Review

Plant volatiles in polluted atmospheres: stress responses and signal degradation

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ABSTRACT

Plants emit a plethora of volatile organic compounds, which provide detailed information on the physiological condition of emitters. Volatiles induced by herbivore feeding are among the best studied plant responses to stress and may constitute an informative message to the surrounding community and further function in plant defence processes. However, under natural conditions, plants are potentially exposed to multiple concurrent stresses with complex effects on the volatile emissions. Atmospheric pollutants are an important facet of the abiotic environment and can impinge on a plant’s volatile-mediated defences in multiple ways at multiple temporal scales. They can exert changes in volatile emissions through oxidative stress, as is the case with ozone pollution. The pollutants, in particular, ozone, nitrogen oxides and hydroxyl radicals, also react with volatiles in the atmosphere. These reactions result in volatile breakdown products, which may themselves be perceived by community members as informative signals. In this review, we demonstrate the complex interplay among stresses, emitted signals, and modification in signal strength and composition by the atmosphere, collectively determining the responses of the biotic community to elicited signals.

Key-words: abiotic stress; biotic stress; cross-stress tolerance; induced volatiles; interactive stresses; multiple stresses; volatile-mediated interactions.

INTRODUCTION

Plants communicate with other community members by emitting a blend of volatile organic compounds (VOCs). The chemical composition and ratios of compounds in the blend constitute the plant scent. The constitutively emitted VOC blend and ratios of compounds in that blend are often considered to be driven by species taxonomy with a large degree of differentiation among species (Bruce et al. 2005). However, there is sometimes a large degree of within-species variation in volatile emissions, which can include general heterogeneity, geographical heterogeneity or chemotype diversification (Loreto et al. 2009; Bäck et al. 2012; Fineschi et al. 2013; Kännaste et al. 2013). Variation in the scent of plants can be induced by different stresses, both biotic and abiotic, and can provide detailed information on plant physiological condition and phenology (Takabayashi et al. 1994; Kuhn et al. 2004; Loreto & Schnitzler 2010). Furthermore, the severity of both biotic and abiotic stresses can modulate the intensity of VOC emissions (Toome et al. 2010; Brilli et al. 2011; Copolovici et al. 2011; Niinemets et al. 2013; Opris et al. 2013), implying that stress alters both the plant scent bouquets and intensity.

Volatiles are transported from the emitting to the receiving organisms in air currents, and upon arrival and reception by the receiver organisms they affect numerous ecological processes. The dilution of volatiles in air clearly modulates the distance over which they can act as effective signals. However, the composition of the air that carries volatiles and its direct impacts on the emitter plants and on their volatile emissions have long been overlooked. Several studies have investigated the quantitative effects of atmospheric pollutants on generation of oxidative stress and induction of plant volatile emissions (Vuorinen et al. 2004; Beauchamp et al. 2005), while additional work has described the degradation of volatiles by oxidizing pollutants, and have addressed the question of how that process can reduce signalling efficiency (Himanen et al. 2009; McFrederick et al. 2009; Pinto et al. 2010).

In this review, we take an approach based on the idea that emission of VOCs constitutes a form of ‘plant language’. We outline some of the volatiles induced by oxidizing pollutants, the phenomenon of degradation of volatiles by atmospheric pollutants, and the known impact of the process on volatile-mediated interactions. We go on to examine how differences in the reactivity of volatiles within a blend may result in evolution of a volatile signal. We also examine how the responses of plants to oxidizing pollutants overlap with those that are regulated by volatiles in the process of inter- and intra-plant signalling. Additional insight as to the implications of atmospheric pollutants on chemical ecology will be provided, with particular focus on Brassicaceae and their associated community. We argue that for gaining full mechanistic insight into plant–plant and plant–insect interactions, understanding volatile signal reactions in the ambient atmosphere is of paramount significance.
PLANT LANGUAGES

When thinking of VOCs emitted by plants and their ecological functions, we can liken the emission of volatiles to a plant ‘language’. Similar analogies have been coined previously, such as the ‘cry for help’ synonym for indirect defence, which is the attraction of herbivore natural enemies to herbivore-damaged plants (Dicke et al. 1990; Dicke 2009). The phenomenon of ‘talking trees’ also draws upon a linguistic analogy to describe plants altering their defences upon exposure to volatiles from biotically stressed neighbours (Baldwin & Schultz 1983; Rhoades 1983; Fowler & Lawton 1985). The intricacies of such interactions have been subject to immense scrutiny from ecologists and evolutionary scientists. The degree of ‘speaking’ and ‘listening’ and the capabilities of plants to do both constitute a focal issue of contemporary chemical ecology and chemical biology (Dicke et al. 2003; Baldwin et al. 2006; Paschold et al. 2006; Dicke & Baldwin 2010).

The intended recipient of volatile signals is also a critical issue from an evolutionary point of view. This is particularly true for plant–plant interactions, for which it has been posited that between-plant interactions – the original ‘talking trees’ hypothesis – are actually an example of eavesdropping, whereby the neighbouring plants are responding to a message meant for a different recipient (Karban & Baxter 2001; Karban & Maron 2002). Even when merging these few simple analogies, it starts to become clear that plant languages are complex. To this end, plant defence and the interactions among plants, neighbouring plants and the multitude of cohabiting arthropods have been the focus of numerous studies and reviews (Dicke 2009; Kant et al. 2009; Dicke & Baldwin 2010; de Rijk et al. 2013). VOCs have been shown to be either attractive or repellent to arthropods foraging for food or hosts. These may be herbivorous, predatory, parasitic or pollinating arthropods, which may all utilize VOCs as cues for locating their required feeding resource, prey or oviposition target, but can also serve for avoidance of inappropriate or hazardous situations (Turlings et al. 1990; Vet & Dicke 1992; Oluwafemi et al. 2011; Kessler et al. 2013). Plant-emitted VOCs may also signal to vascularly distant parts of the same plant, resulting in systemic responses (Frost et al. 2007; Heil & Silva Bueno 2007; Niinemets et al. 2013), and in doing so warn of impending attacks to the neighbouring plants, which may eavesdrop on the signal (Baldwin & Schultz 1983; Rhoades 1983; Karban & Maron 2002; Karban et al. 2006; Kost & Heil 2006). Yet, an important aspect often overlooked is that plants in a community form their specific microclimate characterized by greater humidity and milder temperatures (Niinemets & Anten 2009; Kegge & Pierik 2010; Gommers et al. 2013). Induced defences in eavesdropping receiver plants make these plants more resistant to ongoing and future attacks. Thus, signalling to neighbours can play an important role in preserving the integrity of the vegetation and reducing the risk of more severe abiotic stresses in the community, implying that ‘eavesdropping’ can benefit the signal sending plants as well.

