

REVIEW

published: 19 February 2016 doi: 10.3389/fpls.2016.00135



# Broomrape Weeds. Underground Mechanisms of Parasitism and Associated Strategies for their Control: A Review

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OPEN ACCESS

**Edited by:** Richard S. Winder, Natural Resources Canada, Canada

#### Reviewed by:

Pascal Labrousse, Université de Limoges, France Tadao Asami, University of Tokyo, Japan

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#### Specialty section:

This article was submitted to Crop Science and Horticulture, a section of the journal Frontiers in Plant Science

Received: 07 October 2015 Accepted: 12 January 2016 Published: 19 February 2016

#### Citation:

Fernández-Aparicio M, Reboud X and Gibot-Leclerc S (2016) Broomrape Weeds. Underground Mechanisms of Parasitism and Associated Strategies for their Control: A Review. Front. Plant Sci. 7:135. doi: 10.3389/fpls.2016.00135 Broomrapes are plant-parasitic weeds which constitute one of the most difficult-tocontrol of all biotic constraints that affect crops in Mediterranean, central and eastern Europe, and Asia. Due to their physical and metabolic overlap with the crop, their underground parasitism, their achlorophyllous nature, and hardly destructible seed bank, broomrape weeds are usually not controlled by management strategies designed for non-parasitic weeds. Instead, broomrapes are in current state of intensification and spread due to lack of broomrape-specific control programs, unconscious introduction to new areas and may be decline of herbicide use and global warming to a lesser degree. We reviewed relevant facts about the biology and physiology of broomrape weeds and the major feasible control strategies. The points of vulnerability of some underground events, key for their parasitism such as crop-induced germination or haustorial development are reviewed as inhibition targets of the broomrape-crop association. Among the reviewed strategies are those aimed (1) to reduce broomrape seed bank viability, such as fumigation, herbigation, solarization and use of broomrapespecific pathogens; (2) diversion strategies to reduce the broomrape ability to timely detect the host such as those based on promotion of suicidal germination, on introduction of allelochemical interference, or on down-regulating host exudation of germination-inducing factors; (3) strategies to inhibit the capacity of the broomrape seedling to penetrate the crop and connect with the vascular system, such as biotic or abiotic inhibition of broomrape radicle growth and crop resistance to broomrape penetration either natural, genetically engineered or elicited by biotic- or abioticresistance-inducing agents; and (4) strategies acting once broomrape seedling has bridged its vascular system with that of the host, aimed to impede or to endure the parasitic sink such as those based on the delivery of herbicides via haustoria, use of resistant or tolerant varieties and implementation of cultural practices improving crop competitiveness.

Keywords: integrated pest management, *Orobanche*, *Phelipanche*, parasitism, germination, haustorium, plant recognition, seed bank

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#### INTRODUCTION

The broomrapes are obligate plant-parasitic plants from the genera Orobanche and Phelipanche in the Orobanchaceae family (Bennett and Mathews, 2006; Tank et al., 2006; Joel, 2009). Due to their achlorophyllous nature, broomrapes are constrained to obtain their nutritional resources by feeding off other plants using the haustorium, an organ unique in parasitic plants through which the parasite diverts water and nutrients from the host (De Candolle, 1813; Kuijt, 1969; Musselman and Dickison, 1975; Westwood, 2013). The majority of broomrape species are botanical wonders parasitizing wild host plants in natural ecosystems. However, seven broomrape species, Orobanche crenata, O. cernua, O. cumana, O. foetida, O. minor, Phelipanche aegyptiaca, and P. ramosa have specialized on attacking crops causing trouble in agriculture along Mediterranean, central and eastern Europe, and Asia (Parker, 2009). The crops affected depend on the host range of the broomrape species considered but in general, those in the Asteraceae, Brassicaceae, Apiaceae, Fabaceae, or Solanaceae such as sunflower, oilseed rape, carrot, faba bean, or tomato among many others, sustain the major attacks (Parker and Riches, 1993). The damage induced in the crop by broomrape parasitism differs for each broomrape-host association. In general, parasitized crops suffer from reductions in total biomass at the greatest expense to the reproductive tissue (Barker et al., 1996; Manschadi et al., 1996; Lins et al., 2007). In some crops, the biomass loss equals to that accumulated by the parasite indicating that damage in the crop is directly attributed to the parasitic sink activity (Barker et al., 1996; Manschadi et al., 1996; Hibberd et al., 1998). However, in other broomrapecrop associations the damage induced by broomrape extends beyond assimilate diversion. In those cases, broomrape displays a pathogenic nature promoting disease in the crop mainly through negative effects on the crop photosynthetic machinery and hormonal balance (Stewart and Press, 1990; Mauromicale et al., 2008).

There are not figures based on rigorous data for the total area affected by broomrape weeds (Parker, 2009). Sauerborn (1991) estimated that 16 million ha in Mediterranean and West Asia regions risked being infested. In this regard, France is doing valuable work through the Technical Center for Oilseed Crops and Industrial Hemp, Terresinovia, where a nationwide survey of infested fields is actualized online on real time by the farmers with new cases emerging every year and recently toward new regions such as the French Centre region¹. Several studies suggest that large areas of new territory are at risk of invasion by broomrape (Mohamed et al., 2006; Grenz and Sauerborn, 2007), and in fact, invasions in completely new regions are already emerging in countries such as Spain, UK, France, Algeria, Ethiopia, Egypt, Sudan (Reda, 2006; Babiker et al., 2007; Babiker, 2008; Rubiales et al., 2008; Abu-Irmaileh and Labrada, 2009; Parker, 2014).

Several factors contribute to the fact that broomrape weeds remain an uncontrolled agricultural problem. Control strategies designed for non-parasitic weeds such as cultural and chemical methods do not necessarily achieve the required level of control for broomrape due to its mixed traits as weed and as root parasite. Biological traits in broomrape such as achlorophyllous nature, underground parasitism, the physical connection and growth synchronization with the crop, and the exclusive uptake of resources via crop vascular system rather than from the soil make broomrape control a challenging agricultural task. In addition, the biological similarity between host and parasite characterizing broomrape-crop interactions is higher than in other plant pathosystems, which complicates the development of selective methods to control broomrape, without harmful effect in the crop from which it is feeding (Eizenberg et al., 2006; Hearne, 2009; Yoder and Scholes, 2010; Pérez-Vich et al., 2013).

