



Sveriges lantbruksuniversitet Swedish University of Agricultural Sciences

This is an author produced version of a paper published in Trends in Ecology & Evolution. This paper has been peer-reviewed and is proof-corrected, but does not include the journal pagination.

Citation for the published paper: Inderjit; Wardle, David; Karban, Richard; Callaway, Ragan M. (2011) The ecosystem and evolutionary contexts of allelopathy. *Trends in Ecology & Evolution*. Volume: 26, Number: 12, pp 655-662. http://dx.doi.org/10.1016/j.tree.2011.08.003.

Access to the published version may require journal subscription. Published with permission from: Elsevier.

Standard set statement from the publisher: Posting and dissemination of authors accepted manuscript is allowed to personal websites, to institutional repositories, or to arXiv.

Epsilon Open Archive http://epsilon.slu.se

1	Review
2	
3	The ecosystem and evolutionary contexts of allelopathy
4	
5	Inderjit*, David A. Wardle <sup>1</sup> , Richard Karban <sup>2</sup> , and Ragan M. Callaway <sup>3</sup>
6 7 8 9 10 11	Department of Environmental Studies, Centre for Environmental Management of Degraded Ecosystems (CEMDE), University of Delhi, Delhi, India; <sup>1</sup> Department of Forest Ecology and Management, Swedish University of Agricultural Sciences; Umeå, Sweden; <sup>2</sup> Department of Entomology, University of California Davis, Davis, CA, USA; <sup>3</sup> Division of Biological Sciences, The University of Montana, Missoula, MT, USA.
12 13	*Corresponding author: Inderjit (inderjitdu@gmail.com)
14 15 16	Short title: Allelopathy: ecosystem-dependent interactions
17 18	Type of article: Review
19 20 21 22	Number of words in the abstract = 97 Number of words in the manuscript = 3855 Number of references = 91 Number of Boxes = 1
22 23 24	Number of Tables $= 1$

25 Abstract

27	Plants can release chemicals into the environment that suppress the growth and establishment of
28	other plants in their vicinity, a process known as 'allelopathy'. However, chemicals with
29	allelopathic functions have other ecological roles, such as plant defense, nutrient chelation, and
30	regulation of soil biota in ways that affect decomposition and soil fertility. These ecosystem-scale
31	roles of allelopathic chemicals can augment, attenuate or modify their community-scale
32	functions. In this review we explore allelopathy in the context of ecosystem properties, and
33	through its role in exotic invasions consider how evolution might affect the intensity and
34	importance of allelopathic interactions.
35	
36	Key words: allelopathy, allelochemicals, community ecology, evolution, exudates, herbivory,
37	invasion, soil microbes
38	
38 39	Allelochemical interactions in the context of communities and ecosystems
	Allelochemical interactions in the context of communities and ecosystems
39	Allelochemical interactions in the context of communities and ecosystems How populations are organized into higher units, or "communities", is a central issue in ecology
39 40	
39 40 41	How populations are organized into higher units, or "communities", is a central issue in ecology
<ol> <li>39</li> <li>40</li> <li>41</li> <li>42</li> </ol>	How populations are organized into higher units, or "communities", is a central issue in ecology [1]. The Russian ecologist T.A. Rabotnov [2] hypothesized that adaptation of plant species to the
<ol> <li>39</li> <li>40</li> <li>41</li> <li>42</li> <li>43</li> </ol>	How populations are organized into higher units, or "communities", is a central issue in ecology [1]. The Russian ecologist T.A. Rabotnov [2] hypothesized that adaptation of plant species to the chemistry of other species was crucial to this organization. Rabotnov focused on allelopathic
<ol> <li>39</li> <li>40</li> <li>41</li> <li>42</li> <li>43</li> <li>44</li> </ol>	How populations are organized into higher units, or "communities", is a central issue in ecology [1]. The Russian ecologist T.A. Rabotnov [2] hypothesized that adaptation of plant species to the chemistry of other species was crucial to this organization. Rabotnov focused on allelopathic interactions, which involve biochemically based suppression of the establishment and growth of
<ol> <li>39</li> <li>40</li> <li>41</li> <li>42</li> <li>43</li> <li>44</li> <li>45</li> </ol>	How populations are organized into higher units, or "communities", is a central issue in ecology [1]. The Russian ecologist T.A. Rabotnov [2] hypothesized that adaptation of plant species to the chemistry of other species was crucial to this organization. Rabotnov focused on allelopathic interactions, which involve biochemically based suppression of the establishment and growth of one plant by another. But plant-released secondary chemicals also have powerful effects on
<ol> <li>39</li> <li>40</li> <li>41</li> <li>42</li> <li>43</li> <li>44</li> <li>45</li> <li>46</li> </ol>	How populations are organized into higher units, or "communities", is a central issue in ecology [1]. The Russian ecologist T.A. Rabotnov [2] hypothesized that adaptation of plant species to the chemistry of other species was crucial to this organization. Rabotnov focused on allelopathic interactions, which involve biochemically based suppression of the establishment and growth of one plant by another. But plant-released secondary chemicals also have powerful effects on decomposition [3], herbivory [4], trophic interactions [5] and nitrogen cycling [6,7] (Figure 1).

49 effects [8-14]. Consideration of allelopathy in this integrated community and ecosystem context 50 requires the recognition of the large number of different processes that can be affected by the 51 same chemical or its derivatives, and the potential for the direct allelochemical effects of plants 52 on each other to be augmented, attenuated, modified or offset [11]. These other interactors can 53 enhance or reduce allelochemical production, change the persistence or effectiveness of 54 allelochemicals in soil, and select for higher or lower allelochemical concentrations over 55 evolutionary time. Understanding allelopathy in the context of communities and ecosystems can 56 be further developed by comparing the potential allelopathic effects of invasive species between 57 their native and introduced ranges [15-18]. Such biogeographic comparisons suggest that 58 evolutionary relationships among plants, and between plants and soil biota, may affect the role of 59 allelopathy in community organization [16].

Mere production of chemicals by a plant is not sufficient to ensure their allelopathic potential. 60 61 Abiotic and biotic environmental conditions determine the allelopathic potential of chemicals in 62 soil [10]. Recent studies have advanced our understanding of allelopathy by examining it in 63 environmental [12,19-21], biogeographic [15,16,22] and evolutionary [23,24] contexts. Our goal 64 is to discuss how (i) biotic and abiotic environmental conditions and (ii) evolutionary history 65 affect the production, fate, and effectiveness of allelopathic compounds in soils (Figure 1). 66 Specifically, we consider how habitat or site-specific characteristics, non-native ecosystems, and 67 environmental variables all influence the release, accumulation, and function of chemicals, and 68 thus affect the organization of natural systems.

