when damaged via chewing or
85 digestion ⁵⁹⁻⁶⁹. It is the biological activity of the ITC hydrolysis products in humans that are of
86 most interest in rocket ⁵⁰. GSLs can be hydrolyzed within the intestinal tract by gut microflora
87 that are known to have specific myrosinase activity ⁷⁰⁻⁷³, but the efficacy of their action is not
88 yet well determined.

89 GSL concentrations can vary and change over time depending on environmental
90 conditions and stress ⁷. Other factors affecting GSL profiles include the plant age, organ type,
91 developmental stage, ambient air temperature, level of water stress, photoperiod, agronomic
92 practice, degree of wounding, and geographical origin of the variety/species ⁷⁴⁻⁸¹. These can
93 often affect the profiles of *all* phytonutrients contained within tissue, not just GSLs ⁸², and they
94 are all factors that plant breeders aim to mitigate through development of genetically
95 advanced and uniform breeding lines.

96 GSLs and the ITC derivatives have been an integral part of the human diet for millennia
97 because of the presence of them in the family *Brassicaceae* ^{64-66,83-89}. GSLs are evolutionarily
98 recent secondary metabolic products having arisen 10-15 million years ago ^{90,91}, acting to
99 prevent pathogen attack and dissuade herbivory. They are known in only a few angiosperm

100 families of the order *Brassicales*, which includes the *Brassicaceae* ⁹²⁻¹⁰⁰, and of which *Eruca*
101 and *Diplotaxis* are members.

102 A study by Pasini et al. ³³ of 37 rocket accessions (*Diplotaxis* and *Eruca*) showed that
103 GSL profiles were all very similar, regardless of the species. In total, twelve GSL compounds
104 were found across all accessions; Table 1 illustrates all known GSL compounds identified to-
105 date in rocket. These include 4-mercaptobutyl GSL (glucosativin) ²¹, 4-methylthiobutyl GSL
106 (glucoerucin) ¹⁰¹, and 4-methylsulfinylbutyl GSL (glucoraphanin) ²⁸, which constitute the
107 three most abundant GSLs in rocket.

108 **Flavonols**

109 Flavonols are diphenylpropanes (C₆-C₃-C₆) ¹⁰² and are another important group of chemicals
110 found within rocket species. Flavonols in rocket are found with sugar conjugates, and
111 typically occur in relatively large quantities ¹⁰³. The aglycones found (such as quercetin and
112 kaempferol) are glycosylated and acylated, which in turn affects their biological properties ¹⁸.
113 A study by Martínez-Sánchez et al. ¹⁸ identified over 50 different flavonol compounds across
114 four different species. Watercress, mizuna and two species of rocket were all found to
115 accumulate very different compounds within their leaves, and in varying quantities. Wild
116 rocket showed high levels of quercetin-3,3',4-triglucosyl (43.5 mg per 100g⁻¹ fw) and salad
117 rocket had mostly kaempferol-3,4'-diglucosyl (97.8mg per 100g⁻¹ fw). The group also showed
118 a correlation between quercetin derivatives and high antioxidant activity, despite the
119 significant variations seen between species.

120 Studies conducted on rocket tissues have identified significant concentrations of
121 polyglycosylated flavonols. The core aglycones of these are kaempferol, quercetin and
122 isorhamnetin ¹⁰²; Table 2 provides an up-to-date list of all flavonol compounds identified in
123 rocket to-date. Martinez-Sanchez et al. ¹⁸ showed that *Eruca* species accumulate kaempferol
124 derivatives, whereas *D. tenuifolia* accumulates predominantly quercetin instead, meaning that

125 the two chemicals could be used as an identification marker between the two species ¹⁰⁴.
126 Isorhamnetin aglycones are common to both species but typically in much lower
127 concentrations ³³. The specific aglycones also infer varying degrees of anti-oxidant activity.
128 For example, quercetin derivatives have a higher activity than kaempferol and isorhamnetin.
129 The differences in structure (the arrangement of hydroxyl groups and glycosylation) affect
130 anti-oxidant activity by allowing the molecules to act as hydrogen/electron donors, single
131 oxygen scavengers, or as reducing agents ¹⁰⁵.

132 133 **Phytochemicals and the relation with quality: taste and aroma**

134 It is thought that the presence of glucosativin, glucoerucin and their hydrolysis products
135 within rocket leaves is what gives them a characteristic flavor ⁴⁴. Many of the health beneficial
136 GSLs and ITCs are thought to be responsible for strong tastes that *some* consumers find
137 repellant ¹⁰⁶. It seems that to many people, these compounds contribute very little to a
138 pleasurable eating experience and are actively avoided ⁸³. Conversely however, some people
139 *do* prefer these strong tastes and aromas, and will actively seek to consume rocket when it is
140 available. Growers in Italy often prefer the subsequent cuts because of the more intense tastes
141 and aromas that are produced ¹⁰⁷ and some will even ‘sacrifice’ the first cut in favour of the
142 subsequent leaf growth. This highlights a divide between consumers that may be indicative of
143 underlying genotype(s) for taste perception and preference.

144 The breeding process in rocket varieties to-date has effectively made the species
145 ‘milder’ in taste when compared to plants that grow naturally in the wild. Whether this has
146 been intentional or as a result of selecting for other unrelated traits (such as leaf morphology)
147 is debatable. Some recent commercial varieties have been bred for a ‘hotter’ taste, such as
148 ‘*Wildfire*’, by Tozer Seeds (Surrey, UK).

149 A study by Pasini et al. ¹⁷ demonstrated how breeding for sensory traits could be
150 achieved, by highlighting which glucosinolates contributed to specific taste and aroma
151 elements in rocket. It was found that progoitrin/epiprogoitrin is responsible for bitter taste
152 attributes, despite being only a minor component of the overall GSL profile of rocket (4.3-
153 11.4% of total GSL concentration). The perceived pungency of leaves was positively related to
154 the overall GSL content of accessions, and the levels of glucoraphanin negatively contributed
155 to the typical 'rocket' flavour. The study also highlighted an important difference between
156 rocket and other *Brassica* sensory studies ¹⁰⁸, in that bitterness was perceived as a favorable
157 characteristic according to panelists. The flavonol compound kaempferol-3-(2-sinapoyl-
158 glucoside)-4'-glucoside also significantly and positively contributed to flavor attributes in
159 *Eruca* accessions. This would indicate that GSL compounds are not totally responsible for
160 flavor in rocket. The study itself stopped short of saying how or if the information obtained
161 would be used in breeding programs, but with study into rocket flavor components, milder
162 (and/or stronger) varieties could be bred more efficiently once the responsible compounds
163 are properly identified ²⁶.