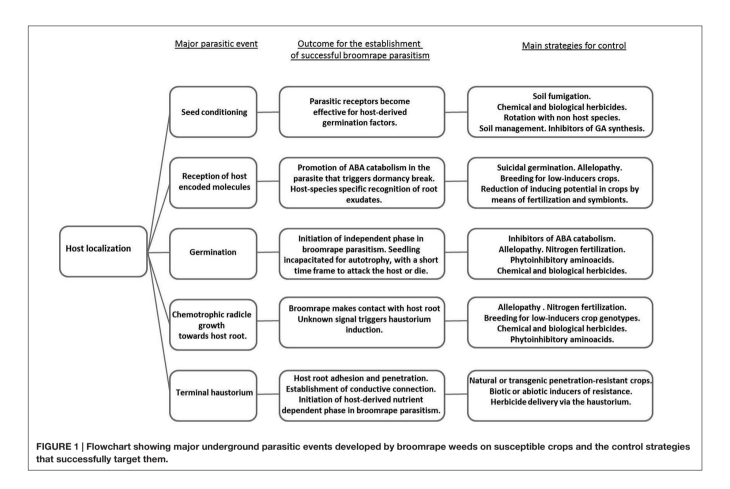
Besides the difficulty of selectively controlling broomrape in the form of host-attached parasite, eradication of broomrape seed bank is extremely difficult due to prolific production of parasitic seeds, their easy dispersal, physiological dormancy, seed longevity, and germination synchronized with specialized range of host cultivation. New infestations can occur through the use of contaminated seeds or machinery and their prevention is essential. Despite of this fact, Seed Certification Services in some of the countries affected, do not include in their certification standards, inspection of crop seed samples for broomrape inoculum. Effective broomrape control should target the underground mechanisms of crop parasitism in order to meet both the short-term productivity expectations of the farmer and reduction of soil bank in the long run (Figure 1). Promising new control strategies have been investigated though the majority of them are under development or remain as prototypes to which farmers have not access. Although some examples of successful control do exist for some crops, the majority of commercially available control methods are either not fully effective or not applicable to many of the affected crops, especially in the case of low-input crops (Joel, 2000).

This paper reviews relevant facts about the biology of broomrape weeds, the key mechanisms they employ to attack crops and the control methods already developed or in development that directly target those mechanisms.

## SPECIALIZED MECHANISMS IN BROOMRAPE WEEDS FOR A PARASITIC MODE OF LIFE

The evolution from autotrophic to heterotrophic mode of nutrition carried a reduction of the main broomrape vegetative organs toward vestigial versions, non-functional for autotrophy. Root system in mature broomrape plants is reduced to short adventitious parasitic roots with functions of anchorage and stabilization in the soil and their leaves are reduced to small achlorophyllous scales (Parker and Riches, 1993). In return they develop haustoria to feed off other plants (Kuijt, 1969; Musselman and Dickison, 1975). The haustorium is the key feature of plant parasitism which has evolved independently at least 11 times in angiosperms (Barkman et al., 2007; Westwood et al., 2012; Yang et al., 2015). Haustorium allows broomrape to attack crops by successive functions, first as host-adhesion

<sup>&</sup>lt;sup>1</sup>http://www.terresinovia.fr/orobanche/carte.php



organ, and subsequently as invasive organ toward host vascular system where finally establishes vascular continuity allowing the parasite to withdraw water and nutrients from the host (Riopel and Timko, 1995; Joel, 2013). The terminal haustorium develops at the apex of the seedling radicle upon host recognition (Musselman, 1980; Joel and Losner-Goshen, 1994). This is a short and delicate stage where the parasite either connects with the host or dies due to nutrient exhaustion. As a consequence of the high risk of establishment failure in the seedling, broomrapes have evolved germination strategies that "predict" establishment potential based on host chemodetection (Vaucher, 1823). In the following sections we describe the key developmental stages in the subterranean broomrape life cycle. These stages constitute sites of broomrape metabolism at which it is possible to design successful strategies to inhibit its sophisticated parasitism.

#### **Host Localization**

### **Seed Conditioning**

Broomrape seed bank remains viable in the soil for many years until germination is triggered by the coincidence of several physical and chemical factors that are indicative of environmental conditions for successful seedling establishment: i.e., the nearby growth of a host plant in a physiological stage susceptible for broomrape invasion and subsequent parasitic reproductive growth (Linke and Saxena, 1991; López-Granados and García-Torres, 1996, 1999). Although hard seed coat has

been described as dormancy mechanism in newly formed broomrape seeds (López-Granados and García-Torres, 1996), water uptake and imbibition are performed quickly by mature seeds through the micropyle without the need of scarification (Bar-Nun and Mayer, 1993; Joel et al., 2012). Broomrape seed bank presents annual cycles of non-deep physiological dormancy induced by seasonal changes in climatic conditions. In non-parasitic plants, physiological dormancy can be relieved through stratification but in the case of broomrape weeds, two consecutive processes are required to release dormancy: an environment-dependent first step of warm stratification called the conditioning phase, and a host-dependent second step of chemodetection. The first step of conditioning promotes in the parasitic seed receptors the required sensitivity for the second step of host detection (Musselman, 1980; Kebreab and Murdoch, 1999; Lechat et al., 2012, 2015; Murdoch and Kebreab, 2013).