71

73 The production, storage, and release of allelochemicals are key mechanisms of plant behavior 74 which affect almost all aspects of a plant's ecology [9]. These processes are affected by the 75 abiotic and biotic properties of the ecosystems in which plants grow [25], and chemicals 76 produced by plants in turn have strong effects on ecosystem properties. We propose that by 77 explicitly recognizing and integrating these ecosystem level effects, we will better understand 78 the various allelopathic, defensive, foraging, and signaling roles of chemicals in the organization 79 of natural communities (Figure 1). 80 Under natural conditions, allelopathic effects can result from interactive effects among 81 multiple compounds [26-29]. One of the best understood allelopathic systems involves the root 82 exudates of Sorghum bicolor which can contain up to 85% sorgoleone [30,31], However, is now 83 recognized that these exudates often contain both sorgoleone and its analogue (the lipid 84 resorcinol) in a 1:1 ratio [31], yielding the opportunity for studying potential interactive effects 85 among these two compounds. 86 Many chemicals released from the roots of plant species function to make nutrients available, 87 often through chelation, and can be quite substrate specific. Some chelators also appear to be 88 allelopathic. Chelating chemicals can degrade either slowly or rapidly, and this can increase or 89 decrease their biological activity [9,12,14]. However, many chelators are non-specific and hence 90 will bind with any of the metal ions with affinity decreasing along a lypotropic series. Most 91 natural soils are abundant in metal ions and hence it is difficult to find an uncomplexed chelator 92 under natural conditions. This aspect therefore needs more attention.

Consumer, competitor and soil microbe effects on allelochemical production and activity

93 There is considerable evidence for the direct inhibitory effects of specific allelochemicals
94 isolated from root exudates, leaf leachates and leaf volatiles of plants on other species. However,

95 in many cases, substantial variation has been found in the field concentrations and production of 96 the chemical, responses of target species, and the chemical's interactions with environmental 97 conditions, other phytochemicals, and other biota [10,12,32]. Such variation in allelochemical-98 environment interactions makes allelopathy difficult to consistently demonstrate in the field [but 99 see 16,33,34], and has led to conflicting evidence for the ecological relevance of particular 100 chemicals (Box 1) [19,32,35-37]. However, variation in the allelopathic potential of chemicals 101 among environments allows for more realistic appraisals of the role of ecological context in 102 driving allelopathic interactions [12]. Such processes provide alternative hypotheses for the direct 103 effects of allelochemicals on other species, and a broader understanding of the conditional effects 104 of allelopathy. Here we discuss how interactions between chemicals and ecosystem factors affect 105 the production, release, accumulation and activity of allelochemicals (Figure 1). 106 107 *Above-ground ecosystem influences on allelopathy* 108 109 Biotic components of the ecosystem such as herbivores, competitors, pathogens and belowground 110 decomposers can alter concentrations of chemicals already in plant tissues or released from 111 plants, or stimulate the production of chemicals that are otherwise not present or occur at very 112 low levels [38,39]. Here we discuss above-ground biotic influences of ecosystems on allelopathic 113 effects of herbivory-induced volatile chemicals in various environments.

Many allelochemicals can be induced by low concentrations of soil nutrients (although the ultimate cue is likely to be low concentrations in tissues). For example, iron deficiencies stimulate highly complex exudation responses [14]. Under iron limitation the roots of *Centaurea diffusa* prolong the release of 8-hydroxyquinoline that also mobilizes metals and makes them available for plant uptake [14]. Thus, the metal content of soils from different ecosystems is 119 likely to strongly influence the production and soil availability of 8-hydroxyquinoline, and 120 complex interactions between this allelochemical and metals may also determine its biological 121 activity [14]. Light intensity increases the root exudation of 8-hydroxyquinoline [14], which 122 exhibits a diurnal rhythm and reaches a maximum after 6 hours of exposure to light. Evaluating 123 the role of an allelochemical in the context of its abiotic environment should aid our 124 understanding on its release and allelopathic activities.

125 Induced secondary metabolite-based defenses are common in plants [40], and if the same 126 secondary metabolites or their derivatives are also allelopathic, herbivory might substantially 127 modify allelopathic interactions [see 11]. Karban [8] found that volatiles produced by 128 experimental clipping of sagebrush also inhibited germination and establishment of neighboring 129 plant species, thus providing experimental evidence of an herbivore-enhanced allelopathic effect. 130 The effects of allelochemicals depend not only on environmental conditions but also the genetic 131 landscape. For example, effects of herbivore-induced volatiles on neighboring sagebrush plants 132 were greater when the plants were genetically identical than when genetically different [41]. 133 Herbivory induces plant defenses that trigger the release of volatile organic compounds [38,42] 134 and accumulation of polyphenolics [43], and some of these chemicals may be allelopathic in 135 nature. Consistent with this, Bi et al. [44] found that exogenous application of methyl jasmonate, 136 a chemical that induces herbivore defenses in many plant species, led to the accumulation of 137 phenolics in rice and increased its allelopathic effects on other plants.

138

161

142 Below-ground influences of ecosystem processes driven by soil biota, genetic effects on root 143 interactions, and complex interactions among different root exudates appear to shape allelopathic 144 interactions. The general importance of soil communities in influencing the qualitative and 145 quantitative availability of allelochemicals is well established [45,46]. Microbial transformation 146 of biologically active chemicals commonly degrades their function, and evaluation of the activity 147 of an allelochemical in microbe-free substratum may therefore not be ecologically relevant. For 148 example, allelopathic effects of m-tyrosine, a metabolite exuded by the roots of *Festuca rubra* 149 ssp. *commutata*, have been demonstrated through filter paper bioassays free from naturally 150 occurring microbes [47]. However, Kaur et al. [19] showed that allelopathic effects of m-151 tyrosine were only evident in sterilized soil and diminished sharply in non-sterile soil with an 152 intact microbial community. Even this type of comparison must be interpreted with caution 153 because the scale of ecological interactions among roots, microbes and allelochemicals is 154 microscopic and ephemeral. For example, Bertin et al. [48] found that the predicted half-life of 155 m-tyrosine in soil in laboratory conditions was less than 1 day, indicative of rapid microbial 156 degradation. Sorgoleone, a major component of root exudates of Sorghum bicolor, is a potent 157 allelochemical [30] and microorganisms present in North American soils readily use it as a 158 carbon source [49]. It has been shown that the methoxy group of sorgoleone, which is responsible 159 for much of its activity, degrades rapidly in soil [49]. 160 In addition to the direct effects of allelochemicals on plant growth, their indirect effects may

162 allelochemical fractions from a phenolic-rich alpine forb, *Acomastylis rossii*, on soil respiration