164

165 **Health promoting properties of glucosinolate-myrosinase products and flavonols of** 166 **rocket**

167 **Isothiocyanates**

168 ITC hydrolysis products have been identified in rocket ⁴⁵, such as 4-(methylthio)butyl ITC
169 (erucin) ^{109,110} which is known to show anti-proliferative activity in human lung carcinoma
170 A549 cells, hepatoma (HepG2) cells, colon cancer cells, prostate cancer cell lines (PC3, BPH-1
171 and LnCap) and leukemia cells ¹¹¹. Erucin is a structurally reduced analog of sulforaphane,
172 (which is predominantly found in broccoli) and has shown promising anti-cancer properties
173 *in vitro* (e.g. anti-proliferation of human erytroleukemic K562 cells) ¹¹². Research into the

174 chemopreventative and anti-genotoxic nature of ITCs has shown promising results ¹¹³ (see
175 Figure 2). Other studies involving chemically induced genotoxicity have shown very strong
176 anti-genotoxic effects of *E. sativa* extracts ¹³ which is in agreement with other *Brassicaceae*
177 studies ^{114,115}. Identifying specific cultivars of rocket with elevated levels of erucin and
178 glucoraphanin would be an important first-step in developing superior varieties from a
179 human nutrition standpoint.

180 The results of GSL/ITC research prompted an investment in broccoli breeding in the
181 last decade. A similar concerted effort could be made for rocket which contains similar
182 compounds, and which are potentially just as efficacious in humans ¹¹⁶. Erucin for example,
183 has been shown to have very similar, and even superior, biological activity to sulforaphane ¹¹⁷.
184 One paper has specifically demonstrated that the concentrations of rocket ingested in an
185 average daily diet is significant enough to infer a cancer preventative effect ¹³. The
186 metabolism of ITCs in humans via the mercapturic acid pathway has been investigated. ITCs
187 are conjugated with glutathione and degraded by N-acetylation, initiating an increase of phase
188 II detoxification enzymes; see Figure 3 for detailed pathway breakdown of erucin ¹¹³.

189 **Nitriles**

190 Along with ITCs, nitriles are the most abundant bioactive compounds produced by GSL
191 hydrolysis ¹¹⁶. The hydrolysis of glucoraphanin for example, yields predominantly
192 sulforaphane and sulforaphane nitrile. The ratio in which the two are formed depends greatly
193 upon the environmental conditions and the plant cultivar that is used ¹¹⁷. A low pH medium
194 tends towards the formation of nitriles, whereas high pH forms ITCs ^{118,119}. The presence of
195 thiol and iron ions favors nitriles, and high temperature and hydration produce more ITCs
196 ^{120,121}. This can have substantial consequences for any potential health benefits that might be
197 inferred from eating rocket ¹¹⁹. The nitrile form is approximately three orders of magnitude
198 less efficacious than the ITC in inducing quinone reductase (phase II enzyme), and thus infers

199 a reduced enzymatic and anticarcinogenic response. Nitriles also compete with ITCs in this
200 induction, and reduce potential positive effects further ³⁸. As the ratio of these compounds
201 may depend on plant variety, care must be taken in rocket breeding when selecting plants for
202 GSL content, as this may not be reflective of the bioactives produced in subsequent hydrolysis
203 reactions ¹²². Other underlying genetic factors may influence which degradation pathway is
204 taken.

205 **Indoles**

206 Indoles are the predominant autolysis product of indole glucosinolates such as glucobrassicin,
207 as their ITC counterparts are unstable ⁸⁵. Glucobrassicin has been detected as a minor GSL in
208 rocket species ³³, and the predominant indole species produced is indole-3-carbinol. This
209 compound is known to be cancer-preventative ^{125,126}, particularly in reproductive organs *in*
210 *vitro* and *in vivo*. A condensation product of indole-3-carbinol, 3,3'-diindolymethane, is also
211 responsible for beneficial physiological effects. Both compounds have been shown to reduce
212 cell proliferation in breast, prostate, cervical and colon cancer cell lines. They also show
213 distinct differences from ITCs such as sulforaphane ¹²⁷, and inhibition of tumor development
214 in the stomach, breast, uterus, tongue and liver of rodents ¹²⁸⁻¹³⁵. Experiments in rodents have
215 shown an increase in drug-metabolizing enzymes in the stomach, liver and small intestines of
216 individuals consuming both ITCs and indoles. This is suggestive of enhanced detoxification
217 phase II enzymes (such as quinone reductase, glutathione reductase and glutathione
218 transferase) ¹³⁴, and a mechanism by which these phytochemicals infer chemopreventative
219 effects ^{135,136}.

220 Typically indoles inhibit cell proliferation through cytostatic mechanisms, whereas
221 ITCs induce cytotoxicity within cell lines (at above 12.5µM concentrations), which ultimately
222 leads to increased apoptosis ^{137,138}. This indicates that both types of compound could act and
223 be effective at different stages of cancer development ¹¹. Indoles have been shown to induce

224 programmed cell death in prostate, breast and osteocarcinoma cell lines ¹³⁹ and G₁ cell cycle
225 arrest in breast and prostate cancer cell lines ^{142,143}. It is these cytostatic effects on cell
226 proliferation that has been suggested as the mechanism responsible for the lack of apoptosis
227 effects in indoles ¹⁴¹.

228 Using information on GSL content in rocket, the ITC and indole effects can be
229 potentially maximized in new varieties, and be of a greater benefit to human health when
230 considered in tandem, rather than separately ¹²⁷.