Complexity of ‘language’ versus ‘loudness’ of talk

To expand upon the language analogy, we may think of the different compounds emitted by plants as words and the blend of compounds emitted constituting sentences. In general, plants emit compounds belonging to a few ubiquitous compound classes such as terpenoids, benzenoids, aliphatic alcohols and aldehydes (Pichersky & Gershenzon 2002; Niinemets et al. 2013). On the other hand, individual plant species frequently emit volatile blends that are specific to the given species, and often specific to the stress acting upon it (Bruce et al. 2005; Niinemets et al. 2013). The ratios of compounds in the blend provide sufficient information for herbivorous insects to locate their host plants within complex environments (Bruce et al. 2005), while individual components of a blend may constitute non-host cues if other blend components are not present (Webster et al. 2010). Furthermore, the ‘loudness’ of the talk, the magnitude of the emissions induced in response to stress, is often quantitatively linked to the severity of the stress, for example, to the amount of leaf area consumed by herbivores (Copolovici et al. 2011; Niinemets et al. 2013) or damaged by pathogens (Steindel et al. 2005; Toome et al. 2010; Niinemets et al. 2013). As well as the ratios of compounds, the qualitative blend of VOCs can be rather specific with some compounds common to almost all plants, while others may be specific to one or few related taxa (Pichersky & Gershenzon 2002; Karban 2011; McCormick et al. 2012). However, species-specific constitutive blends of volatiles may have substantial variation in the ratio of dominating compounds when populations of a larger geographical scale are compared (Semiz et al. 2007), and the responses of the blends induced by elicitors can also widely vary (Semiz et al. 2012), collectively constituting a large evolutionary pool for stress response and adaptation. In the case of the Brassicaceae, a range of volatile glucosinolate breakdown products are emitted when plants are damaged (e.g. Gols et al. 2009). Isothiocyanates are one such breakdown product and represent a major component of the odour characteristic of damaged brassicaceous plants. Individual isothiocyanates, and particularly 3-butenyl isothiocyanate, have been shown to attract aphid parasitoids without the presence of other compounds (Blande et al. 2007a). Clearly, there is a huge variability in the herbivory-driven volatile blend composition and their emission rates within and among species, and this is further accompanied by important co-evolutionary modifications in receiver organisms.

Signal fidelity and longevity

Two critical issues relating to the efficiency of communication between one organism and another are the fidelity and longevity of the signal (Fig. 1). The majority of previous studies reporting the responses of arthropods and plants to VOCs have focused on emitter plants subjected to either no controlled stress or else to single stresses (Glinwood et al. 2009; Holopainen & Gershenzon 2010). However, natural environments are exceedingly more complicated, incorporating
multiple biotic and abiotic factors that act simultaneously on plants and have the potential to induce blends of VOCs that are different in composition to those induced by the individual stressors (Niinemets 2010b; de Rijk et al. 2013). The effect of multiple stresses may be to induce different volatile blends, with the consequence being that the ‘listening’ arthropod or plant community will respond differently to volatile blends from plants stressed by multiple factors compared with the response to the volatiles induced by each stress occurring individually (Dicke et al. 2009; Zhang et al. 2009; Ponzio et al. 2013). There may also be an overlap in the volatiles induced by different stresses. It has recently been shown that signalling in Arabidopsis in response to egg deposition by Pieris brassicae is similar to that triggered by recognition of pathogen-associated molecular patterns (Gouhier-Darimont et al. 2013), which highlights a common defence response to multiple biotic factors. Modification of induced volatile blends constitutes one way by which abiotic factors may critically disturb volatile-mediated interactions. Induction of volatile emissions by oxidizing pollutants, modification of not-yet-airborne volatiles by oxidants within tissues, and degradation of airborne volatiles by oxidants in the atmosphere are potentially co-occurring effects of the abiotic environment. In the following, each of these areas will be considered in terms of how they affect a volatile blend and the related interactions between organisms.