- and the growth of the grass *Deschampsia caespitosa*. They found that some fractions had a direct

be mediated by microbial activity. Meier and Bowman [50] compared the effects of several

164 phytotoxic effect (i.e., which did not increase soil respiration but killed *D. caespitosa*) while 165 others appeared to work indirectly through the soil microbial community (i.e., which stimulated 166 soil respiration and reduced plant growth and plant N concentration). Their results provide a 167 compelling example of how phenolic compounds can inhibit root growth directly as well as 168 through interacting with soil biota. In another example, Alliaria petiolata can have negative 169 impacts on arbuscular mycorrhizal (AM) fungi and regeneration of seedlings native to North 170 America in soil from North America [51], but much weaker effects on AM fungi in soils from 171 Europe where it is native. Cantor *et al.* [52] showed that even very low field concentrations of 172 allyl isothiocynate (ca. 0.001mM) produced in the presence of A. petiolata strongly inhibited the 173 spore germination of the AM fungus Glomus clarum. However, Barto et al. [53] did not find 174 effects of A. petiolata extracts on the AM fungal colonization of roots or soils, and suggested that 175 potential alleopathic effects of A. petiolata might be due to direct inhibition of plant seedlings 176 and fungus before the formation of symbiosis.

177 The impacts of seasonal variation on the production and accumulation of allelochemicals [54] 178 and soil microbial communities [55] also contribute to the context-specificity of allelopathic 179 effects. For example, Alliaria petiolata accumulates glucotropaeolin three times more rapidly in 180 autumn than in spring, while accumulation of alliarinoside is highest in spring [54]. Fungal 181 communities and ectomycorrhizal colonization rates showed linear and curvilinear responses to 182 alliarinoside and glucosinolate concentrations, respectively [24]. Increasing concentrations of 183 alliarinoside were found to alter AM fungal communities, leading to a decline in AMF 184 colonization of *Quercus rubra* roots [24].

Belowground interactions among plants may also be genotype or ecotype dependent. For example, when the roots of different *Ambrosia dumosa* plants make contact they often stop growing, but there is a geographic and genotypic aspect to this response. For example, roots of 188 the plants from the same region show strong contact inhibition, but roots from plants from 189 different regions do not [56,57]. Cakile edentula plants allocate biomass differently to roots if 190 they are grown in the same pots shared by genetic relatives (kin) compared to pots shared by 191 strangers [58]. Lankau [59] reported that investment in high tissue concentrations of sinigrin 192 produced by *Brassica nigra* gave it an advantage in interspecific competition but a disadvantage 193 in intraspecific competition. Further, selection for *B. nigra* individuals that produced high levels 194 of sinigrin was stronger when grown with other species than with other individuals of its own 195 species.

196 Coexisting plant species can differ greatly in their growth response to allelochemicals 197 produced by a given plant species, and allelopathic effects can be highly species-specific 198 [16,22,60]. As such, there is a wide range of abilities (and perhaps mechanisms) among species 199 to protect themselves from chemical effects of their neighbors. Weir *et al.* [61] found that 200 Gaillardia grandiflora and Lupinus sericeus secrete oxalate in response to catechin exposure, 201 which could make these two species resistant to C. stoebe invasion. Exogenous application of 202 oxalate blocks the production of reactive oxygen species in the target plants, minimizing 203 oxidative damage caused by catechin. Such variation in the species-specific response of target 204 species may play a crucial role in the organization or assembly of plant communities in a similar 205 manner that it does for microbial communities [62], and provides an opportunity for allelopathy 206 to drive natural selection [63]. Variation in the ecological roles of secondary compounds is better 207 understood for consumer defense than for allelopathy, but for both types of interactions variation 208 is an important aspect of the effects of chemicals on communities and populations. 209 Issues of spatial scale and patchiness make studies of the roles of allelochemicals in soils

210 difficult to interpret. The effects of allelochemicals in soils are generally examined using "bulk

soils", where allelochemicals are added to a volume of soil that is orders of magnitude greater

212 than the soil volume in which the interactions occur. 'Realistic' concentrations of 213 allelochemicals are estimated for the average of the large soil volume. However, the action of 214 root-exuded chemicals often takes place at root-root interfaces. The use of estimated soil 215 concentrations is just one way to explore allelopathy in a reasonably realistic manner, but they 216 have limitations for the determination of the allelopathic functions of chemicals. If an 217 allelochemical is experimentally applied to soil in such a way as to allow it to transform before 218 contact with roots [12,19,34,37], then the failure to find an effect cannot be taken as evidence that 219 effects do not occur when roots are in close proximity to each other. This issue is, however, less 220 relevant when allelochemicals enter the soil through release from foliage or decomposition of 221 plant litter. 222 223 **Biogeographic comparisons of allelopathy: evolutionary changes in allelochemical effects** 224 225 The effects of allelopathy are also dependent on the evolutionary history of the interaction. 226 Understanding the mechanisms by which many exotic invasive plants strongly suppress their 227 neighbors in invaded but not native ranges has attracted growing recent attention. Allelopathy 228 and other biochemically driven interactions may contribute to the success of some exotic invasive 229 plants, and when either specific allelochemicals or general allelopathic effects are stronger 230 against potentially evolutionarily naïve species in invaded ranges, we gain insight into how 231 evolutionary history affects biological organization [64]. Biogeographical comparisons of the 232 ecological and biochemical traits of species in introduced and native ranges have proven useful 233 for evaluating mechanisms of invasion [65]. Examining the production and/or accumulation of 234 allelochemicals in novel and native environments, and the sensitivity of native residents and soil 235 communities to novel chemicals, can help understand these mechanisms.

236 The Novel Weapons Hypothesis (NWH) provides a possible explanation for biogeographic 237 patterns of interactions in different ecosystems. The NWH was first proposed in the context of 238 allelopathy as a potential mechanism for the success of *Centaurea diffusa* as an invader in North 239 America [66], and subsequently as a component of invasion by C. stoebe [67]. Recent studies on 240 biogeographic comparisons of exotic species in native and introduced ranges have shown some 241 support for NWH [15,16,18,68,69]. A recent meta-analysis of hypotheses for invasions, focusing 242 on trees, found that published evidence for the NWH resulted in a stronger effect size in support 243 of the idea than the effects sizes of six other hypotheses [70]. Barto et al. [71] provided evidence 244 in support of NWH by showing that the allelochemical profile of invasive A. petiolata was not 245 shared by any native Brassicaceae in North America. Further, Callaway and Ridenour [67] 246 suggested that stronger allelopathic effects in invaded regions could lead to selection for greater 247 allelopathic production and thus increased competitive ability. 248 Biogeographic differences in the effects of particular compounds between native and invaded 249 ranges may occur in part through a lack of adaptation by species and soil communities in the 250 invaded ranges. However, these types of biogeographic differences may also emerge or intensify 251 because of particular conditions in the novel environment. As such, soil biota can be powerful 252 ecosystem mediators of biogeographic differences in allelopathic effects [46]. For example, soil 253 microbial taxa that metabolize specific chemicals are likely to have undergone evolution to do so, 254 or at least to utilize a related group of chemicals. If plants that occur in a given region do not 255 produce a particular allelochemical, then those soil microbes that are required to metabolize it 256 may not be present when it is introduced by an invader. Thus, novel chemicals produced by 257 invaders may have prolonged resident times in invaded ranges and therefore be more biologically 258 active. Such indirect processes may reinforce biogeographic differences in plant-soil feedbacks 259 involving invasive species [72].