The length and temperature required to promote seed conditioning depends on the broomrape species but are usually described under laboratory conditions in a range of 4–12 days at a temperature of 19–23°C, in dark and humid conditions (Kebreab and Murdoch, 1999; Gibot-Leclerc et al., 2004; Lechat et al., 2012). This treatment in the lab mimics the soil conditions in climatically suitable regions for broomrape such as Mediterranean non-irrigated agrosystems where the onset of warm and wet season coincides with the growth of juvenile stages

of many annual crops (López-Granados and García-Torres, 1996; Grenz and Sauerborn, 2007). It is important for broomrape to initiate parasitism in young crops otherwise host reproductive organs in the rapid seed-filling stage will be able to endure a delayed parasitism by establishing a stronger competition with parasitic sinks (Manschadi et al., 1996; Fernández-Aparicio et al., 2009a, 2012a). Therefore broomrape seeds timely gain sensitivity for host chemodetection by means of conditioning (López-Granados and García-Torres, 1996). In absence of host detection the continuation of wet conditions allows broomrape seeds to enter again in deeper levels of dormancy, from which they will emerge upon the new onset of sequenced dry/wet seasons carrying new opportunities to encounter suitable hosts (Kebreab and Murdoch, 1999; López-Granados and García-Torres, 1999). Interestingly, experimentation carried out on broomrape species specialized on summer crops revealed their lower requirement for conditioning when compared with species specialized in winter annual crops highlighting the ecological adaptation of broomrape weeds to the cropping system in which they become specialized (Plakhine et al., 2009).

The metabolic activity of the seed conditioning in broomrape has been characterized in terms of patterns of respiration, synthesis and turnover of proteins, metabolism of nitrogen, carbohydrates and lipids and hormonal balance. Seed respiration patterns during conditioning indicate a strong activation of metabolism. Neither nitrogen nor lipid content change significantly during conditioning, while carbohydrate metabolism and protein synthesis seems to be crucial (Bar-Nun and Mayer, 1993, 2002; Mayer and Bar-Nun, 1994, 1997). It is well-established in autotrophic plants that abscisic acid (ABA) acts as a positive regulator of induction of seed dormancy and its maintenance and gibberelins (GAs) antagonizes with ABA, promoting dormancy release and subsequent germination (Finch-Savage and Leubner-Metzger, 2006). Accordingly, broomrape seed conditioning induces a decrease in ABA levels (Chae et al., 2004; Lechat et al., 2012) and GA synthesis (Joel et al., 1991; Zehhar et al., 2002). The reduction of ABA:GA ratio induced by stratification (conditioning) is enough to break dormancy and promote germination in dormant seeds of non-parasitic weeds but it is not enough for broomrape, which requires a further decrease in ABA levels induced by the activation of the ABA catabolic gene PrCYP707A1 (Lechat et al., 2012). This gene remains silenced during conditioning phase and its activation occurs mediated by host-encoded germination stimulants, i.e., strigolactones, only after the conditioning phase is complete. Gain of host sensitivity in broomrape seeds at the end of the conditioning phase is mediated by demethylation of PrCYP707A1 promoter. Besides the demethylation of PrCYP707A1 promoter required for host-dependent PrCYP707A1 expression, the high levels of global DNA demethylation observed at the end of conditioning period suggest that the epigenetic process occurring during the conditioning phase may be targeting other unknown molecules during conditioning. Whether the demethylation and host stimulation are independent or related processes remains to be clarified (Lechat et al., 2015).

#### Host-Induced Germination

The timing of germination is the most crucial event that obligated parasitic plants face along their life cycle (**Figure 2C**). Their absolute dependence on host-derived nutritive resources for successful seedling establishment and consequent growth makes necessary the synchronization of parasitic germination with the growth of its host. In order to achieve such synchrony they evolved mechanisms that release seed from dormancy triggering germination upon detection of specific molecules contained in host root exudates (Vaucher, 1823).

Broomrape species display high diversity with regard to their host range. Host specificity in broomrape species is usually indirectly related to the predictability of nutritive resources. The predictability of establishment on perennial hosts is high and therefore wild broomrape species feeding off perennial plants have narrow host ranges. On the contrary, weedy broomrape species are usually generalists attacking annual crops (Schneeweiss, 2007). The first mechanism involved in host specialization is displayed during broomrape germination and is mediated by the broomrape recognition of host root exudates in a species-specific manner. For each broomrape-crop association, broomrape germination potential is defined by the combination of both, the stimulatory capability of crop root exudates and the sensitivity of parasitic receptors to recognize specific forms of germination-inducing factors (Fernández-Aparicio et al., 2008a, 2009Ь, 2011).

The stimulatory capability of crop root exudates is defined by the qualitative and quantitative content of germinationinducing factors and varies across crop species and cultivars. In addition it also varies considerably in crops growing under different physiological status, growth stages and growing seasons, allowing broomrape to synchronize its germination with physiologically suitable hosts (López-Granados and García-Torres, 1996; Yoneyama et al., 2007a,b; Fernández-Aparicio et al., 2009b, 2014; Xie et al., 2010). Several classes of germination stimulants have been identified in root exudates such as strigolactones (Xie et al., 2010), peagol and peagoldione (Evidente et al., 2009), peapolyphenols A-C (Evidente et al., 2010), soyasapogenol B, trans-22-dehydrocampesterol (Evidente et al., 2011), dehydrocostus lactone (Joel et al., 2011), or isothyocyanates (Auger et al., 2012). The best studied group of germination-inducing factors are strigolactones, a group of terpenoid lactones. They are exuded by the crop to the rhizosphere under nutrient deficient conditions in order to promote symbiotic interactions (Akiyama et al., 2005). Parasitic plants eavesdrop the plant-to-symbiont communication to sense their hosts and germinate (Xie et al., 2010). Besides their role as extraorganismal signaling, recent research is uncovering new functions for strigolactones as plant hormone controlling crop development in response to the environment (Gomez-Roldan et al., 2008; Umehara et al., 2008). Postgermination development in broomrape could be probably regulated by their own broomrape-encoded strigolactones as it occurs in the close related parasite Striga hermonthica or in non-parasitic plants (Liu et al., 2014; Das et al., 2015). Although analytical chemistry methods have failed to detect strigolactones in parasitic plants (Liu et al., 2014), transcriptome