231 **Oxazolidine-2-thiones & goitrogens**

232 The hydrolysis of β -hydroxy-alkyl GSL compounds (e.g. progoitrin; a minor GSL in rocket) can
233 produce oxazolidine-2-thiones such as goitrin (5-vinyloxazolidine-2-thione) ¹⁴²⁻¹⁴⁸. It is these
234 compounds that are largely attributed to the thyroid condition of goiter in mammals ¹⁴⁹, but
235 the action of microflora in the gut is thought to mediate the problems associated with high
236 oxazolidine-2-thione intake ^{150,151}. That being said, oxazolidine-2-thiones interfere with
237 thyroxine synthesis ¹⁵⁴ and are therefore likely to have an adverse biological effect regardless
238 of gut microflora action or bodily iodine status ³. A study by Nishie and Daxenbilcher ¹⁵⁵
239 showed that these compounds are not teratogenic or embryotoxic however.

240 These molecules contribute significantly to the bitter taste of rocket that some people
241 perceive quite strongly ¹⁵⁴. The detection of these compounds may be mediated in a similar
242 genetic fashion as PROP (propylthiouracil), for example ^{155,156}. By using phytochemical data in
243 rocket breeding programs these oxazolidine-2-thione components could be reduced,
244 potentially improving consumer acceptance (depending on the target consumer) and avoiding
245 any possible adverse health effects associated with over-consumption.

246 **Ascorbigens**

247 Ascorbigens are formed via the reaction of indole-3-carbinol and 3,3'-diindolymethane with
248 ascorbic acid in the stomach during myrosinase-catalyzed degradation of indoly-3methyl

249 glucosinolates ^{157,158}. In this manner it is thought that ascorbigens have a role in cancer-
250 modulation ¹⁵⁹ via quinone reductase induction ¹¹⁴. As has been highlighted previously, this
251 has important implications for breeding for plant varieties with enhanced chemopreventative
252 effects.

253 **Epithioalkanes**

254 Epithioalkanes are formed as part of the myrosinase reaction with GSLs at low pH with
255 epithiospecifier protein and ferrous ions. These GSLs typically have a side-chain with a double
256 bond, such as sinigrin ^{160,161}. It is uncertain whether these compounds produce any significant
257 bioactive effect in humans, but the ratio in which they are produced alongside ITCs, nitriles
258 and indoles may impact on these compounds' efficacy as anti-carcinogens.

259 **Flavonols**

260 The antioxidant and anti-inflammatory function of flavonols in the human diet are well known
261 and include protecting the colonic epithelium from free radical damage ¹⁶⁴⁻¹⁶⁷. They can
262 induce the up-regulation of enzymes (such as cytochrome P450), that may lead to a decreased
263 risk of cancer, cardiovascular disease, immune dysfunction, atherosclerosis and chronic
264 inflammation ^{168,169}.

266 **Factors affecting phytochemical content**

267 **Breeding and cultivation**

268 Rocket has been consistently shown to be a good dietary source for flavonols, GSLs and anti-
269 oxidants. However, there can be large differences between plants of the same germplasm
270 accession due to a combination of genetic and environmental variability. This is probably due
271 to the outbreeding nature of the species ¹⁰⁴ and a lack of overall uniformity in varieties.
272 Commercial varieties cannot be considered truly domesticated because of this tendency for
273 outcrossing, and the susceptibility of plants to inbreeding depression (a loss of genetic

274 variability due to repeated self-pollination or crossing with a closely related individual).
275 Development of advanced open-pollinating breeding lines (lines that are allowed to cross-
276 pollinate freely in a population of selected individuals), or even F₁ hybrids (superior varieties
277 produced by crossing distinctly different, elite inbred lines), could potentially minimize such
278 variation.

279 Throughout the food chain there are many aspects that can have an adverse effect on
280 GSL levels within leaves (Figure 4). These include the cultivar choice, cultivation practice,
281 climatic conditions, photoperiod, sulphur and nitrogen availability, harvest date, time spent in
282 storage, the temperature of wash water, levels of physical damage to leaves, packaging
283 atmosphere and food preparation methods ^{30-32,170-173}.

284 **Harvesting**

285 Rocket species have the ability to re-grow their leaves repeatedly after cutting, which allows
286 for several harvests to take place under optimal conditions ¹⁰⁷. In parts of southern Italy, it is
287 not unheard of for up to seven harvests to occur from a single planting. This has obvious cost-
288 saving benefits for growers, but multiple harvests also induce stress responses in rocket that
289 may be detrimental to the flavor and aesthetics of the crop. Stress drives up the production of
290 secondary metabolites such as GSLs and anthocyanins, which will produce very strong, bitter
291 tastes. There are other detrimental effects of multiple harvests; leaves become progressively
292 smaller and more 'skeletal' in appearance with each cutting, for example. High anthocyanin
293 levels also affect the color of leaves, turning them an undesirable pink, purple or red. Color
294 has been found to be one of the most important characteristics consumers look for in rocket
295 ¹⁷⁴, and so the loss of fresh appearance can ultimately lead to rejection of crops by
296 supermarkets and processors.

297 **Industrial and culinary processing**

298 There are five main influences that have been identified in affecting GSL levels during
299 processing ⁹⁴. These are the action of myrosinase hydrolysis, myrosinase inactivation, the
300 lysis and leaching of GSLs into wash-water, thermal degradation of GSLs, and the loss of
301 ascorbic acid, iron and other enzyme co-factors. Myrosinase inactivation and thermal
302 degradation of GSLs is probably less of an issue in rocket species, as the leaves are not
303 typically cooked. The leaves are not ordinarily frozen, and so freeze-thaw hydrolysis is not
304 likely to be a major factor either. Other factors almost certainly play a significant role in GSL
305 and phytochemical loss in rocket. Verkerk et al. ⁹⁴ highlighted four key areas that affect GSL
306 levels before reaching the end consumer. These are:

- 307 1. The variety / cultivar used
- 308 2. Storage and packaging (post-harvest, post-processing & in shops/supermarkets)
- 309 3. Industrial processing
- 310 4. Consumer preparation methods

311 If each of these areas can be mitigated through breeding superior varieties, consumers
312 will receive an end product that is of higher nutritive quality and thus provide increased
313 health benefits.