Soil communities from non-native ranges have also been shown to eliminate allelopathic effects of exotic plants. For example, the invader *A. petiolata* exerts allelopathic effects through glucosinolate exudation on the native species *Platanus occidentalis* in sterilized soil but not in non-sterile soil from the invaded range [73]. Future research would be required to determine whether soil microbial communities from locations that differ in their invasion history of *A. petiolata* also differ in their ability to degrade glucosinolate.

266

Potential evolutionary relationships: temporal declines in allelochemicals from invasive
 species

269

270 Plant species that are introduced into a novel environment would likely evolve in response to new 271 conditions over time, and other species that are native to that environment may in turn evolve in 272 response to the introduced species [16]. Such evolutionary responses have been reported for 273 populations of *Trifolium repens* that have co-adapted to (and with) local competitors [74], and for 274 populations of native soapberry bugs (*Leptocoris tegalicus*) that have adapted to various 275 introduced host plants [75]. Some native residents in the naturalized range of C. stoebe have 276 exhibited tolerance to it relative to individuals of other native species that have not previously 277 encountered the invader [76]. Individuals grown from seeds of parents that have survived 278 exposure to allelochemicals from C. stoebe have become more resistant to its invasion. This is 279 consistent with the NWH, and suggests that allelopathy may play a role in evolution between 280 neighbors in the non-native ranges.

Biogeographic variation in the production of volatile sesquiterpenes in particular could be due to differences in herbivore densities between the native and introduced ranges [77]. Recently, it has been shown [15] that lower amounts of volatile chemicals were released by plants from

exotic populations of the invasive plant *Ageratina adenophora* than by plants from native
populations grown in a common environment. However, it is not known whether such differences
in volatile emissions are evolutionary consequences of interactions with other species or due to
founder effects.

288 An allelochemical produced by a species can provide multiple ecological functions, making its 289 effects highly dependent on specific environmental conditions. Further, allelochemicals with 290 multiple functions should be selected for because this spares the plant the cost of producing 291 several different compounds [11]. Glucosinolates and their derivatives have been found to have 292 multiple functions as mediators of plant–plant, plant–microbe, and plant–insect interactions [59]. 293 Lankau and Kliebenstein [78] found that competition and herbivory determined the accumulation 294 and fitness consequences of sinigrin for *B. nigra*. Further, it has been shown that the fitness costs 295 and benefits of sinigrin conformed to optimal defense theory only in the absence of competition, 296 apparently due to its multiple functions [11,78]. Further, Oduor *et al.* [79] found that invasive 297 populations of *B. nigra* had higher levels of sinigrin which defends the invader against generalist 298 herbivores. An increase in resistance against generalist herbivores and growth performance of B. 299 *nigra* in its introduced ranges compared to its native range further supports the hypothesis that 300 defenses have shifted [79]. Sinigrin from *B. nigra* is also reported to possess allelopathic 301 activities, which provide a competitive advantage to *B. nigra* over heterospecific neighbors [59]. 302 Lankau et al. [23,24] examined the production, release and impact of glucosinolates from A. 303 petiolata along a gradient of invasion history i.e., from early invaded to recently-invaded 304 populations. They found a significant decline in the production of glucosinolates and an increase 305 in the community's resistance to A. petiolata invasion over time. Following an initial decline in 306 the number of operational taxonomic units (OTUs) of bacteria, fungi and AM fungi, an increase 307 was observed in older invaded sites [24]. The observed development of resistance to exotic

308 invasion in late invasion stages could lead to more species rich native communities. However, the 309 eventual outcome of the evolutionary changes is still unclear. Lankau et al. [80] found that the 310 higher production of sinigrin by introduced *B. nigra* suppressed mycorrhizal abundance, which 311 adversely affected the growth of heterospecific competitors but not non-mycorrhizal conspecifics. 312 Such rapid selection based on tradeoffs between competitive advantages against either 313 conspecifics or heterospecifics contributes to intransitive competitive networks which affect 314 genetic and species diversity in communities [80]. Studying evolutionary relationships between 315 native and non-native communities and ecosystems along gradients of invasion history has 316 significant potential for improving understanding of the role of allelopathy in community 317 organization.

318

010	$\alpha$	•
4 I U	l 'one	ngiong
319	COLU	lusions
/		

320

321 It is important to identify how variation in the environment establishes conditionality in 322 allelopathic interactions. Sources of such variation include (1) the impact of soil chemistry on 323 production and effects of allelochemicals, (2) the impact of consumers, competitors, and soil 324 microbes on production and effects of allelochemicals, (3) evolutionary changes in 325 allelochemical effects, and (4) declines in allelochemical production and activity from invasive 326 species over time. A major gap in current allelopathy research involves the role of conditional 327 ecosystem factors that drive allelopathic processes and how these change over space and time 328 (Figure 1). Further, despite recent advances, we still have a limited understanding of the role of 329 evolution over time in the production, release and eventual loss of activity of biogeographically 330 novel chemicals.

331 The production, fate, and effectiveness of allelopathic compounds in soils is influenced by 332 environmental conditions and evolutionary history, generating a need for allelopathic interactions 333 to be studied across spatial and temporal scales (Figure 1). Over very small scales (microns to 334 millimeters; seconds to hours), processes in the rhizosphere, such as microbial-driven breakdown 335 of allelochemicals or metal chelation, dominate the influences of allelochemicals. Over small 336 scales (millimeters to meters; hours to months), organismal responses are important, for instance, 337 the increased production of chemicals following herbivore attack. At the medium scale (meters to 338 kilometers; months to years), variation in the plant and soil communities, and abiotic soil 339 conditions become increasingly important, if different species are more or less susceptible to the 340 allelochemicals. Finally, at the large scales (kilometers and beyond; years and beyond), the 341 evolutionary history of the allelopathic plant and the recipient soil and plant community assumes 342 increasing significance (Figure 1).