**ABIOTIC STRESS – VOLATILE INDUCTION**

Plant responses to environmental factors

Typically, abiotic stresses, especially long-term sustained stress events, weaken plants and can make them more vulnerable to any subsequent or simultaneous stress such as pathogen or herbivore attack (e.g. Niinemets 2010b). A range of abiotic factors are known to have significant effects on the volatile emissions of plants. Among the key environmental and stress factors, drought, humidity, light intensity and quality, ozone, CO₂, temperature and nutrient availability all have some impact on the volatile emission dynamics, or on the ratios of compounds in a volatile blend (Gouinguéné & Turlings 2002; Pinto et al. 2010; Staudt & Lhoutellier 2011). These factors may also impact on plant quality, which can affect herbivore performance and also the performance of predators and parasitoids. Consequently, the abiotic environment has enormous potential to interfere with multitrophic interactions, including those mediated by volatiles, blends of which can be altered in plants subjected to oxidative stress (Blande et al. 2007b). Some abiotic factors may be classified as stresses, such as enhanced levels of ozone, drought or flooding, while others are more difficult to classify based on their effects. Humidity and increased CO₂ are environmental factors, which may alter the metabolism of a plant, but do not necessarily constitute a stress. Even enhanced UV-B, long thought to be a plant stress, may not be as unequivocally stressful as previously postulated, a hypothesis that is presented in recent reviews (Hideg et al. 2013; Wargent & Jordan 2013). Therefore, both stressful and other abiotic factors affecting plant metabolism, especially volatile metabolism, should be considered as obstacles to the fidelity of volatile-mediated interactions. In the following, we consider the effects of some prevalent air pollutants on plant volatile emissions, but recognize the significance of other facets of the abiotic environment as drivers of volatile emissions. Previous reviews have provided detailed accounts of the roles of abiotic factors as inducers and regulators of volatile emissions and should be referred to for additional information (e.g. Niinemets et al. 2010a,b; Grote et al. 2013).

**Impact of key air pollutants on volatile emissions**

Air pollutants are mostly chemical compounds of anthropogenic origin that affect biotic systems due to their enrichment in areas of human activity. Many of the air pollutants also have natural origins (e.g. sulphurous compounds released during volcanic activity, CO₂, and oxides of nitrogen and methane released as a result of microbial activity) and participate in normal atmospheric processes. Higher concentrations of these compounds are often a result of human activity, but they can be dispersed to remote areas or undergo atmospheric reactions to form breakdown products or secondary pollutants such as in the photochemical formation of ozone from oxides of nitrogen (Sitch et al. 2007).

Ozone has been shown to both induce and reduce volatile emissions from different plant species depending on the severity and duration of exposure (Calfapietra et al. 2013). Exposing Brassica napus plants to 100 nmol mol⁻¹ ozone reduced the emission of two monoterpens, sabine and δ-3-carene (Himanen et al. 2009), but this had virtually no effect on the tritrophic interaction incorporating Plutella xylostella as a herbivore and the parasitoid Cotesia vestalis. Hybrid poplar (Populus deltoides × maximowiczii) clones...
have been shown to have increased emissions of hydrocarbons and oxygenated sesquiterpenes upon exposure to 80 nmol mol$^{-1}$ ozone for 5 h $^{-1}$ for 10 d (Pellegrini et al. 2012). In a hybrid aspen clone, Populus tremula L. × P. tremuloides Michx., moderate increases in ozone of 1.3–1.4 times the ambient level induced an increase in total monoterpane emissions (Blande et al. 2007b). On the other hand, strong elicitation of emissions of stress volatiles, including green leaf volatiles (GLVs), methyl salicylate and sesquiterpenes, has been observed in tobacco (Nicotiana tabacum) (Beauchamp et al. 2004, 2005). These emissions were quantitatively associated with ozone dose according to a threshold-type response, characterized by moderately elevated emissions at lower ozone doses and massively enhanced emissions when a certain stress threshold was exceeded (Beauchamp et al. 2005). Such quantitative responses can serve as a valuable resource for constructing induced emission models quantitatively linking stress severity and induced emission responses (Niinemets 2010a; Grote et al. 2013; Niinemets et al. 2013).

Nitrogen oxides – or NO$_X$ – are chiefly comprised of nitric oxide, NO, and nitrogen dioxide, NO$_2$. NO$_X$ are major components of vehicle exhaust fumes and are involved in the reactions leading to the formation of tropospheric ozone. However, vegetation, especially under stress conditions, can also release NO (Wildt et al. 1997; Copolovici & Niinemets 2010). Exposure to NO$_X$ and NO in fumigation studies showed that there can be variable effects on plant growth depending upon species (Bell et al. 2011). However, there is very limited information on the effects of NO$_X$ on volatile emissions of plants. There is indication that NO might induce emissions of some terpenoids from Lima bean in exposure studies, but the degree of induction was quite low (Souza et al. 2013). Fumigation of plants with NO prior to oxidative stress induced by either ozone or singlet oxygen generated by rose bengal resulted in reduced emission of lipoxygenase (LOX) compounds produced by the octadeecanoid pathway and reduced levels of hydrogen peroxide (H$_2$O$_2$) and malondialdehyde (MDA), which are indicative of oxidative stress (Velikova et al. 2008; Souza et al. 2013).

The above examples are non-exhaustive, but highlight that atmospheric pollutants can induce plants to emit volatile compounds, whereas the intensity of the emissions and emission composition importantly depend on past stress history and stress severity. Although the complexity of abiotic influences makes evaluation of abiotic stress effects difficult, we argue that the quality and quantity of these induced volatiles do have an effect on the volatile-mediated interactions between different biotic components of a plant-based community.