314 **Post harvest storage**

315 Studies on both *Diplotaxis* and *Eruca* species have been conducted to determine the effects of
316 post harvest storage conditions on chlorophyll content and respiration rates ¹⁵. Both species
317 of rocket have been found to have high respiration rates ¹⁰⁷ leading to rapidly impaired visual
318 quality, such as stem browning, tissue yellowing and general decay ¹⁷⁵. Provided initial GSL
319 loss can be mitigated through breeding, ITC formation has been shown to increase over nitrile
320 formation during the storage period ¹⁷⁶.

321 Time, temperature, humidity and atmospheric conditions are all optimized for specific
322 crops within the logistics chain, but these factors are often only designed to prevent visual

323 degradation and not phytochemical breakdown ¹⁰⁰. Getting producers, packagers and
324 transporters to change their current practices in order to better preserve the health-
325 promoting compounds in rocket would be a difficult task. Treatments and storage conditions
326 are often integrated parts of protocols and procedures, and changing these would require
327 significant testing on a commercial scale.

328

329 **New selection tools for breeders**

330 **Phytochemical selection**

331 It should not be forgotten that some GSLs and their breakdown products are thought to be
332 toxic, and even carcinogenic, at high concentrations ¹²⁸. Breeders and researchers should be
333 mindful that more of a certain compound does not necessarily mean ‘better’ ¹⁷⁷. Humans seem
334 to be able to tolerate GSLs much better than pigs, rats and rabbits for example; but
335 overconsumption of these compounds may have serious health consequences ⁶⁴ as high dose-
336 effect relationships are as yet unknown in humans ⁹⁴. Few papers in GSL research (regardless
337 of species) have acknowledged the potential for plant breeders to utilize HPLC/UPLC/LC-
338 MS/GC-MS methods within breeding programs to ‘monitor’ and select plants for their
339 phytochemical content in this manner. These techniques would provide valuable information
340 on breeding lines relatively rapidly, especially for GSL and flavonol breeding ¹⁷⁸. It is not
341 common practice to select rocket plants based on their phytochemical profile at present, but
342 as interest in these compounds increases it will be necessary for breeders to modify their
343 selection criteria and information sources in order to remain competitive in the salad
344 vegetable market ⁹⁴. This has been achieved with ‘*Beneforte*’ broccoli (Seminis Vegetable
345 Seeds; subsidiary of Monsanto Company, St. Louis, Missouri, USA; www.beneforte.com) for
346 example. It has also been indicated in hybrid varieties of *Brassica* that ITC/nitrile ratios can be
347 selected for ¹⁷⁹.

348 **Genetic resources and Marker Assisted Breeding**

349 European initiatives (such as the EU GENRES project ‘Leafy vegetables germplasm,
350 stimulating use’; <http://documents.plant.wur.nl/cgn/pgr/leafyveg/>) have included rocket
351 species within their remit, indicating the rising prominence of the species, and the desire for
352 more work to be conducted on them. The germplasm accessions stored in gene banks are a
353 valuable genetic resource for breeders to take advantage of ¹⁸⁰. The accessions contained
354 within these collections are highly variable and have unique visual and sensory
355 characteristics that could be introgressed into breeding lines relatively easily ¹⁸¹.

356 Genetic information about rocket within the published literature is very scarce. Some
357 molecular marker techniques such as Random Amplification of Polymorphic DNA (RAPD),
358 Inter-Simple Sequence Repeats (ISSR) and Amplified Fragment Length Polymorphisms
359 (AFLP) have been used to analyze morphological characteristics of *Eruca vesicaria* ²⁹. ISSR and
360 AFLP are relatively robust for screening variable populations and discriminating between
361 cultivars ¹⁸⁰ but RAPDs are notoriously unreliable and suffer from a lack of reproducibility
362 and resolution. Perhaps one of the most underutilized marker types is SRAP (Sequence
363 Related Amplified Polymorphism). The forward and reverse primers are designed to target
364 arbitrary GC and AT rich sequences of the genome respectively, and are therefore more likely
365 to anneal to active genomic regions ¹⁸². This could be of use in understudied crops such as
366 rocket, as it provides a simple, repeatable and reliable way of screening large populations.

367 These techniques are now for the most part however, obsolete in advanced molecular
368 plant breeding, as NGS (Next Generation Sequencing) and SNP (Single Nucleotide
369 Polymorphism)/QTL (Quantitative Trait Loci) analyses are far more specific, reliable and
370 cost-effective. SNPs are the most abundant marker type within genomes, and their high
371 density is ideal for studying specific regions in detail ¹⁸³. NGS techniques are now relatively
372 affordable, even for relatively small companies. They are widely available in academic

373 institutions, but many companies are bypassing these in favor of dedicated private
374 commercial services ¹⁸⁴ or are developing their own in-house facilities. The inability of some
375 research institutions to provide adequate customer service, cost-effectiveness, data storage,
376 and results on time is jeopardizing how much knowledge is in the public domain. Increasingly,
377 both large and small breeding companies are collaborating privately and advancing
378 techniques far beyond those found in academic institutions. Future work by institutes in
379 advanced genomics, sequencing and genotyping is likely to be obsolete in some cases because
380 private research is already finding new innovations, e.g. for data storage and bioinformatics.
381 Because private companies have no obligation to share their knowledge, many of these
382 advances may be unobserved by the mainstream scientific community. Institutes and
383 Universities need to do more to attract business from industry in order to keep up with the
384 pace of private advances in this area.

385 Transcriptome sequences are now (generally) adequate for breeders to use and make
386 huge advances in only a few years. Linkage mapping and QTL analyses can be conducted on
387 desktop computers, making integration into breeding companies relatively straightforward
388 from an IT point of view, even if the actual sequencing and genotyping are outsourced. Again,
389 this may typically be to private companies providing a dedicated service. The availability of
390 software licenses and advanced training courses from private companies also means plant
391 breeders do not necessarily need the expertise found in Universities and research institutes in
392 order to attain their goals.

393

394 **Summary**

395 Of all the research papers concerning rocket species and their phytochemistry, none have
396 directly addressed how information could be used within a working breeding population.
397 Often it is explained or postulated purely as theory rather than actual practice, or only given a

398 cursory mention. Only very rarely is a plant breeding program reflective of theory, due to the
399 large number of environmental factors affecting plant growth, development and reproduction.
400 The progressive selection of rocket plants through conventional/molecular breeding would
401 be a valuable tool for the research community as well as providing an excellent incentive for
402 breeding companies to fund research. The actual monitoring and quantification of
403 GSL/flavonol levels through successive generations (i.e. not just one as has been the case with
404 most studies) would not only validate the heritability of such traits in rocket, but would also
405 provide a 'roadmap' for how other minor crops might be developed for commercial use.