343 Continuing to quantify various aspects of how ecosystem factors influence allelopathy is key 344 to better understanding of how plants interact with each other. Other important steps would 345 include greater focus on conducting experiments under natural conditions, comparing single 346 chemical effects to whole-exudate effects, profiling metabolites, and conducting bioassays in 347 search of unidentified compounds that mediate these interactions. More generally, there is a 348 greater need for understanding of how biotic and abiotic environmental conditions and 349 evolutionary history affect the production, fate, and effectiveness of allelopathic compounds in 350 soils. Recent work linking chemical ecology to biogeography and evolutionary biology has 351 provided new perspectives on biochemical processes in ecosystems. Expanded use of 352 biogeographical and evolutionary approaches will improve our understanding of the release of 353 allelochemicals over a range of abiotic and biotic conditions and how those conditions determine 354 the outcomes of allelochemical interactions.

### 356 Acknowledgements

357

- 358 Inderjit acknowledges research funding from the University of Delhi and Council of Scientific &
- 359 Industrial Research (CSIR). Ragan M. Callaway thanks the National Science Foundation and
- 360 DoD SERDP for support, and David A. Wardle acknowledges support from a Wallenberg
- 361 Scholars award. We thank two reviewers for their valuable comments.

363	63 <b>References</b>		
364			
365	1.	Lortie, C.J. et al. (2004) Rethinking plant community theory. Oikos 107, 433-438	
366	2.	Rabotnov, T.A. (1982) Importance of the evolutionary approach to the study of	
367		allelopathy. <i>Ékologia 3</i> , 5–8	
368	3.	Hättenschwiler, S., Coq, S., Barantal, S. and Handa, I.T. (2010) Leaf traits and	
369		decomposition in tropical rainforests: revisiting some commonly held views and towards a	
370		new hypothesis. <i>New Phytol.</i> 189, 950-965	
371	4.	Karban, R., Shiojiri, K., Huntzinger, M. and McCall, A.C. (2006) Damage-induced	
372		resistance in sagebrush: volatiles are key to intra- and interplant communication. <i>Ecology</i>	
373		87, 922-930	
374	5.	Hättenschwiler, S. and Jørgensen, H.B. (2010) Carbon quality rather than stoichiometry	
375	5.	controls litter decomposition in a tropical rain forest. J. Ecol. 98, 754-763	
376	6.	Northup, R.R., Dahlgren, R.A. and McColl, J.G. (1998) Polyphenols as regulators of plant-	
377	0.		
		litter-soil interactions in northern California's pygmy forest: a positive feedback?	
378	7	Biogeochemistry 42, 189-220	
379 380	7.	Hättenschwiler, S. and Vitousek, P. (2000) The role of polyphenols in terrestrial ecosystem nutrient cycling. <i>Trends Ecol. Evol.</i> 15, 238-243	
381	8.	Karban, R. (2007) Experimental clipping of sagebrush inhibits seed germination of	
382	0.	neighbours. Ecol. Lett. 10, 791-797	
383	9.	Metlen, K.L., Aschehoug, E.T. and Callaway, R.M. (2009) Plant behavioural ecology:	
384	<i>.</i>	dynamic plasticity in secondary metabolites. <i>Plant Cell Environ</i> . 32, 641-653	
385	10.	Inderjit and Weiner, J. (2001) Plant allelochemical interference or soil chemical ecology?	
386	10.	Persp. Plant Ecol. Evol. Systemat. 4, 3-12	
387	11.	Siemens, D., Garner, S., Mitchell-Olds, T. and Callaway, R.M. (2002) The cost of defense	
388	11.	in the context of competition. <i>Brassica rapa</i> may grow <i>and</i> defend. <i>Ecology</i> 83, 505-517	
389	12.	Pollock, J.L., Callaway, R.M., Thelen, G.C. and Holben, W.E. (2009) Catechin-metal	
390	12.	interactions as a mechanism for conditional allelopathy by the invasive plant, <i>Centaurea</i>	
391		maculosa. J. Ecol. 97, 1234-1242	
392	13.	Lankau, R.A. and Strauss, S.Y. (2008) Community complexity drives patterns of natural	
393	15.	selection on a chemical defense of <i>Brassica nigra</i> . <i>Am. Nat.</i> 171, 150-161	
393 394	14.	Tharayil, N., Bhowmik, P.C., Alpert, P., Walker, E., Amarasiriwardena, D. and Xing, B.	
39 <del>4</del> 395	14.	(2009) Dual purpose secondary compounds: phytotoxin of <i>Centaurea diffusa</i> also	
395 396			
	15	facilitates nutrient uptake. <i>New Phytol.</i> 181, 424-434	
397	15.	Inderjit <i>et al.</i> (2011) Volatile chemicals from leaf litter are associated with invasiveness of	
398	16	a neotropical weed in Asia. <i>Ecology</i> 92, 316-324 Thomas A.S. Tholan, C.C. Discours, A. and Callenver, P.M. (2000) Boot any data is	
399	16.	Thorpe, A.S., Thelen, G.C., Diaconu, A. and Callaway, R.M. (2009) Root exudate is	
400		allelopathic in invaded community but not in native community: field evidence for the	
401	17	novel weapons hypothesis. J. Ecol. 97, 641-645	
402	17.	Ens, E.J., French, K. and Bremner, J.B. (2009) Evidence for allelopathy as a mechanism of	
403		community composition change by an invasive exotic shrub, <i>Chrysanthemoides monifera</i>	
404	10	ssp. rotundata. Plant Soil 316, 125-137	
405	18.	Zhang, D., Zhang, J., Yang, W. and Wu, F. (2010) Potential allelopathic effect of	
406	10	<i>Eucalyptus grandis</i> across a range of plantation ages. <i>Ecol. Res.</i> 25, 13-23	
407	19.	Kaur, H., Kaur, R., Kaur, S., Baldwin, I.T. and Inderjit (2009) Taking ecological function	
408		seriously: soil microbial communities can obviate allelopathic effects of released	
409		metabolites. PLoS ONE, 4, e4700	