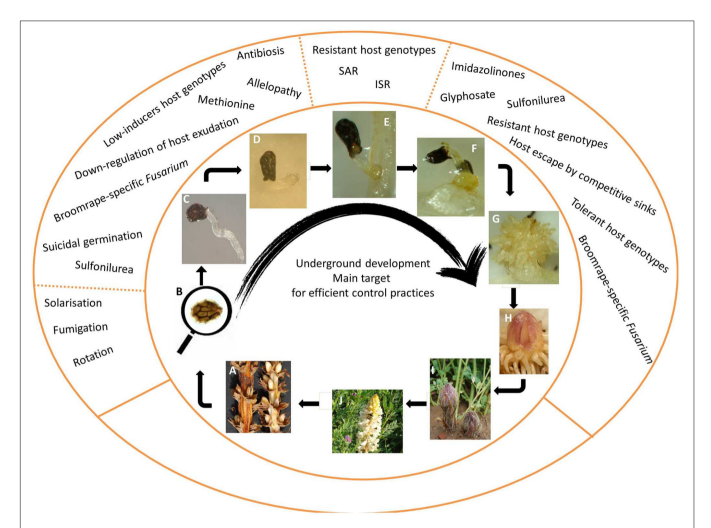


FIGURE 2 | Ilustration of broomrape life stages and mechanisms of control. (A) Fructification and dehiscence of capsules containing mature seeds; (B) microscopic view of a seed (size ranging 0.2–2 mm) that undergoes sucessive dispersal, primary dormancy and annual release of secondary dormancy; (C) broomrape embryo does not develop morphologycaly identified cotyledons or shoot meristem and upon host-induced germination, only a radicle emerges from the seed with the function of searching and contacting the host root; (D) upon haustorial induction, the radicle stops elongating and a single terminal haustorium is differentiated. The first function of haustorium is as adhesion organ to host root surface mediated by a papillae cell layer; (E) adhesion to the root 3 days after germination induction; (F) upon vascular connection with the host, broomape initiates the development of the tubercle, the broomrape storage organ for host-derived nutrients. A swelling of the host root at the penetration point is also observed due the parasitic stimulation of host tissue proliferation; (G) tubercle develops a crown of adventicious roots; (H) tubercle differentiates apical shoot meristem (single shoot meristem for Orobanche species and several shoot meristems for Phelipanche species); (I) the underground shoot eventually emerges through the root surface; (J) flowering and pollination occur. Some broomrape species are outcrossers while others are self-pollinating. Reviewed in Joel et al. (2007).

sequencing reveals that all known strigolactone genes, both synthesis and perception are present in broomrapes with apparently full-length proteins (Péron et al., 2012; Das et al., 2015). The presence of strigolactone biosynthetic system in broomrapes raises the question on how the parasite performs diversified stimulant recognition in order to set the timing of germination. How broomrapes make the distinction not only between host-derived and their own-encoded strigolactones but also how they sense diversified strigolactone profiles in root exudates across species correlated with host ranges. Gene expression analysis could be indicating that parasitic plants down-regulate their synthesis of strigolactones at the end of conditioning period, and perhaps the creation of

that internal deficit for broomrape-encoded strigolactones contributes to the broomrape sensitivity for external, host-derived strigolactones at the time of host detection (Das et al., 2015). In addition, the parasitic-specific receptor KAI2d that enables host detection in broomrapes has recently been identified. Multiple *KAI2d* genes across broomrape species genomes may allow diversified recognition of root exudates corresponding with suitable hosts (Conn et al., 2015). Additional mechanisms that could contribute to the selective action of host-derived strigolactones in broomrape germination could be (1) variations of molecular structure between host-derived and parasite-encoded strigolactones conferring different specificity for different biological functions or (2) different spatial

localization inside the broomrape seed for functions of strigolactone detection and strigolactone synthesis (Das et al., 2015).

The requirement for germination-inducing factors in order to break dormancy in parasitic seeds are bypassed by ethylene or cytokinins (which promotes ethylene biosynthesis) in Striga sp. a close related parasitic weed genus, but these hormones are ineffective in promoting germination of broomrape weeds (Lieberman, 1979; Logan and Stewart, 1995; Berner et al., 1999; Joel, 2000; Toh et al., 2012). GA acts positively on germination in dormant non-parasitic species by counteracting ABA (Seo et al., 2009). However, exogenous application of GA alone is not sufficient to promote broomrape germination (Takeuchi et al., 1995; Chae et al., 2004) and strigolactone-mediated ABA catabolism in conditioned seeds is required to trigger germination (Lechat et al., 2012). A better understanding in the roles of major hormones in the process of broomrape germination would facilitate the design of feasible control strategies based on either inhibition of broomrape germination during crop cultivation or promotion of suicidal germination in the absence of the crop. In order to increase their applicability in low-input crops, the development of synthetic analogs of hormones would constitute a cheap alternative to natural bioregulators for seed bank control of weeds in general and parasitic weeds in particular.

#### Radicle Elongation and Haustorium Differentiation

The embryos in broomrapes have not morphologically identified cotyledons or shoot meristems and upon germination, only a radicle emerges through the seed coat with the only function of reaching and invading the host. The broomrape radicle shows no gravitropism and grows toward the host as a result of cell elongation. It has no root cap and does not develop procambium or conductive tissues (Joel and Losner-Goshen, 1994). Being deprived of the initiation of autotrophic mode of life, the growth of broomrape seedling toward the host is only sustained by water absorption and remobilization of reserve nutrients from the seed perisperm and endosperm (Joel, 2000; Joel et al., 2012). The seedling absorbs water both from the soil and from the seed endothelium, the later ensuring radicle development even in dry soil (Joel et al., 2012). The maximum radicle elongation is limited (1-5 mm) and its viability in the absence of host connection only last a few days after germination has been triggered (Veronesi et al., 2007). This spatial/temporal frame defines the maximum hostreaching distance for successful broomrape parasitism. Upon host detection, the broomrape radicle stops elongating and terminal haustorium is differentiated as an anchoring device. In this process, cellular expansion of the root meristem is redirected from longitudinal to radial and the root apex changes its form from conical to spherical. The external cell layer at the root tip differentiates into a papillate cell layer forming an adhesion epithelium (Figure 2D). The papillae form a crown around the apical cells that remain non-papillate but later will become intrusive cells with an essential function in the penetration process. This surface is covered by carbohydrate secretion that sticks the haustorium to the host surface. This

structure is described as the external anchorage device of the pre-penetrated haustorium to the host surface (Joel and Losner-Goshen, 1994).