406 Attention must be paid to the phytochemical content of varieties within breeding
407 populations of rocket. By focusing solely on morphological traits, important phytochemical
408 genotypes may be inadvertently lost from populations; this could be said of all *Brassicaceae*
409 species, not just rocket. The balance of glucosinolate-myrosinase degradation products does
410 seem to have a genetic component to it and so could be selected for also. Utilising genetic
411 resources, the falling costs of sequencing and bioinformatics can produce nutritively superior
412 varieties of rocket in the near future. Plant breeding typically takes longer than the average
413 research project allows for, even with the use of advanced genomic selection methods. This is
414 a situation that could be remedied by long-term industrial collaboration and sponsorship by
415 plant breeding firms.

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Figure captions

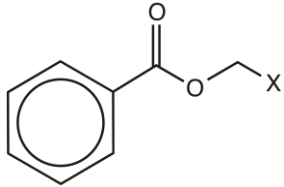
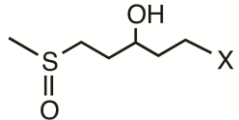
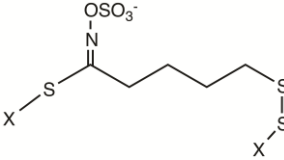
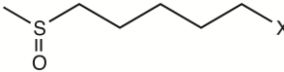
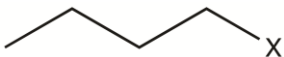
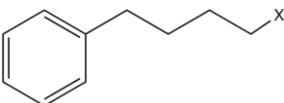
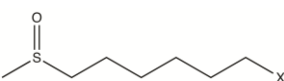
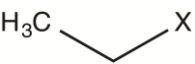
Figure 1: – The glucosinolate-myrosinase reaction and some of the subsequent compounds produced under different conditions, such as pH and the influence of epithiospecifier proteins (ESP) (Adapted from Zhang ⁹ and Hall et al. ¹⁸⁵).

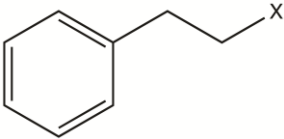
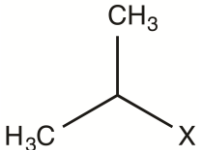
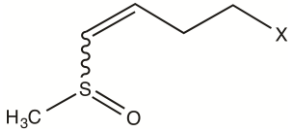
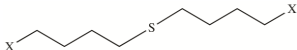
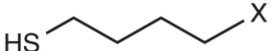
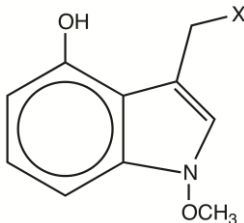
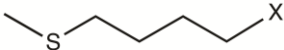
Figure 2: – Pathways of documented ITC action in tumorigenic cells. See Wu et al. ¹¹³ for a detailed review of the roles ITCs play in cancer prevention.

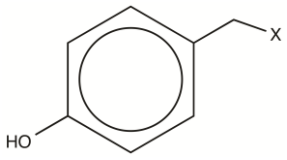
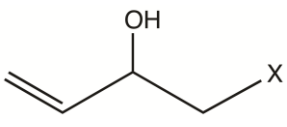
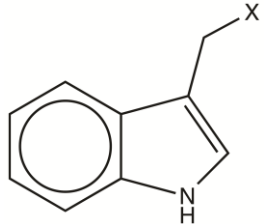
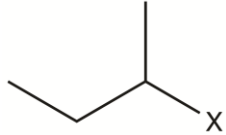
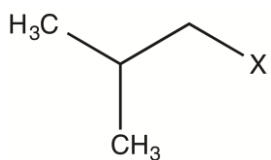
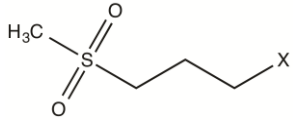
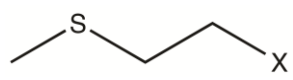
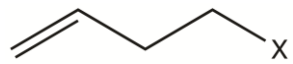
Figure 3: – The mercapturic acid pathway of ITC metabolism in the human body. After ingestion of rocket leaves glucoerucin is hydrolyzed by myrosinase to form erucin. This is released and absorbed in the ileum, where it is transported in the blood to cells around the body. ITCs initiate Phase II detoxification enzymes in this pathway, and are known to aid in cancer prevention. (Adapted from Wu et al. ¹¹³).

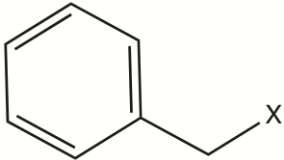
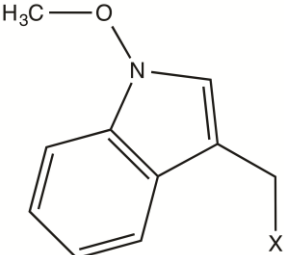
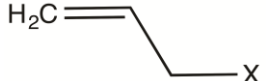
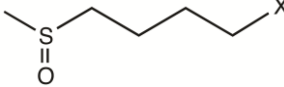
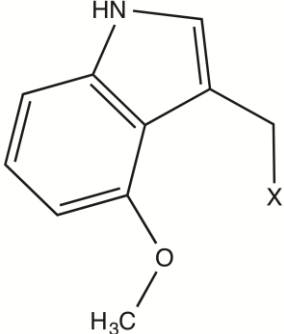
Figure 4: – Factors and conditions within the commercial supply chain that affect GSL and flavonol levels within rocket leaves.