**Overlap of abiotically induced volatiles and biotically induced volatiles**

The volatiles emitted by plants in response to abiotic factors may constitute either informative cues to other community members, or alternatively may act as a false signal to receivers searching for a particular resource. Under certain abiotic stress conditions, plants may become a better quality host for herbivores than under regular conditions. An example is leaves that have accelerated senescence caused by excess water stress, which in silver birch (Betula pendula) were observed to be colonized more by aphids than the non-senescing green leaves (Holopainen et al. 2009). The reallocation of nutrients during the senescence process could be the reason for the preferences displayed by the aphids (White 1993; Holopainen et al. 2009), but it could also be that reduced defence in the yellowing leaves underlies the observation. It is likely that in this case, visual cues are utilized by the aphids in the foraging process as yellow coloration is known to be much more attractive than green coloration to numerous aphid species (Debarro 1991; Döring et al. 2009). However, if volatiles induced by the abiotic stress are perceived by aphids, they may also constitute a useful cue to the surrounding community. Alternatively, volatile profiles that closely resemble those induced by herbivores – but with no herbivores present – could be misleading to predators and parasitoids searching for prey or hosts. Certain abiotic and biotic stresses induce plants via the same phytohormonal pathways. In fact, there is a convergence of various stress pathways at the level of oxidative signalling (Fujita et al. 2006; Mittler 2006; Koornneef & Pieterse 2008), resulting in development of induced resistance that is effective against a broad variety of attackers (Koornneef & Pieterse 2008). For example, ozone and aphid feeding both induce the salicylic acid pathway, which may lead to overlap in stress-induced volatile blends. Furthermore, both ozone (Beauchamp et al. 2004, 2005) and aphid feeding (Blande et al. 2010b) also induce emissions of sesquiterpenes, further underscoring the overlap of stress pathways. In addition, exposure to ozone at concentrations sufficient to cause visible symptoms on leaves and exposure to spider mite feeding both induce the emission of GLVs and homoterpenes in lime bean, but emission of the monoterpenic β-cimene was only induced by spider mite feeding (Vuorinen et al. 2004). When exposed to both stresses simultaneously, there was a direct additive effect in stress-induced volatile blends (Vuorinen et al. 2004).

In addition to the overlap between volatiles induced by abiotic factors and those induced by herbivores and other biotic agents, a crosstalk between phytohormone-regulated pathways induced by these different agents could result in novel blends of compounds that are different from those induced when either of the factors acted independently. Interestingly, the changes in volatile emissions may not always be a cumulative response to an increasing stress load, but may be indicative of a reduction in stress. This process is known as a cross-stress tolerance, in which prior exposure to one stress provides a degree of resistance to another. As mentioned above, exposure to NO and ozone in sequence results in a lower volatile emission and less oxidative stress than exposure to the individual pollutants in isolation (Souza et al. 2013). However, it is sometimes important which stress comes first (Zhang et al. 2009; Niinemets 2010b), and the effect can also be significantly altered by the timing and severity of the subsequent stress (Niinemets 2010b). Any degree of cross-stress tolerance involving abiotic factors and herbivore feeding

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could affect the volatile emissions of the plant and again impact on the fidelity of volatile-mediated interactions.

However, sometimes, sequential stresses can act almost independently. For instance, ozone exposure of bean (*Phaseolus vulgaris*) did not alter the vulnerability to *Botrytis cinerea* infections (Tonneijck 1994). Furthermore, effects of simultaneous exposure to multiple stresses are generally additive or even interactive, but again the effects may be importantly determined by past stress history resulting in either priming or exhaustion of plant defences due to reduction of soluble and storage carbohydrate pools (Richardson *et al.* 2004; Myers 2005; Niinemets 2010b). Especially, sustained exposure to some abiotic stresses such as drought may predispose plant stands to insect damage or fungal attacks (Schoeneweiss 1983; Appel 1984; Mattson & Haack 1987; Wargo 1996; Bigler *et al.* 2007).

### THE ECOLOGICAL EFFECTS OF VOLATILE UPTAKE AND DEGRADATION

Once volatiles have been released from plants, the plant’s control over those volatiles is effectively over. The volatiles have numerous potential fates based on their reactivity with atmospheric pollutants and their degree of volatility. In the following, we focus on the uptake of volatiles by vegetation, degradation of volatile compounds by pollutants, the effects of volatile degradation on volatile-mediated interactions, and some of the biological and ecological effects of degradation products.

**Uptake, modification and release of volatiles by vegetation**

The passage that volatiles take after emission from the plant takes them first to the boundary layer, or the leaf–atmosphere interface (Fig. 2). The physical and chemical properties of this zone, which envelopes the aerial parts of plants, have been assigned particular importance to exchange of chemical signals between organisms (Riederer *et al.* 2002). The vapour pressure of a compound largely governs the passage of compounds from the plant into the boundary layer, while volatility, lipophilicity and stomatal opening affect whether volatiles will remain in the boundary layer, be taken up by the plant either through stomata or diffusion, or enter the turbulent atmosphere (Riederer *et al.* 2002).

Semi-volatile compounds, such as sesquiterpenes, are more likely to remain in the boundary layer or to be readsorbed to the surfaces of vegetation according to equilibrium partition coefficients (Noe *et al.* 2008; Fowler *et al.* 2009; Himanen *et al.* 2010; Niinemets *et al.* 2011), where they may have further active roles in between-species interactions (Himanen *et al.* 2010).

Abiotic factors also have a decisive role in the route that volatile emissions will take. Humidity affects the deposition of atmospheric constituents such as ozone to surfaces, which may react with volatiles within the boundary layer (Altimir *et al.* 2006), while temperature is a critical factor in the volatility of chemical compounds (Copolovici & Niinemets 2005).

Sesquiterpenes are known to be ‘sticky’ compounds that adsorb to surfaces at low temperatures and are re-released as temperatures increase (Schaub *et al.* 2010). This adsorption phenomenon has been directly linked to the observations of associational resistance in birch (*Betula* spp.) neighbouring *Rhododendron tomentosum* (Himanen *et al.* 2010) and may also have a role in recent observations of associational resistance to oviposition by *Spodoptera littoralis* in cotton (*Gossypium hirsutum*) and alfalfa (*Medicago sativa*) plants adjacent to *S. littoralis*-damaged cotton plants (Zakir *et al.* 2013). It is clear that there is great potential for different abiotic factors to alter the atmospheric behaviour of volatiles, and in doing so to contribute to a change in volatile-mediated processes. Indeed, the adsorption of sesquiterpenes could change the mode of action of the molecule from being a component of a volatile blend to be perceived by foraging predators and parasitoids into a plant surface constituent that may have an impact on host selection by herbivores and their subsequent feeding.