Close related parasitic plants of Orobanchaceae such as Striga and Triphysaria use host derived phenolic derivatives to induce haustorium differentiation (Riopel and Timko, 1995; Albrecht et al., 1999; Bandaranayake and Yoder, 2013). Hydrogen peroxide generated by parasitic radicles activates host peroxidases that catalyze the conversion of host cell walls into haustorium-inducing quinones (Keyes et al., 2000, 2007). Haustorium-inducing factors are structurally similar to allelopathic phytotoxins and gene expression of parasitic radicles exposed to haustorium-inducing factors is similar to that after radicle is exposed to phytotoxins (Tomilov et al., 2006). Parasitic plants probably evolved to recruit plant defense molecules as host recognition cues (Atsatt, 1977; Matvienko et al., 2001; Bandaranayake and Yoder, 2013). In broomrape species, the chemistry of host recognition for haustorium initiation remains uncharacterized. They have been traditionally considered the exception in parasitic Orobanchaceae that do not require host factors for haustorium initiation (Joel and Losner-Goshen, 1994; Bandaranayake and Yoder, 2013). However, results recently arisen from a molecule screening identified phytotoxins that induce the development of anchoring device in broomrape radicles (Cimmino et al., 2014, 2015). The significance of this structure in broomrape parasitism requires further investigation. A better understanding of the biochemistry of host recognition in broomrape will facilitate the generation of control strategies targeting the haustorium development.

#### **Host Invasion**

After host adhesion to host root surface the haustorium develops its invasive function of penetrating the host root (Figure 2E). The apical cells in the radicle apex develop into intrusive cells, which successively invade host root cortex, endodermis, and the central cylinder. During the host penetration process, broomrape does not dissolve the host cells in its way toward vascular cylinder. Mechanical force exerted by the haustorium development toward host vascular cylinder combined with enzymatic secretion promotes the separation of host cells without their lysis (Privat, 1960; Ben-Hod et al., 1993; Sholmer-Ilan, 1993; Singh and Singh, 1993; Antonova and Ter Borg, 1996; Bar-Nun et al., 1996; Losner-Goshen et al., 1998; Veronesi et al., 2005). Although broomrape pre-vascular connections benefits from host nutrients, the growth of broomrape in its way toward vascular cylinder is mainly sustained by consumption of seed reserves (Aber et al., 1983; Joel and Losner-Goshen, 1994; Joel, 2000). If the vascular connection is not successfully performed in few days the parasitic seedling dies of inanition and therefore quick invasion of the host is of advantage to avoid loss of viability. When resistant crops impose barriers to stop the parasitic development at this stage, broomrape exhausts and parasitism is quickly aborted. The chemical characteristics of the barriers of resistance to broomrape penetration have been extensively studied in Fabaceae crops (Pérez-de-Luque et al., 2009) and are reviewed in this article in Section "Resistant Crops to Broomrape Invasion"

# Establishment of Vascular Continuity and Parasitic Sink

Shortly after host penetration and connection, the parasite begins its heterotrophic growth at the expense of host resources. Broomrape acts as a strong sink, depriving the host from water, mineral, and organic nutrients with the consequent negative impact on the growth of the host plant (Manschadi et al., 1996; Hibberd et al., 1998; Joel, 2000; Abbes et al., 2009). Few days after host vascular connection, the part of the broomrape seedling that remains outside the host root develops into a storage organ called tubercle. As the tubercle matures a crown of adventitious roots will emerge from this tubercle carrying capacity of developing lateral haustorial connections. Underground shoots will also develop from the tubercles that will eventually emerge through the soil surface leading into the development of reproductive organs (Figures 2F–J).

The transfer of nutrients from host to broomrape is performed through a continuous vascular system at the host-parasite interface. A continuous phloem system between broomrape and its host has been microscopically observed at the terminal haustoria. Sieve elements of both organisms are already interconnected by interspecific sieve pores at early stages of parasitism. It allows the parasite to quickly start tapping carbohydrates, amino acids, and organic acids from its host (Dörr and Kollmann, 1995; Nandula et al., 2000; Abbes et al., 2009). Although host phloem supplies the majority of nutrients including minerals, open xylem connections developed at the host-parasite interface allow additional mineral and water flow toward the parasite (Abbes et al., 2009; Westwood, 2013). These connections are probably developed from simultaneous differentiation of adjacent host and parasite cells to xylem elements (Dörr, 1997). The differentiation of xylem elements in the parasite are under the control of polar auxin transport (Harb et al., 2004; Bar-Nun et al., 2008). During the grafting between host and parasite, broomrape assumes the role of a root, orientating vascular tissues from the host shoot into itself (Bar-Nun et al., 2008). Acquisition of water is driven by a lower water potential in broomrape tissues (Ehleringer and Marshall, 1995). This is maintained by accumulation of solutes mainly potassium at higher concentrations than in the corresponding host tissues (Abbes et al., 2009). Regarding carbon assimilation broomrape takes it from the host phloem mainly in the form of sucrose (Aber et al., 1983; Hibberd et al., 1999). Once in the parasite system, sucrose is not accumulated but metabolized to other compounds. Cleavage of sucrose into glucose and fructose doubles the osmotic potential of the parasite. Sucrose is also metabolized to starch that is accumulated in the broomrape storage organ, the tubercle (Abbes et al., 2009; Draie et al., 2011).

Nitrogen metabolism remains largely unknown in broomrape. Activity of some nitrogen assimilating enzymes has been reported low in broomrapes. Nitrate reductase is not detectable (Lee and Stewart, 1978) and activity of glutamine synthetase is very low (McNally et al., 1983). Broomrape tubercles accumulate host-derived nitrogen in the form of either arginine or in the arginine and aspartate pair (Nandula et al., 2000; Abbes et al., 2009). Besides arginine and aspartate, other major forms of amino acids translocate from the host phloem but they are rapidly utilized by broomrape. Maintenance of relative low levels of those amino acids in tubercles either by low levels of synthetase activities (McNally et al., 1983) and or their rapid turnover of host-derived amino acids, establishes a decreasing concentration gradient that favors the unloading of amino acids into the parasite (Abbes et al., 2009). Despite the reports of broomrape inefficient machinery for nitrogen assimilation and broomrape dependence for host-derived organic forms of nitrogen demonstrated by the fact that broomrape growth is arrested when feeding on host cultivars with decreased amino acid-phloem levels (Abbes et al., 2009), inhibition of enzymes at the top of amino-acid biosynthetic pathway by means of either direct inhibitory action of herbicides (Gressel, 2009) or by feedback inhibition induced by amino-acid end-products (Vurro et al., 2006) are able to kill broomrape. This seems to indicate contribution of amino acid synthesis in broomrape mediated by broomrape-encoded enzymes although their identification and characterization remain unknown (Gressel, 2009; Eizenberg et al., 2012).