Table 1: – Intact Glucosinolates Identified Within Leaves Of Rocket, *Eruca* and *Diplotaxis* Species, By LC-MS (Negative Ion Mode)

R-group	Common name	R-group structure ^x	Mass parent ion	MS ² spectrum ions (signature ions in bold)	Reference
2-(benzoyloxy) ethyl	-		466	386	33
3-hydroxy-5-(methylsulfinyl) pentyl	-		482	403	
4-(β-D-glucopyranosyldisulfanyl) butyl	Diglucothiobeinin		600	521	33,186
5-(methylsulfinyl) pentyl	Glucoalyssin		450	371	
N-butyl	Dihydrogluconapin		374	294	
4-phenylbutyl	Glucoamoracin		450	371	28
7-(methylsulfinyl) heptyl	Glucoibarin		494	414	
Ethyl	Glucolepiidin		346	266	

2-phenylethyl	Gluconasturtiin		422	343	
1-methylethyl	Glucoputranjivin		360	280	28
4-(methylsulfinyl)-3-butenyl	Glucoraphenin		434	354	
Dimeric 4-mercaptobutyl	DMB		811	731, 569, 405	21,33,186
4-mercaptobutyl	Glucosativin		406	259 , 209, 194, 138 97, 96	
4-hydroxy-3-indolymethyl	4-Hydroxyglucobrassicin		463	383, 285 , 275, 267, 259 , 240	33,178
4-(methylthio) butyl	Glucoerucin		420	340, 291, 275, 259 , 242, 227, 195, 178, 163	

4-hydroxybenzyl	Glucosinalbin		424	344, 291, 275, 261, 259 , 246, 231, 228, 182	33,178
(R,S)-2-hydroxy-3-butenyl	Progoitrin/epiprogoitrin		388	332, 308, 301, 275, 259, 210, 195, 136	
3-indolymethyl	Glucobrassicin		447	275, 259 , 251, 205	33,178,187
1-methylpropyl	Glucocochlearin		374	294	85,186
2-methylbutyl	Glucojiaputin		388	308	
5-(methylsulfonyl) pentyl	Glucoerysienin		466	386	28,33
3-(methylthio) propyl	Glucoiberberin		406	326, 275, 259 , 228, 145	85,178
3-butenyl	Gluconapin		372	292, 275, 259 , 227, 195, 194, 176	28,178

Benzyl	Glucotrapaeolin		408	328, 275, 259 , 241, 230, 212, 195, 166	
1-methoxyindol-3-ylmethyl	Neoglucobrassicin		477	447, 466 , 284, 259	28,178
2-propenyl	Sinigrin		358	278, 275, 259 , 227, 195, 180, 162	
4-(methylsulfinyl) butyl	Glucoraphanin		436	372 , 291, 259 , 97, 96	21,33,178,187
4-methoxy-3-indolymethyl	4-Methoxyglucobrassicin		477	291, 275, 259 , 235, 227, 195	178, 188

^x = standard GSL molecule according to IUPAC nomenclature

Table 2: – List Of Flavonol Compounds Identified In Leaves Of *Eruca* And *Diplotaxis* Species, By LC-MS (Negative Ion Mode).

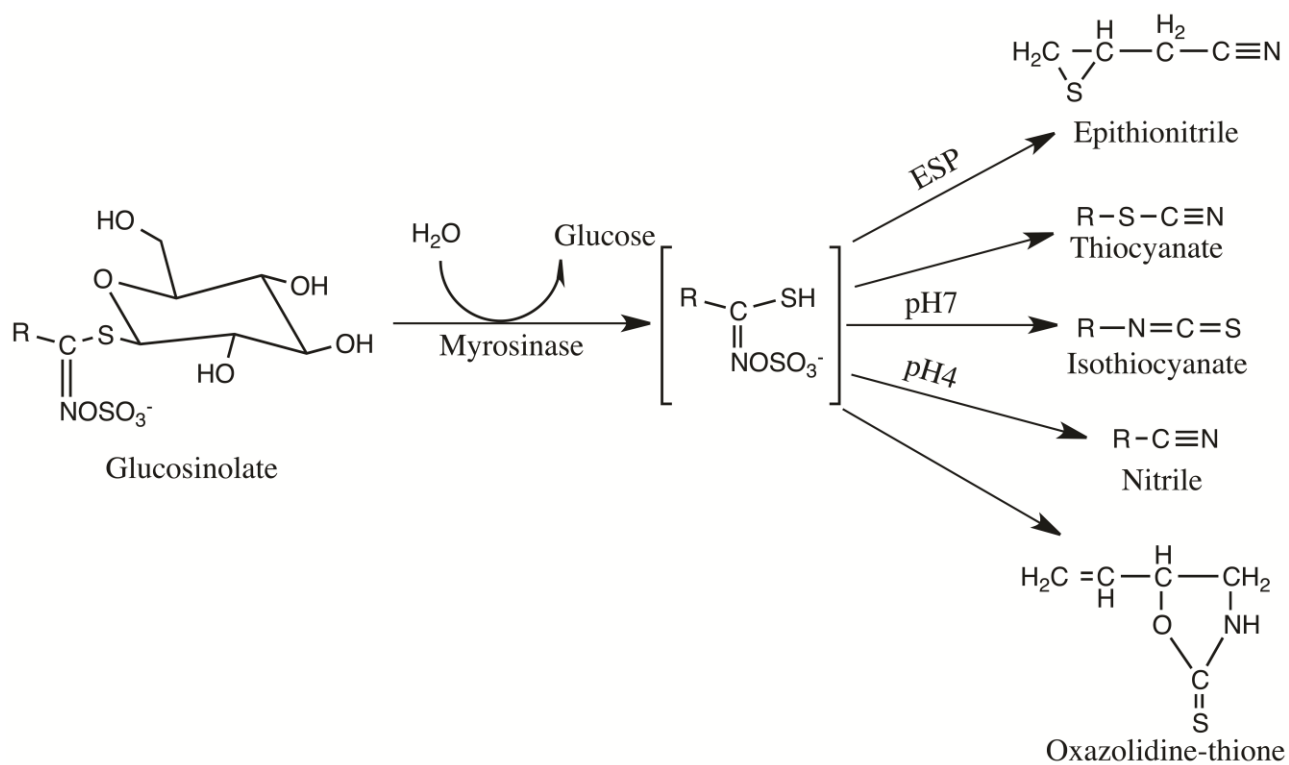
Flavonol compound ^a	<i>Eruca</i> ^p	<i>Diplotaxis</i> ^p	Mass parent ion	MS ² spectrum ions (signature ion in bold)	Reference
I 3,4'-diGlc	✓	✓	639	477	
I 3-Glc	✓		477	-	
K 3-(2-Sinp-Glc)-4'-Glc	✓		817	-	18,33
K 3,4'-diGlc	✓	✓	609	-	
K 3-Glc	✓		447	285	
Q 3-Glc	✓		463	301	
K 3-diGlc-7-Glc	✓		771	609	
K 3-Sinp-triGlc-7-Glc	✓		1139	977, 771, 609, 429	33
Q 3,4'-diGlc-3'-(6-Caf-Glc)		✓	949	787, 625 , 463, 301	
M	✓		317	151	
Q	✓		301	151	189
R	✓		609	300	
Q 3-(2-Caf-Glc)-3'-(6-Sinp-Glc)-4'-Glc		✓	1155	993, 831, 787, 669, 625 , 463, 301	18,25
Q 3-(2-Mcaf-Glc)-3'-(6-Sinp-Glc)-4'-Glc		✓	1185	1023, 817, 669, 655	
Q 3-(2-Fer-Glc)-3'-(6-Fer-Glc)-4'-Glc		✓	1139	977, 639, 463	
Q 3-(2-Fer-Glc)-3'-(6-Sinp-Glc)-4'-Glc		✓	1169	1007, 831, 669, 639, 463, 301	18,25,33