Uptake of volatile compounds into plants could have a role to play in the process of between-plant signalling. One of the most intriguing molecules with respect to interplant signalling is methyl jasmonate (MeJA), which has a key role in regulation of plant defence responses. MeJA has been shown to be taken up by plants and metabolized into jasmonic acid and jasmonoyl-isoleucine, and to a lesser extent jasmonoyl-leucine (Tamogami *et al.* 2008). These metabolic conversions lead to the activation of VOC products, MP denotes products of metabolism, MP-VOC denotes volatile products of metabolism, and the SOA drawing represents secondary organic aerosol. The lines with round ends represent adsorption to surfaces. The line with a diamond at the end represents VOC uptake.

Figure 2. Transport, reaction, deposition and metabolism of volatiles after emission from plants. Volatile emission begins with the red arrow and movement of volatiles is represented by the dashed blue arrows. Reaction of volatile organic compounds (VOCs) with oxidants is represented by the line and plus sign. Solid line arrows represent resulting products, RP denotes reaction products, MP denotes products of metabolism, MP-VOC denotes volatile products of metabolism, and the SOA drawing represents secondary organic aerosol. The lines with round ends represent adsorption to surfaces. The line with a diamond at the end represents VOC uptake.
emissions, which are part of a plant’s indirect defence response. In addition to uptake of signalling molecules, such as MeJA, plants may take up other volatile compounds, which may be re-emitted or transformed into other by-products. This process has been explored with respect to phytoremediation of indoor air by common houseplants, with aldehydes and ketones shown to be taken up (Tani et al. 2007; Tani & Hewitt 2009). The catabolism of volatiles in plant and emission of reaction products can have a significant impact on the composition of volatiles emitted. VOC catabolism and degradation can affect a range of processes including plant C balance, tolerance to environmental stress, plant signalling and plant–atmosphere interactions (Oikawa & Lerdau 2013). However, the effects on foraging arthropods have not been studied extensively. Clearly, this is an important gap that needs filling in the near future.

Degradation of herbivore-induced plant volatiles (HIPVs) and effects on volatile-mediated interactions

Oxidizing pollutants and atmospheric constituents, including ozone, OH radicals and NO3 radicals, can react with VOCs in the atmosphere and impact the dynamics and fidelity of interactions between volatile-emitting plants and the volatile-receiving community (Pinto et al. 2007a, 2010) (Fig. 1). Nitrate radicals (NO3) formed by oxidation of NO2 by O3 are considered to be the most important night-time oxidant of VOCs (Monks 2005). NO3 radicals can initiate, but not catalyse, removal of VOCs from the atmosphere, which leads to removal of NO3 radicals in the presence of highly reactive VOCs. In daytime reactions, the abundant OH radicals catalyse several atmospheric reactions leading to the rapid removal of numerous volatile compounds (Atkinson & Arey 2003; Monks 2005). Changes to the volatile blend can potentially render the ‘sentence’ conveyed by VOCs distorted or indecipherable to some or all recipients in the community. In this situation, the altered blend may just be inactivated, or else it may provide different information to that of the volatile blend actually emitted by the plant. The reaction products formed in atmospheric reactions can be extraordinarily diverse, with ozonolysis of limonene alone shown to result in nearly 1200 different organic compounds, although with some having very short lifetimes (Kundu et al. 2012). In a recent review (Holopainen & Blande 2013), the various fates of VOCs after release to the atmosphere and the potential for atmospheric oxidants to degrade a volatile signal and thus confer temporal and spatial dimensions upon a volatile blend were outlined, underscoring the importance of the aerial environment in signal propagation and composition.

The majority of studies addressing the effects of oxidizing air pollutants on volatile-mediated interactions have focused on the role of tropospheric ozone, with a number of the most widely studied volatile-mediated interactions shown to be reduced in efficiency by elevated ozone levels. The process of indirect defence, whereby herbivore-damaged plants emit volatiles that attract natural enemies of the herbivores, has been shown to be reduced in efficiency in a system comprising cabbage (Brassica oleracea), P. xylostella and the foraging parasitoid C. vestalis (Pinto et al. 2007b), while a further study from the same group (Pinto et al. 2007c) showed no disruption of foraging in the same system, nor in a system comprising lima bean (Phaseolus lunatus), two-spotted spider mites (Tetranychus urticae) and the foraging predatory mite (Phytoseiulus persimilis), possibly due to the degree of volatile degradation not being sufficient to completely eliminate foraging behaviours in laboratory-based tests. Gate et al. (1995) conducted one of the earliest studies on the effects of air pollutants on the searching behaviour of parasitoids using a system comprising Drosophila subobscura larvae and the braconid parasitoid Asobara tabida. They examined the efficiency of host searching by A. tabida for differing densities of hosts feeding on an artificial diet and the effects of three different pollutants, ozone, NO2 and sulphur dioxide (SO2), on parasitism rates and searching efficiency. The key observation was that ozone significantly reduced searching efficiency, while there also appeared to be a negative effect of NO2 on the abilities of parasitoids to distinguish between patch sizes of their hosts. Interestingly, Himanen et al. (2009) observed that ozone affected the orientation of C. vestalis searching for P. xylostella feeding on transgenic Brassica napus ssp. oleifera containing a synthetic Bacillus thuringiensis (Bt) cry1Ac gene, but had no effect on non-transgenic lines.

The host-finding behaviour of the striped cucumber beetle (Acalymma vittatum) has also been shown to be reduced in efficiency by ozone (Fuentes et al. 2013). This herbivore feeds on the foliage, flowers and pollen of cucurbit crops, and is attracted by the floral volatiles (Andrews et al. 2007). However, mixing those volatiles with 80 nmol mol−1 ozone appeared sufficient to disrupt orientation towards the floral volatiles in Y-tube bioassays. Beyond the effects of ozone on plant–arthropod interactions, ozone has also been shown to reduce the distance of communication between plants (Blande et al. 2010a), with a concentration of 80 nmol mol−1 having a significant effect in reducing volatile-mediated signalling between lima bean plants in chamber experiments. The value of 80 nmol mol−1 has been identified as important in two laboratory studies of different systems and using different methods, which is suggestive of it being a potential threshold value that results in rapid degradation of biologically important compounds in the plant volatile blends. It is feasible that under field conditions with greater volatile dilution in air currents that lower ozone concentrations have a similar impact.