- Tharayil, N., Bhowmik, P.C. and Xing, B. (2006) Preferential sorption of phenolic
  phytotoxins to soil: implications for the availability of allelochemicals. *J. Agric. Food Chem.* 54, 3033-3040
- Tharayil, N., Bhowmik, P.C. and Xing, B. (2008) Phyto-availability of allelochemicals as
  affected by companion compounds in soil-microbial systems. *J. Agric. Food Chem.*, 56,
  3706 3713
- 416 22. He, W.M. *et al.* (2009) Novel weapons and invasions: biogeographic differences in the
  417 competitive effects of *Centaurea maculosa* and its root exudates (±)-catechin. *Oecologia*418 159, 803-815
- Lankau, R.A., Nuzzo, V., Spyreas, G. and Davis A.S. (2009) Evolutionary limits ameliorate
  the negative impact of an invasive plant. *Proc. Natl. Acad. Sci. USA* 106, 15362-15367
- 421 24. Lankau, R.A. (2011) Resistance and recovery of soil microbial communities in the face of
   422 *Alliaria petiolata* invasions. *New Phytol.* 189, 536-548
- 423 25. Wardle, D.A., Nilsson, M.C., Gallet, C. and Zackrisson, O. (1998) An ecosystem level
  424 perspective of allelopathy. *Biol. Rev.* 73, 305-319
- Blum, U., Dalton, B.R. and Shann, J.R. (1985) Effects of various mixtures of ferulic acid
  and some of its microbial metabolic products on cucumber leaf expansion and dry matter in
  nutrient culture. J. Chem. Ecol. 11, 619-641 2
- Blum, U., Gerig, T. M. and Weed, S.B. (1989) Effects of mixtures of phenolic acids on leaf
  area expansion of cucumber seedlings grown in different pH Portsmouth A1 soil materials. *J. Chem. Ecol.* 15, 2413-2423
- 431 28. Gerig, T. M. and Blum, U. (1993) Modification of an inhibition curve to account for effects
  432 of a second compound. *J. Chem. Ecol.* 19, 2783-2790
- 433 29. Inderjit, Striebig, J. and Olofsdotter, M. (2002) Joint action of phenolic acid mixtures and
  434 its significance in allelopathy research. *Physiol. Plant.* 114: 422-428 6
- 435 30. Baerson, S.R. *et al.* (2008) A functional genomics investigation of allelochemical
  436 biosynthesis in *Sorghum bicolor* root hairs. *J. Biol. Chem.* 283, 3231-3247
- 437 31. Dayan, F.E., Howell, J. and Weidenhamer, J.D. (2009) Dynamic root exudation of
  438 sorgoleone and its *in planta* mechanism of action. *J. Exp. Bot.* 60, 2107-2117
- 439 32. Inderjit, Callaway, R.M. and Vivanco, J.M. (2006) Plant biochemistry helps to understand
  440 invasion ecology. *Trends Plant Sci.* 11, 574-580
- 33. Nilsson, M-C. (1994) Separation of allelopathy and resource competition by the boreal
  dwarf shrub *Empetrum hermaphroditum* Hagerup. *Oecologia* 98, 1-7
- 443 34. Inderjit, Pollock, J.L., Callaway, R.M. and Holben, W. (2008) Phytotoxic effects of (±)444 catechin *in vitro*, in soil, and in the field. *PLoS One* 3, e2536.
- Blair, A.C., Nissen, S.J., Brunk, G.R. and Hufbauer, R.A. (2006) A lack of evidence for an
  ecological role of the putative allelochemical (+/-)-catechin in spotted knapweed invasion
  success. J. Chem. Ecol. 32, 2327-2331
- 448 36. Chobot, V., Huber, C., Trettenhahn, G. and Hadacek, F. (2009) (±)-catechin: chemical
  449 weapon, antioxidant, or stress regulator? *J. Chem. Ecol.* 35, 980-996
- 450 37. Duke, S.O. *et al.* (2009) Is (-)-catechin a novel weapon of spotted knapweed (*Centaurea stoebe*)? *J. Chem. Ecol.* 35, 141-153
- 452 38. Baldwin, I.T., Halitschke, R., Paschold, A., von Dahl, C.C. and Preston, C.A. (2006)
  453 Volatile signaling in plant-plant interactions: "Talking trees" in the genomics
  454 era. *Science* 311, 812-815
- 455 39. Lohmann, M., Scheu, S. and Müller, C. (2009) Decomposers and root feeders interactively
  456 affect plant defence in *Sinapis alba. Oecologia* 160, 289-298

457 40. Agrawal, A.A. (1998) Induced responses to herbivory and increased plant performance. 458 Science 279, 1201–1202. 459 41. Karban, R. and Shiojiri, K. (2009) Self-recognition affect plant communication and 460 defense. Ecol. Lett. 12, 502-506 461 42. Karban, R. (2010) The ecology and evolution of induced resistance against herbivores. 462 Func. Ecol. 25, 339-347 463 Roitto, M. et al. (2008) Induced accumulation of phenolics and sawfly performance in 43. 464 Scots pine in response to previous defoliation. Tree Physiol. 29, 207-216 465 44. Bi, H.H., Zheng, R.S., Su, L.M., An, M. and Luo, S.M. (2007) Rice allelopathy induced by methyl jasmonate and methyl salicylate. J. Chem. Ecol. 33, 1089-1103 466 467 Inderjit. (2005) Soil microorganisms: an important determinant of allelopathic activity. 45. 468 Plant Soil 274, 227-236 469 Inderjit and van der Putten, W.H. (2010) Impacts of soil microbial communities on exotic 46. 470 plant invasion. Trends Ecol. Evol. 25, 512-519 471 Bertin, C. et al. (2007) Grass roots chemistry: meta-tyrosine, an herbicidal nonprotein 47. 472 amino acid. Proc. Natl. Acad. Sci. USA 104, 16964-16969 473 Bertin, C., Harmon, R., Akaogi, M., Weidenhamer, J.D. and Weston, L.A. (2009) 48. 474 Assessment of the phytotoxic potential of m-tyrosine in laboratory soil bioassays. J. Chem. 475 Ecol. 35, 1288-1294 476 49. Gimsing, A.L., Bælum, J., Dayan, F.E., Locke, M.E., Sejerø, L.H. and Jacobsen, C.S. 477 (2009) Mineralization of the allelochemical sorgoleone in soil. Chemosphere 76, 1041-478 1047 479 Meier, C.L. and Bowman, W.D. (2008) Phenolic-rich leaf carbon fractions differentially 50. 480 influence microbial respiration and plant growth. Oecologia 158, 95-107 481 51. Callaway, R.M. et al. (2008) Novel weapons: invasive plant suppresses fungal mutualists in 482 America but not in its native Europe. Ecology 89, 1043-1055 483 Cantor, A., Hale, A., Aaron, J., Trow, M.B. and Kalisz, S. (2011) How allelochemical 52. 484 concentrations detected in garlic mustard-invaded forest soil inhibit fungal growth and 485 AMF spore germination. *Biol. Invs.* (in press) 486 Barto, K., Friese, C. and Cipollini, D. (2010) Arbuscular mycorrhizal fungi protect a native 53. 487 plant from allelopathic effects of an invader. J. Chem. Ecol. 36, 351-360 488 54. Barto, E.K. and Cipollini, D. (2009) Half-lives and field soil concentrations of Alliaria 489 petiolata secondary metabolites. Chemosphere 76, 71-75 490 55. Lankau, R.A. (2011) Intraspecific variation in allelochemistry determines an invasive 491 species' impact on soil microbial communities. Oecologia 165, 453-463 492 Mahall, B.E. and R.M. Callaway. (1996) Geographic and genetic contributions to root 56. 493 communication in Ambrosia dumosa. Amer. J. Bot. 83, 93-98 494 Mahall, B.E. and R.M. Callaway. (1991) Root communication among desert shrubs. Proc. 57. 495 Natl. Acad. Sci. USA 88, 874-876 496 58. Dudley, S.A. and File, A.L. (2007) Kin recognition in an annual plant. Biol. Lett. 3, 435-497 438 498 59. Lankau, R. (2008) A chemical trait creates a genetic trade-off between intra- and 499 interspecific competitive ability. Ecology 89, 1181-1187. 500 Gómez-Aparicio, L. and Canham, C.D. (2008) Neighbourhood analysis of the allelopathic 60. 501 effects of the invasive tree Ailanthus altissima in temperate forests. J. Ecol. 96, 447-458