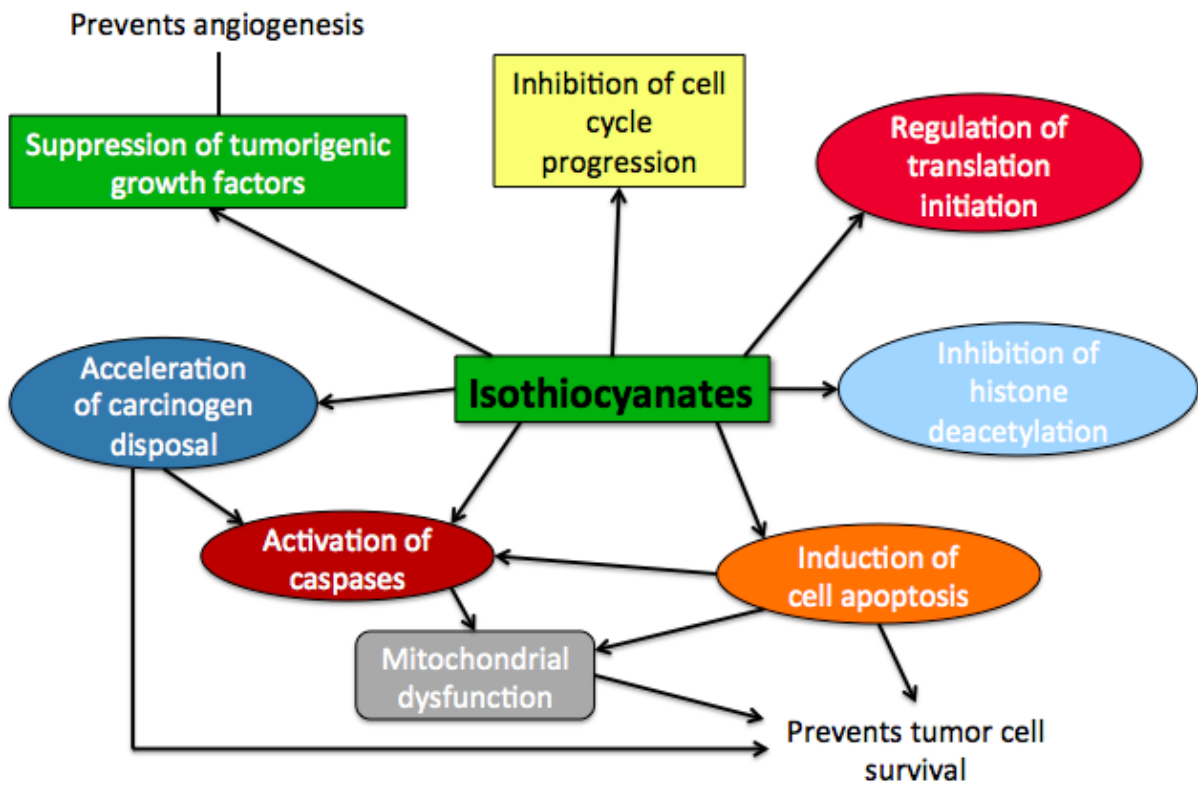
Q 3-(2-Sinp-Glc)-3'-(6-Sinp-Glc)-4'-Glc	✓	1199	1037, 831, 669, 463, 301	
Q 3,3',4-triGlc	✓	787	625, 463, 301	
Q 3,4'-diGlc-3'-(6-Fer-Glc)	✓	963	801, 639, 463, 301	18,25,33
Q 3,4'-diGlc-3'-(6-Mcaf-Glc)	✓	979	817, 655, 463, 301	
Q 3,4'-diGlc-3'-(6- <i>p</i> .Coum-Glc)	✓	933	771, 609, 463, 301	
Q 3,4'-diGlc-3'-(6-Sinap-Glc)	✓	993	831, 669, 463, 301	