In addition to degradation of plant volatiles, ozone also degrades the aggregation pheromone of Drosophila melanogaster with negative effects on the attractiveness of the pheromone (Arndt 1995), which emphasizes the broader biological consequences of increasing concentrations of atmospheric pollutants.

While the majority of past studies on the effects of pollutants on volatile-mediated interactions have focused on the effects of ozone, a recent study by Girling et al. (2013) concentrated on the effects of diesel engine exhaust fumes on the degradation of volatiles in the blend emitted by oilseed rape (B. napus) flowers and the responses of the honeybee Apis

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mellifera to intact and modified blends. They found that exhaust fumes rapidly and extensively degraded the volatile blend, with two terpenes, α-terpinene and α-farnesene, subject to particularly rapid degradation. The fractions of the exhaust fumes responsible for the degradation were primarily NO\textsubscript{2} and NO, which at concentration ratios of 1 μmol mol\textsuperscript{-1}: 10 μmol mol\textsuperscript{-1} and 10 μmol mol\textsuperscript{-1}:10 μmol mol\textsuperscript{-1} resulted in substantial degradation of both terpenes. By conditioning the proboscis extension reflex of honeybees to the intact volatile blend, the authors were able to determine that removal of the two reactive terpenes from the volatile blend significantly reduced the ability of honeybees to recognize the odour. This could have profound effects on the abilities of honeybees to utilize volatile cues in urban areas, while the subsequent reactions between NO\textsubscript{x} and VOCs will, in the presence of sunlight, lead to the formation of ground-level ozone which could further degrade volatiles with further ecological implications. It should be noted that the products of reactions between the terpene and NO\textsubscript{x} were not included in the assessment of odour recognition by Girling et al. (2013), which should be explored further in future studies. The need to consider the ecological effects of both primary and secondary pollutants is emphasized by this work and should be a key future area of research.

**Biological effects of plant VOC degradation products**

As already mentioned, we are unaware of studies that have directly elucidated the effects of VOC degradation products on insects and other herbivorous arthropods. However, some first-phase reaction products of VOCs are known to be synthesized directly by various organisms, and their biological functions could hint at the potential biological effects of the same compounds forming after atmospheric transformation of plant emissions. The isoprene oxidation product methacrolein is emitted by mechanically damaged sagebrush (Artemisia tridentata) plants, and this compound has the capacity to prime trypsin proteinase inhibitor (TPI) activity and reduce herbivory in nearby Nicotiana attenuata plants (Kessler et al. 2006). Another isoprene degradation product, formaldehyde, when encapsulated in fertilizer pellets and released from soil, has been shown to reduce activity of herbivorous slugs (Schuder et al. 2006). In contrast, addition of formaldehyde to artificial diet of moth larvae has been shown to stimulate their growth (Assemi et al. 2012).

Some monoterpane oxidation products are known to elicit biological responses in herbivores. A common monoterpane of conifers and many other plants, α-pinene, is oxidized in atmospheric reactions to form several major first-generation gas or particle phase products such as verbenene, pinic acid, ionic acid and pinonaldehyde (Lee et al. 2006). Verbenone and pinonaldehyde were detected in the secondary aerosol particle mass of conifer forests (Rissanen et al. 2006), but pinonaldehyde formation is related to higher humidity, and it is detectable mostly during nights (Zhao et al. 2013). Verbenone is also internally synthesized by aggregating bark beetles (Gitau et al. 2013) and is formed through microbial processes of soil comprising a fraction of decomposing α-pinene-rich pine needle litter (Kainulainen & Holopainen 2002). Verbenone can act as a repellent anti-aggregation signal chemical for some bark beetle species (Gitau et al. 2013), when populations become too crowded. So far, it has not been shown that natural atmospheric oxidation of α-pinene emission from damaged trees could lead to reduced bark beetle attack in nearby trees, but synthetically produced and released verbenone has been an efficient repellent for certain bark beetle species (Gitau et al. 2013). Interestingly, a common bark beetle-predating beetle, Temnochila chlorodoria, is attracted to high release rates of verbenone (Fettig et al. 2007), suggesting that some of the atmospheric oxidation products of herbivore-induced monoterpenes could act as multitrophic signalling compounds and be efficient attractants of natural enemies of herbivores.

In early oxidation of the monoterpane limonene, formaldehyde and limonoaldehyde have been detected with minor amounts of formic acid, acetone and acetic acid (Lee et al. 2006). Formaldehyde, formic acid, acetone and acetic acid are also common degradation products of other monoterpenes such as β-pinene, myrcene and 3-carene (Lee et al. 2006). Formic acid is an ant venom and it has acaricidal properties (Boncristiani et al. 2012). Acetic acid is a strong attractant of fruit flies (Drosophila sp.) (Landolt et al. 2012).