Understanding the key processes of host recognition, haustorium development and maturation and metabolic regulation of the parasitic sink allow virulence and the design and implementation of predictions highly calibrated, feasible, and durable control strategies leading to the arrest of broomrape parasitism minimizing simultaneously environmental impact and yield losses.

# CONTROL STRATEGIES TARGETING UNDERGROUND BROOMRAPE STAGES

Successful broomrape control should target the underground broomrapes at their earlier life stages, prior attachment or as soon as it attach to the host, because of their highest vulnerability at those stages and the avoidance of yield loss in the current crop. The majority of strategies aimed to manage autotrophic weeds do not necessarily work for broomrapes and those that provide a degree of success for broomrape need to be optimized for each broomrapecrop species combination, local environmental conditions and broomrape population. Due to the high broomrape fecundity, long seed viability and for some weedy broomrape species, broad host range, the seed bank is easily replenished and long lasting. Therefore an integrated and sustained management strategy composed by several control methods acting at different broomrape life stages is highly recommended to keep away the broomrape weed problem in a durable manner (Kebreab and Murdoch, 2001). The following sections and Table 1 review the major feasible control measures for broomrape control.

TABLE 1 | Methods for Orobanche and Phelipanche spp. control.

	Technique	Broomrape stage targeted	Feasibility	Drawbacks or side effects
Cultural control				
Rotation including	Trap crops Catch crops Allelopathic crops	Seed germination Pre-attached seedling Young attachments	To avoid seed bank replenishing the rotation should avoid  • Susceptible crops  • Broomrape-susceptible non-parasitic weeds	Miscalculation of optimal harvesting time for catch crops leads to broomrape multiplication     Avoidance of susceptible crops that in most cases are the most economically advantageous     Slow process requiring long term management perspective     Risk of widen broomrape host range
Intercropping sus	Intercropping susceptible crops with allelopathic species	Seed germination Pre-attached seedling	Requires appropriate selection of sowing density and non-host component for each broomrape species     Requires greater management skills     Once optimized, rather easy to set up	<ul> <li>Yield uncertainty</li> <li>Partial effect within a growing season</li> </ul>
Fertilization	Nitrogen	Direct effect on seed germination and pre-attached stages Indirect effect on seed bank: negative regulation of host synthesis and exudation of germination-inducing factors	Urea and ammonium but not nitrate forms inhibit broomrape seed germination and radicle elongation  Nitrogen regulates host exudation of germination-inducing factors in some species such as sorghum, lettuce, vetch, and wheat but not in important broomrape hosts such as clover, alfalfa, and tomato	Environmental pollution     Potentially inappropriate Nitrogen supply for the crop species     Lower levels of symbiotic interactions in legume crops
	Phosphorus	Indirect effect on seed bank: negative regulation of host synthesis and exudation of germination-inducing factors	At low or high pH the P solubility is reduced     Requires correct placement of the fertilizer both for easy access to crop root (P is immobile) and to avoid toxic effect to the crop seed (DAP-containing fertilizers produce free ammonia)	<ul> <li>Environmental pollution</li> <li>Limited global resources.</li> <li>Lower levels of symbiotic interactions in crops</li> </ul>
Delayed sowing		Ungerminated seed bank (lower stimulatory capability in root exudates of mature crops growing late in the season) Young attachments (lower parasitic sink strength competing with rapid seed-filling stage of host fruits)	<ul> <li>Requires proper timing to obtain the best balance between inhibition of parasitic sink strength and crop productivity</li> </ul>	Shorter growing cycles not only reduces the amount of biomass partitioned to the parasite but also the overall crop productivity
Physical control	_			
Solarization		Ungerminated seed bank	• Requires high solar radiation	Expensive     Plastic waste     For intercropping periods

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	Technique		Broomrape stage targeted	Feasibility	Drawbacks or side effects
Chemical contro	lo.				
Soil fumigation	Methyl bromide		Ungerminated seed bank	Banned by international agreement	Environmental pollution     Expensive     Labor intensive
	Metham sodium Dazomet 1,3-dicloropropene		Ungerminated seed bank	Tested as substitutes of methyl bromide but they are more expensive and less effective	Environmental pollution     Expensive     Many active ingredients removed from the market for their unacceptable environmental profile under recent legislation     Labor intensive
Soil herbigation	Synthetic herbicides	Sulfonylurea	Seed germination Radicle elongation Young attachments	Requires proper timing and application technology.     Requires a crop tolerant to sulfonylurea residues	Environmental pollution.     High risk of appearance of sulfonylurea-resistance races.
	Inducers of suicidal broomrape germination	Synthetic analogs of strigolactones Fluridone GA agonists	Ungerminated seed bank	<ul> <li>Requires proper timing, application technology</li> <li>Should be applied in absence of susceptible crops in order to avoid further increase of parasitism</li> </ul>	<ul> <li>Though suicidal germination has been proved successful for Striga weed control in USA, it has not reached acceptable levels of seed bank reduction in broomrape</li> </ul>
	Broomrape-specific phytotoxic amino acids	Methionine Lysine	Seed germination Radicle elongation Crop invasion	Green and non-toxic     Requires proper timing, application technology, and tolerant crops     Commercially available at large scale as animal feed supplements     Mixed applications of two inhibitory amino acids can bring synergistic effect on weed control and reduced risk of resistance breakdown adaptation	<ul> <li>Soil pH alteration</li> <li>Difficult application at the required broomrape toxic concentration (5 mM)</li> <li>Technique under experimental development</li> </ul>
	Toxins from microbial origin	Deoxynivalenol Diacetoxyscirpenol Fusarenon X HT-2 toxin Neosolaniol Nivalenol Roridin A T-2 toxin Verrucarrins A, B and M	Seed germination Radicle elongation	Molecules of biological origin are biodegradable in comparison with synthetic herbicides     Requires proper timing and application technology     Requires drip-irrigation system for site-specific weed management and avoidance of toxin spread     Broomrape-specific toxins are preferred	Lack of broomrape selectivity  Lack of knowledge on the physiological mechanism responsible for inhibitory action  Lack of knowledge on the persistence in the soil  Technique under experimental development  Lack of large scale production
Foliar application	Foliar application of systemic herbicides	Glyphosate Imidazolinones Sulfonylurea	Attached parasites (herbicide reaches the parasite via the haustorium)	Requires proper timing and application technology     Requires crops with herbicide-resistance not based on metabolic degradation or inactivation     Where herbicide-resistance varieties are not available within a crop, repeated application at low doses are effective for young parasitic stages     Exact knowledge of the underground phenology related with thermal time is required to predict the best application timing	Marginal crop selectivity     Environmental pollution     Injury of reproductive meristems in some crops     Sublethal concentrations of glyphosate suppress phytoalexin-based immunity in legumes toward pathogens     GM crop paradigm
					(Continued)