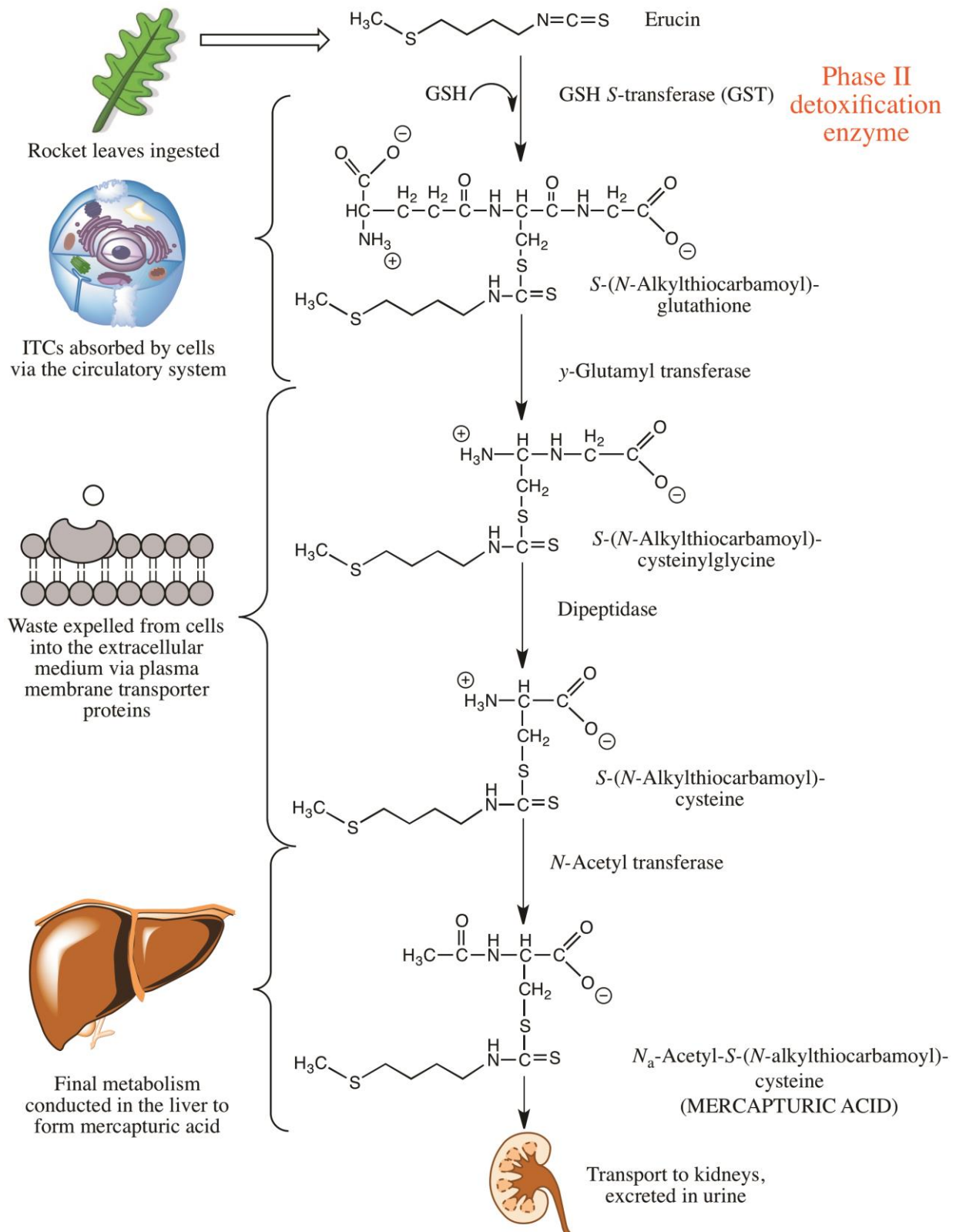
^a = Abbreviations: Caf, caffeyol; Mcaf, methoxycaffeyol; *p*.Coum, *p*-coumaroyl; Fer, feruloyl; Sinp, sinapoyl; Glc, glucoside;

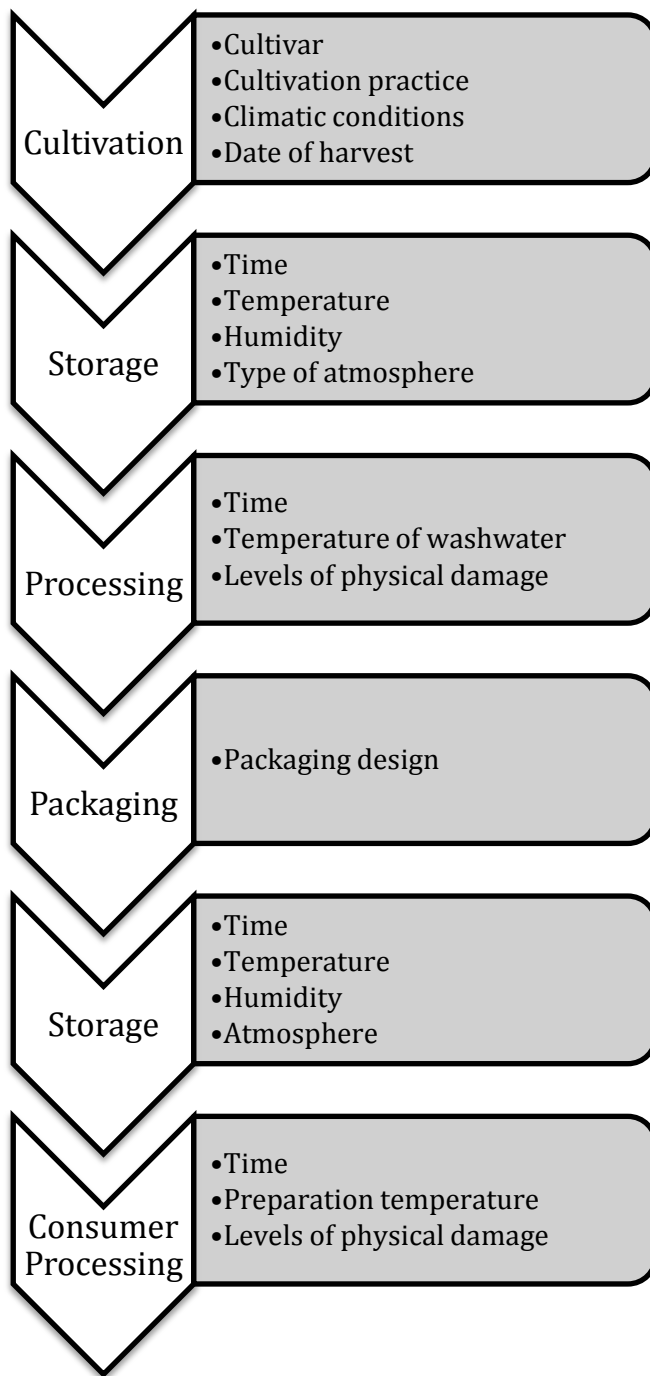
Q, quercetin; K, kaempferol; I, isorhamnetin; M, myricetin; R, rutin

^p = ✓ compound positively identified in species









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