502	61.	Weir, T.L. et al. (2006) Oxalate contributes to the resistance of Gaillardia
503		grandiflora and Lupinus sericeus to a phytotoxin produced by Centaurea maculosa. Planta
504		223, 785-795
505	62.	Callaway, R.M. and Howard, T.G. (2006) Competitive networks, indirect interactions, and
506		allelopathy: a microbial viewpoint on plant communities. Prog. Bot. 68, 317-335
507	63.	Callaway, R.M., Hierro, J.L. and Thorpe, A.S. (2005) Evolutionary trajectories in plant and
508		soil microbial communities: <i>Centaurea</i> invasions and the geographic mosaic of
509		coevolution. In Exotic Species Invasions: Insights into Ecology, Evolution and
510		<i>Biogeography</i> (Sax, D.F., Gaines, S.D. and Stachowicz, J.J., eds.), pp. 341-363. Sinauer,
511		Sunderland
512	64.	Inderjit and Callaway, R.M. (2003) Experimental designs for the study of allelopathy. <i>Plant</i>
513	0	Soil 256, 1-11
514	65.	Hierro, J.L., Maron, J.L. and Callaway, R.M. (2005) A biogeographical approach to plant
515	001	invasions: the importance of studying exotics in their introduced and native range. J. Ecol.
516		93, 5-15
517	66.	Callaway, R.M. and Aschehoug, E.T. (2000) Invasive plant versus their new and old
518	00.	neighbors: a mechanism for exotic invasion. <i>Science</i> 290, 521-523
519	67.	Callaway, R.M. and Ridenour W.M. (2004) Novel weapons: invasive success and the
520		evolution of increased competitive ability. Front. Ecol. Environ. 2, 436-443
521	68.	Kim, Y.O. and Lee, E.J. (2011) Comparison of phenolic compounds and the effects of
522		invasive and native species in East Asia: support forb the novel weapons hypothesis. <i>Ecol.</i>
523		Res. 26, 87-94
524	69.	Ni, G.Y., Schaffner, U., Peng, S.L. and Callaway, R.M. (2011) Acroptilon repens, an Asian
525		invader, has stronger competitive effects on species from America than species from its
526		native range. Biol. Invas. 12, 3653-3663
527	70.	Lamarque, L.J., Delzon, S. and Lortie, C.J. (2011) Tree invasions: a comparative test for
528		the dominant hypotheses and functional traits. <i>Biol. Invas.</i> (in press)
529	71.	Barto, K., Powell, J.R. and Cipollini, D. (2010) How novel are the chemical weapons of
530		garlic mustard in North America forest understories. Biol. Invas. 12, 3465-3471
531	72.	Callaway, R.M., Thelen, G.C., Rodriguez, A. and Holben, W.E. (2004) Soil biota and
532		exotic plant invasion. Nature 427, 731-733
533	73.	Lankau, R.A. (2010) Soil microbial communities alter allelopathic competition between
534		Alliaria petiolata and a native species. Biol. Invas. 12, 2059-2068
535	74.	Turkington, R. (1989) The growth, distribution and neighbor relationships of Trifolium
536		repens in a permanent pasture. V. The coevolution of competitors. J. Ecol. 77, 717-733
537	75.	Carroll, S.P. et al. (2005) And the beak shall inherit – evolution in response to invasion.
538		Ecol. Lett. 8, 944-951
539	76.	Callaway, R.M., Ridenour, W.M., Laboski, T., Weir, T. and Vivanco, J.M. (2005) Natural
540		selection for resistance to the allelopathic effects of invasive plants. J. Ecol. 93, 576-583
541	77.	Mooney, K.A., Halitschke, R., Kessler, A. and Agrawal, A.A. (2010) Evolutionary trade-
542		off in plants mediate the strength of trophic cascades. Science 327, 1642-1644
543	78.	Lankau, R.A. and Klienbenstein, D.J. (2009) Competition, herbivory and genetics interact
544		to determine the accumulation and fitness consequences of a defense metabolite. J. Ecol.
545		97, 78-88
546	79.	Oduor, A.M.O., Lankau, R.A., Strauss, S.Y. and Gómez, J.M. (2011) Introduced Brassica

*nigra* populations exhibit greater growth and herbivore resistance but less tolerance than
 native populations in the native range. *New Phytol.* 191, 536-544

- 549 80. Lankau, R.A., Wheeler E., Bennett, A.E. and Strauss, S.Y. (2011) Plant-soil feedbacks
  550 contribute to an intransitive competitive network that promotes both genetic and species
  551 diversity. *J. Ecol.* 99, 176-185
- 552 81. Calow, P. (1999) Blackwell's Concise Encyclopedia of Ecology. Blackwell, Oxford
- Stermitz, F.R., Hufbauer, R.A and Vivanco, J.M. (2009) Retraction. *Plant Physiol.* 151, 967
- 83. Perry, L.G., Thelen, G.C., Ridenour, W.M., Callaway, R.M., Paschke, M.W. and Vivanco,
  J.V. (2007) Concentrations of the allelochemical (+/-)-catechin in *Centaurea maculosa*soils. *J. Chem. Ecol.* 33, 2337-2344
- Schultz, M.J. (2008) Soil ecological interactions of spotted knapweed and native plant
   species. M.S. Thesis, Colorado State University, Fort Collins, Colorado
- 560 85. Tharayil, N. and Triebwasser, D.J. (2010) Elucidation of a diurnal pattern of catechin
  561 exudation by *Centaurea stoebe*. J. Chem. Ecol. 36, 200–204
- Marler, M.J., Zabinski, C.A., Wojtowicz, T. and Callaway, R.M. (1999) Mycorrhizae and
  fine root dynamics of *Centaurea maculosa* and native bunchgrasses in western Montana. *Northwest Sci.* 73, 217-224
- 565 87. LeJeune, K.D. and Seastedt, T.R. (2001) *Centaurea* species: the forb that won the west.
  566 *Conser. Biol.* 15, 1568-1574
- 567 88. Callaway, R.M., Thelen, G.C., Barth, S., Ramsey, P.W. and Gannon, J.E. (2004) Soil fungi
  568 alter interactions between the invader *Centaurea maculosa* and North American natives.
  569 *Ecology* 85, 1062-1071
- 570 89. Thorpe, A.S., Archer, V. and Deluca, T.H. (2006) The invasive forb, *Centaurea maculosa*,
  571 increases phosphorus availability in Montana grasslands. *Appl. Soil Ecol.* 32, 118-122
- 572 90. Thorpe, A. and Callaway, R.M. (2011) Biogegraphic differences in the effects of
  573 *Centaurea stoebe* on the soil nitrogen cycle: novel weapons and soil microbes. *Biol. Invas.*574 13, 1435-1445
- 575 91. Baldwin, I.T. (2003) Finally, proof of weapons of mass destruction. Sci. STKE pe42
- 576
- 577 578