Sesquiterpenes are rapidly degraded in the presence of ozone (Pinto et al. 2007c). Key reaction products of ozone-initiated oxidation of β-caryophyllene, a common sesquiterpene, are β-caryophyllonic acid, β-caryophyllene aldehyde and formaldehyde (Zhao et al. 2010). β-caryophyllene aldehyde and β-nocaryophyllone aldehyde are degradation products that were identified in ambient secondary organic aerosol (SOA) samples from a forest site in Southern Finland (Parshintsev et al. 2008). Parshintsev et al. (2008) were able to produce these compounds in experimental ozonolysis reactions of β-caryophyllene in laboratory conditions, but the biological effects of these compounds are unknown. Jaouhi et al. (2003) found small amounts of β-caryophyllene oxide, which is a common biogenic VOC emission (e.g. in Betula) as an ozonolysis product of β-caryophyllene. The formation of β-caryophyllene aldehyde and β-caryophyllonic acid has also been reported in ozonolysis studies (Jenkin et al. 2012). Photo-oxidation and β-caryophyllene degradation in the presence of OH radicals can result in formation of a wide range of tertiary and secondary peroxy radicals (Jenkin et al. 2012), which could cause oxidative stress in various ecological systems. First-phase photo-oxidation products of other sesquiterpenes include formaldehyde from aromadendrene and formic acid from longifolene (Lee et al. 2006). Photo-oxidation of humulene did not produce formaldehyde, but a C\textsubscript{15} ketoaldehyde was detected (Lee et al. 2006).

GLVs are an important group of C\textsubscript{6} compounds that are active in plant–plant and plant–carnivore interactions. Their atmospheric behaviour is not very well known, but GLVs such as cis-3-hexenol and cis-3-hexenyl acetate are among the first compounds to disappear when a herbivore-induced VOC plume is exposed to ozone (Pinto et al. 2007c).
Induction of sesquiterpene and monoterpen emissions by herbivore feeding is generally thought to increase the formation of SOA, especially if the result of feeding is an overall increase in emission rather than a qualitative alteration of the volatile blend (Amin et al. 2012; Mentel et al. 2013). However, strong emissions of GLVs rather complicate matters by suppressing the formation of SOAs, which has been presumed to be through the process of scavenging OH radicals (Mentel et al. 2013). This is at least partly supported by the demonstration that in reaction chamber experiments, cis-3-hexenol and cis-3-hexenyl acetate produce a significantly higher SOA yield when exposed to ozone than to OH radicals (Hamilton et al. 2009). The atmospheric chemistry of VOC degradation is extremely complicated, with many gaps in our knowledge regarding the end products (Kroll & Seinfeld 2008) of oxidation reactions and their ecological effects. However, it is clear that attention should be given to the roles that those reaction products play and the potential effects of adsorbed SOA on interactions between plants and their community.

Modelling the effects of atmospheric reactions on infochemical signals

McFrederick et al. (2008) modelled the oxidation of a few common plant-emitted terpenes using Lagrangian diffusion models and indicated that atmospheric oxidants including ozone, hydroxyl and nitrate radicals would significantly reduce the distance over which volatiles would signal to pollinating insects. The empirical studies mentioned earlier tend to lack information about how combinations of different oxidants act, which is the situation encountered in nature (McFrederick et al. 2009). The modelling route is able to incorporate multiple oxidizing species in making predictions, but generally lacks empirical information of how pollinating insects actually respond to the changes in volatile diffusion patterns.

The potential responses of pollinators to reaction products or more stable volatile compounds are not considered in such a model. Therefore, measuring the responses of insects and plants to volatile signals under conditions incorporating multiple oxidants is a key step for future studies. Experiments involving field-scale enrichment of more than one oxidant have not been conducted to date. Therefore, those experiments conducted with enrichment of individual oxidants, such as by Pinto et al. (2008) in the ozone Free Air Concentration Enrichment (O_3-FACE) facility based in Kuopio, Finland, offer the best reference points available to date as to how foraging insects are affected by the presence of ozone in their short-range foraging for hosts. In the case of the parasitoid C. vestalis foraging for P. xylostella feeding on cabbage (B. oleracea), it seems that moderate enrichment of ozone to 1.5 times the ambient level has little effect on foraging success (Pinto et al. 2008). The effects on longer distance foraging are not known to date. However, such effects might be important under low host and parasitoid population densities, which regularly occur for insects with stochastic population dynamics.

In their review, McFrederick et al. (2009) used the available evidence to estimate the ecological interactions most at threat from different oxidizing pollutants. Many of the interactions listed are still untested, but the observations made since their review include the effects of ozone on the use of volatiles by foraging herbivores (Fuentes et al. 2013) and plant–plant interactions (Blande et al. 2010a). The short distance foraging by herbivores, which was best represented in the Y-tube study conducted by Fuentes et al. (2013), and the short distance between-plant signalling investigated by Blande et al. (2010a) were both thought to be at low risk by McFrederick et al. (2009), and were both significantly affected by ozone concentrations of around the 80 nmol mol⁻¹ level. This suggests that the effects of oxidizing pollutants on volatile-mediated interactions could be more dramatic than earlier thought. It also highlights the need for a holistic view of the effects of ozone on ecological systems, whereby the direct effects of ozone and other abiotic stresses on plants are looked at in tandem with the oxidizing effects of the same agents on the volatiles in the air.

Recent advances in the ecology of plant-plant signalling have indicated a degree of complexity that was not previously expected. In both sagebrush (A. tridentata) (Karban & Shiojiri 2009; Karban et al. 2013) and willow (Salix spp.) (Pearse et al. 2013), receiver plants have been shown to respond more vigorously to volatiles from related plants than to unrelated conspecific plants, which demonstrates a degree of self or kin recognition. The deep mechanics underlying such recognition patterns have yet to be elucidated, but the essential information seems to be somehow encoded in the composition or ratios of compounds in a volatile blend. Therefore, small changes to that blend over short distances may result in considerable alteration to the signal quality and may play a large role in determining the distances over which volatile-mediated plant–plant interactions occur in nature.