	Technique		Broomrape stage targeted	Feasibility	Drawbacks or side effects
Biological control	ıtrol				
l and bacterial pathogens	Solid formulation	Wheat, corn or rice seeds Granular formulation (microbial agent+nutrients)	All stages from un-germinated seed bank to broomrape inflorescences	Risk of low uniform distribution and control efficiency Control efficiency can be increased by "multiple pathogen strategy" Control efficiency can be increased with abiotic elicitors of resistance, e.g., BTH or sublethal herbicidal doses	Allows only one application (at planting)     Moisture is limiting factor     Low uniform distribution and control efficiency     Regularity of the result can hardly be ascertained because of biotic interactions prevailing in the local conditions
einetosd bas ignu	Microbigation		All stages from un-germinated seed bank to broomrape inflorescences	Requires drip-irrigation system for site-specific     weed management     Repeated applications of Fusarium suspensions     provide moisture and optimization of     concentration and timing	Still under scientific evaluation. PGPR area to which this technique is related remains largely unexplored
Broomrape-specific p	Hypervirulence transgenes Pathogens overproducing and excreting phytotoxic amino acids	Enzymes that overcome broomrape defense mechanisms Broomrape-specific toxins	All stages from un-germinated seed bank to broomrape inflorescences	<ul> <li>Increased broomrape kill rate in comparison with their wild forms</li> <li>Enhanced virulence should not after specificity for broomrape weed</li> <li>Spread of transgenic organisms could be avoided with the selection of asporogenic mutants</li> </ul>	Still under scientific evaluation. Techniques remain largely unexplored
lnsects	Phytomyza orobanchia		Avoids seed bank replenishing	Broomrape-specific  Efficiency can be lowered by several agricultural practices and parasitoids  Efficient control requires multiple applications over several years  No artificial diet for rearing is available  Easiness to add to other agricultural controlling practices	<ul> <li>Low impact in a short term seed bank management</li> <li>No impact in the productivity of the current crop</li> </ul>
Symbiotic organisms	Rhizobium leguminosarum Glomus mosseae	ш	Seed germination Radicle growth Crop invasion	• Requires optimal control of fertilization	<ul> <li>Strategy in its early developmental stage</li> </ul>

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	Technique	Broomrape stage targeted	Feasibility	Drawbacks or side effects
Host resistance				
Breeding for natural resistance	Broomrape-resistant varieties	Seed germination Host attachment Crop invasion	Low cost of implementation once developed     Environmental compatible     Varieties with pyramiding gene resistance are preferred to avoid resistance breakdown by the parasite     Resistant varieties should be used in an integrated management strategy to avoid resistance breakdown by the parasite	<ul> <li>Not available for all crops</li> <li>Risk of emergence of new virulent races</li> </ul>
	Broomrape-tolerant varieties	Parasites attached to the host	<ul><li>Feasible</li><li>Environmental compatible</li></ul>	Not available for all crops
	Systemic herbicide-resistant varieties	Young parasitic seedling connected to host vascular system	Feasible if herbicide resistance is not based on metabolic degradation or inactivation     Requires proper timing and application technology	Not available for all crops     Risk of emergence of herbicide-resistant broomrape populations
Mutagenised crops	EMS-mutagenesis Fast-neutron-mutagenesis	Seed germination Host attachment Crop invasion	Low cost of farm implementation of resistant varieties     Environmental compatible	Risk of emergence of new virulent races
Transgenic resistance	Transgenes encoding for toxic products to broomrape	Young broomrapes	Low cost of farm implementation of resistant varieties     each series are obtained when using broomrape-responsive gene promoters	Technique under experimental development     Public acceptance in some countries
	RNA interference	Young attached parasites (broomrape-specific RNAi reaches via the haustorium)	Low cost  Environmental compatible Requires previous identification of parasitic genes essential for broomrape virulence and metabolism Resistant varieties should be used in an integrated management strategy to avoid resistance breakdown by the parasite	Genetic redundancy may dilute silencing effect     Technique under experimental development
	Transgenic herbicide resistance	Attached parasites (herbicide reaches the parasite via the haustorium)	Herbicide-resistance mechanism must be based in mechanism other than metabolic degradation or inactivation of herbicide by the transgenic crop     Low herbicide doses are required to kill the young parasite	<ul> <li>Food safety issues</li> <li>Herbicide resistant-gene transfer to wild plants</li> <li>Public acceptance in some countries</li> </ul>

# Management Strategies to Protect Crops from Detection by Broomrape Seeds

Due to the small size of the seeds and their inability to develop autotrophy, the establishment probability of broomrape seedlings is very low. Broomrapes counteract the high risk of failure in establishment on a host with highly evolved mechanisms of survival. Broomrape high fecundity, with thousands of seeds released per broomrape plant (Figures 2A,B), multiplies the chances of the next generation to encounter a host and achieve successful parasitism (Parker and Riches, 1993). In addition long lived seed banks under physiological dormancy ensure that germination will occur when a suitable host in its correct stage of development is present nearby (Rubiales et al., 2009b). Preventing the movement of parasitic seeds from infested to non-infested agricultural fields, by contaminated machinery or seed lots, is crucial (Panetta and Lawes, 2005). Once a field is infested, controlling the broomrape seed bank is very difficult due to its high resilience. A variety of methods have been developed to specifically neutralize broomrape pre-attached development though the majority of them are not commercially implemented because they are still at the stage of development or have not proved enough efficiency or applicability for large scale crops. These methods can be classified as cultural and physical, chemical, biological control, and use of host resistance (Rubiales et al., 2009b).