#### Glossary

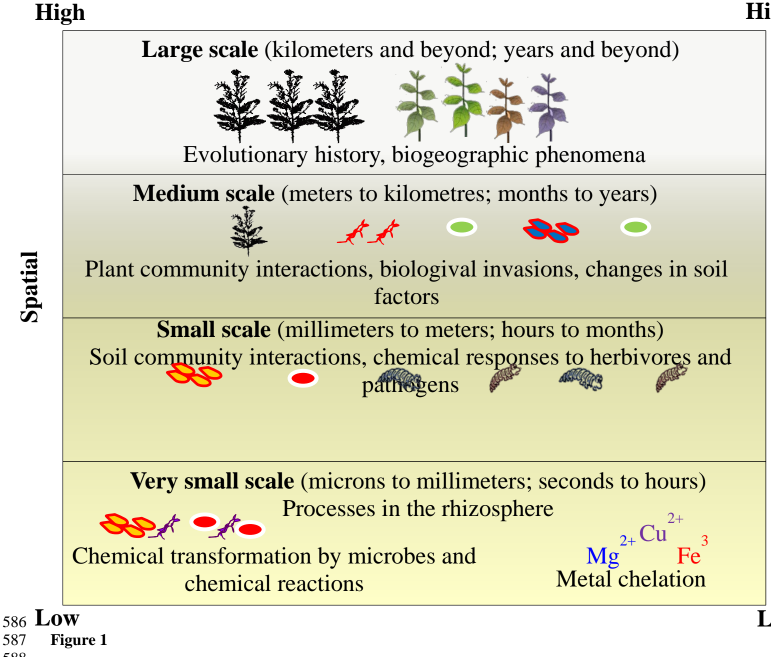
**Allelopathy:** Suppression of the growth and/or establishment of neighboring plants by chemicals released from a plant or plant parts.

Allelochemicals: Secondary compounds of plant origin that interact with their environment and possess allelopathic activities.

**Homeostatis:** The tendency of a biological system (organism, population, community or ecosystem) to resist changes and to remain in the state of equilibrium or change its properties in such a way as to minimize the impact of outside factors [81].

**Novel weapons hypothesis** (**NWH**): The idea that some invasive plant species produce secondary metabolites that are novel in their non-native ranges and that this novelty provides advantages to the invasive species as it interacts with native plants, microbes or generalist herbivores.

- Figure 1. The impact of ecosystem factors, biogeographic variations and coevolutionary
- relationships on the production, release and activity of allelochemicals along spatial and temporal
- scales.



**Box 1. Catechin as a novel weapon.** (-)-Catechin, reported to be exuded from the roots of a Eurasian invader in North America, *Centaurea maculosa* (*C. stoebe*), was the first isolated chemical discussed as a possible 'novel weapon' [67]. Initial work on this compound used (-)- catechin but subsequent experimental studies used  $(\pm)$ -catechin because root exudates of *C. stoebe* contain a racemic mixture of (+)- and (-)- catechin. Early reports of consistently high rates of exudation have not been reproducible using protocols similar to those in the original experiment [see retraction, 82]. Catechin has been reported at very low concentrations in soil in the rhizospheres of *C. stoebe* [35] but high concentrations may occur periodically [83,84]. The phytotoxic effects of the enantiomeric form ( $\pm$ )-catechin, and the (+) form have been demonstrated *in vitro*, in sand culture, in controlled experiments with field soils, and in the field [12,16,22,34 and citations within], but others have not found either the + or the – form to be phytotoxic [36,37].

Tharayil and Triebwasser [85] quantified catechin release at picomolar levels by roots of C. stoebe in hydroponic medium and showed a diurnal rhythm in its exudation in response to light. There is also evidence that this invader's impact is also due to interactions with the soil ecosystem including through effects on nitrogen (N) and phosphorous (P) cycling and on soil fungi [72,86-89]. Recently, Thorpe and Callaway [90] examined biogeographical differences in the responses of soil communities to C. stoebe and catechin by studying the effects of catechin on soil ammonification and nitrification in both native (Romania) and non-native (Montana) ranges. Catechin and *C. stoebe* were linked to similar reductions of resin-extractable nitrates and gross nitrification in Montana soils but not in Romanian soils where C. stoebe is native. As discussed below, we do not know if the consistency and rate of catechin exudation and its concentration at root-root and root-bacteria interfaces is adequate to drive substantial effects in natural systems, but biogeographical differences in ecosystem effects controlled by soil bacteria suggests that novel chemicals might affect soil nutrients by influencing soil communities as well as other plants, and that these effects have an evolutionary context.

#### **INSERT Figure I HERE**

Figure I. Abiotic and biotic ecosystem components influence the release, accumulation and activity of catechin. Unresolved issues regarding whether catechin has an important role as a novel chemical and under which environmental conditions could be addressed by studying the natural release of catechin in different ecosystems, or across gradients of invasion history.

# Factors that may influence production, release or activity of catechin

- Soil biota [32]
- Soil chemistry [12]
- Impact on nitrification [90]
- Cell wall elicitors from soil fungi [91]
- Biogeographic differences in neighbour sensitivity [63,64]
- Abiotic factors (e.g., light) [85]

## Key factors that we don't know

- Phytotoxicity through natural release
- Evolutionary changes in impact over time in non-native ranges
- Potential for other chemicals in exudates or foliar leachates to alter or exceed in importance
- Differences in impact of soil biota from native and non-native ranges
- Seasonal differences in release or impact

593	
594	
595	
596	
597	
598	
599	
600	
601	Figure I (for Box 1)
602	