OVERLAPS BETWEEN PLANT RESPONSES TO OXIDATIVE STRESS AND PLANT RESPONSES TO VOLATILE SIGNALS

Plant–plant interactions mediated by volatiles are among the most sensitive interactions observed in nature (Blande et al. 2010a). Consequently, they are particularly vulnerable to environmental perturbation. While foraging insects have great potential to learn different odours and associate them with a positive experience (Vet & Dicke 1992; Allison & Hare 2009; Hoedjes et al. 2011; Trowbridge & Stoy 2013), plants do not process information in the same way, and consequently they could benefit from an alternative strategy for coping with the breakdown of important signalling chemicals by ozone. It seems that one strategy may be to have a large degree of commonality in response to herbivore-induced volatiles and ozone itself. Numerous studies have been conducted on responses of undamaged plants to volatiles from a damaged neighbour, with several defensive traits shown to be modified in response to such an exposure. Some of the defensive attributes include an increase in LOX gene transcripts, which has been observed in B. oleracea plants exposed to
HIPVs from conspecifics (Peng et al. 2011) and in tomato (Solanum lycopersicum) and potato (Solanum tuberosum) plants exposed to volatiles from damaged emitter plants (Peng et al. 2005). Genes for the pathogenesis-related proteins PR-1 (Peng et al. 2005), PR-2 (Arimura et al. 2000; Yi et al. 2009), PR-3 and PR-4 (Arimura et al. 2000) have all been shown to be up-regulated in response to exposure to HIPVs. Phenylalanine-ammonia lyase (PAL) is also up-regulated in response to HIPV exposure (Arimura et al. 2000; Peng et al. 2005). Exposure to herbivore-induced GLVs has further been shown to induce emissions of other volatiles including the homoterpene (E)-DMNT and acetylated green leaf volatile derivatives (Ruther & Furstenau 2005; Yan & Wang 2006). In addition, lima bean (P. lunatus) plants exposed to HIPVs have a greater quantity of soluble sugars in extrafloral nectar secretions, which is an indirect defence trait that attracts and arrests natural enemies of herbivores (Kost & Heil 2006). However, the concentration threshold for elicitation of such systemic responses and for how long the systemic elicitation of emissions is preserved is still unclear (see Niinemets et al. 2013 for a discussion). Although the systemic elicitation is silenced after the eliciting signal is extinguished, plants may still maintain a certain memory effect, priming, implying a faster and often also a stronger response upon a subsequent volatile signal or biotic attack (Karban & Niino 1995; Conrath et al. 2006; Heil & Silva Bueno 2007; Ton et al. 2007; Choudhary et al. 2008; Frost et al. 2008).

There is also evidence of many of these HIPV-induced defensive traits being directly induced by exposure to ozone. Such an induction could, to some extent, negate the need to receive a signal from a damaged neighbour in order to fine-tune defensive responses, although the blend of ozone-induced emissions may be different and, accordingly, the response may not necessarily be identical to the response elicited upon exposure to plant stress volatiles. In soybean (Glycine max), LOX activity was doubled by 3 h of ozone treatment, with ozone found to modulate LOX expression and to act on the transcription of the LOX genes (Maccarrone et al. 1992). In tomato (S. lycopersicum), exposure to ozone reduced soluble sugars and free amino acids, and increased expression of PAL and PR-1 (Cui et al. 2012), and such an activation of the shikimic acid pathway is also compatible with the evidence of enhanced emissions of methyl salicylate (Beauchamp et al. 2005). Ozone exposure also triggers increased expression of PR-1 in tobacco (N. tabacum) (Pasqualini et al. 2012). Additionally, exposing rice (Oryza sativa) seedlings to enhanced ozone for 2 d significantly induced PR-5 and two PR-10 proteins (Feng et al. 2008). In a study of the effects of ozone on plant-plant signalling, it was found that a level of 80 nmol mol$^{-1}$ was sufficient to significantly reduce the effective distance of plant to plant signalling via HIPVs, but that at 120 nmol mol$^{-1}$ and higher, extrafloral nectar was secreted in response to ozone exposure, the ozone had effectively switched on one of the main induced defences associated with plant-plant signalling, albeit possibly not as a targeted response to the stress but as a metabolic overflow after stoppage of growth. There is a large within-species variability in ozone tolerance associated with the sensitivity of plants to up-regulation of the salicylic acid pathway (Rao & Davis 1999; Koch et al. 2000; Vahala et al. 2003), and it remains to be tested whether this variation in abiotic stress tolerance is also associated with differences in engagement of defence pathways upon biotic stress. In particular, whether more abiotic stress-resistant genotypes are more resistant to biotic attacks or whether reduced level of induced emissions reduces plant fitness due to reduced capacity to attract enemies of herbivores.

**SUMMARY AND FUTURE DIRECTIONS**

There are numerous routes via which abiotic factors and particularly oxidizing atmospheric pollutants may impact on volatile-mediated interactions. Factors that combine to influence the blend of volatiles emitted by plants and make them deviate from those induced by herbivore feeding could have a direct effect on indirect defence and on plant–plant interactions. Atmospheric oxidants that react with a volatile blend will rapidly alter the strength of the signal to other community members. These factors may operate in tandem, with potentially large consequences for the fidelity and longevity of volatile-mediated signalling. However, there is still a lot to be investigated before we really know the effects of such disturbances on interactions between organisms and on a broader ecosystem function. One particularly important but technically challenging focus is the impact of atmospheric pollutants on foraging by pollinators. Long-distance foraging via volatile cues is particularly vulnerable to the effects of oxidizing pollutants. Therefore, it is important to carefully select systems whereby pollinators utilize volatiles during foraging, and also come into contact with varying levels of pollutants. Although the previous study by McFrederick et al. (2008) offers an excellent indication of how volatiles may be degraded and consequently be reduced in concentration at distances from a point source, there is no consideration of the potential for pollinators to utilize the degradation products, which may be more stable and travel further than their precursor molecules. Future research should focus on understanding the effects of multiple stresses on the information wrapped up in the volatile blends emitted by plants and in how foraging arthropods utilize the emissions directly from plants and their degradation products, as well as in understanding how systemic elicitation by volatiles from stressed neighbours is propagated and how these secondary inductions increase the fitness of receiver and transmitter plants.

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