#### **Cultural and Physical Control Practices**

Fertilization can induce soil suppressiveness to initiation of broomrape parasitism. Application of phosphate or nitrogen to deficient soil reduces broomrape parasitism on clover and tomato (Southwood, 1971; Jain and Foy, 1992). Nutrients influence the crop-parasite pre-attached interaction in several ways. Direct toxic effects by urea and ammonium but not nitrate forms inhibit broomrape seed germination and radicle elongation (Jain and Foy, 1992; Abu-Irmaileh, 1994; van Hezewijk and Verkleij, 1996; Westwood and Foy, 1999). Accumulation of ammonium can be toxic to plants and its detoxification occurs via incorporation into organic compounds. The activity of glutamine synthetase in broomrape is very low and therefore carries a reduced broomrape ability to detoxify ammonium. Urea has no detrimental effects in plants but it is toxic to broomrape pre-attached stages probably exercised via ammonium after broomrape urease hydrolyses urea into ammonium. Nitrate is not toxic to broomrape as it lacks the ability to convert nitrate into ammonium (van Hezewijk and Verkleij, 1996). In addition to the toxic effects on broomrape seed and seedling, fertilization can protect crops from broomrape parasitism by means of down-regulating the crop synthesis and exudation of strigolactones, the most potent germinationinducing factors for broomrape. Phosphorous and nitrogen have been described to down regulate strigolactones exudation in some crop species (Yoneyama et al., 2007a,b, 2012). This effect may not be applicable to those broomrape species with preference for classes of germination-inducing factors other than strigolactones (Joel et al., 2011; Auger et al., 2012).

Intercropping systems cultivate simultaneously more than one species in close association to take agronomic advantage of biodiversity, competition, and complementarity between them. For broomrape control, this system seeks the simultaneous cultivation of susceptible host species with inhibitory species of broomrape parasitism. The release of phytochemicals by the roots of the allelopathic component in the intercrop inhibits the broomrape germination and/or radicle elongation toward the host component. Successful reduction of broomrape parasitism in the current crop is obtained by intercropping host species with inhibitory species of cereals, fenugreek, or berseem clover (Fernández-Aparicio et al., 2007, 2008b, 2010a). Careful selection of the non-host component in the intercrop is, however, required as some plant species can act as non-host facilitators and therefore increase the severity of broomrape infection in the host component (Gibot-Leclerc et al., 2013).

A rotation decreasing the frequency of host cultivation is one of the main ways that farmers deal with the broomrape-related problem. This prevents broomrape parasitism from taking place, maintaining the seed bank dormant and reducing the rate of seed bank replenishing. However, it is a long-term strategy due to the long viability of seed bank (Rubiales et al., 2009b), which requires at least a nine-course rotation in order to prevent broomrape seed bank increases (Grenz et al., 2005). Its efficacy for broomrape cultural control can be increased if the farmer includes trap and/or catch crops as components in the rotation (Rubiales et al., 2009b). The concept of trap crops refers to the cultivation of crop species whose root exudates exhibit high germinationinducing activity on broomrape seeds, but these species do not become infected because they are resistant to later stages of the parasitic process indirectly leading to the killing of the young broomrape seedlings due to the lack of proper host. The inductor potential of root exudates from a given species varies with the broomrape considered. Each broomrape species show specificity not only for root exudates in order to germinate but also for host species to invade and feed on, being the germination-stimulatory range usually broader than the actual host range (Fernández-Aparicio et al., 2009b). Potential trap crops have been suggested for broomrape weeds (Parker and Riches, 1993). For instance, root exudates of field pea induces high germination of the very destructive broomrape species O. crenata, O. foetida, O. minor, and P. aegyptiaca, however, it only becomes infected by O. crenata therefore pea may theoretically be a good trap crop against O. foetida, O. minor, and P. aegyptiaca but not for O. crenata infested field (Fernández-Aparicio and Rubiales, 2012). Many other interesting examples of trap crops emerged from a root exudates screening of important crops (Fernández-Aparicio et al., 2009b). The second possibility to increase rotation efficacy for broomrape control is to include catch crops, which are crops that also induce high broomrape germination but they are not resistant to it. On the contrary, they must be highly susceptible, as the farmer is the one with the role of stopping the parasitic process by harvesting the catch crop as a green vegetable before the parasite emerges. For instance, tori (Brassica campestris var. toria) when managed properly as a catch crop can result in up to a 30% reduction in the size of broomrape seed bank (Acharya et al., 2002).

Soil management affects the success of broomrape seeds in becoming established on the host and then the longevity of broomrape seed bank. Minimum tillage reduces the amount of viable seeds incorporated in the soil and then their capacity to reach the crop root system (Ghersa and Martinez-Ghersa, 2000; López-Bellido et al., 2009). The opposite agricultural practice deep-plowing, has been suggested to bring seeds of parasitic weeds to a depth with less oxygen availability and therefore a reduction in its germination capacity (Van Delft et al., 2000). According with pot experiments carried out in the tomato-P. aegyptiaca system, deep-plowing bringing the seeds to depth  $\geq 12$  cm will strongly reduce broomrape infection severity in terms of number of parasites, total parasitic biomass, delayed broomrape emergence and prevention of flower initiation and seed set (Eizenberg et al., 2007).

Solarization is a thermal soil disinfestation method that shows high efficiency reducing the viability of the broomrape seed bank along with other harmful organisms to crops such as plant-parasitic nematodes, disease causing microorganisms and non-parasitic weeds. This method consists in heating the soil through sun energy achieving temperatures above 45°C, by covering a wet soil with transparent polyethylene sheets for a period of 4-8 weeks during the warmest season (Katan, 1981; Mauro et al., 2015). Many beneficial organisms are either able to survive the solarization treatment or able to recolonize solarized soil (Sauerborn et al., 1989; Mauromicale et al., 2001). Not all areas infested by broomrape are suitable for efficient solarization. Hot air temperature and clear skies are required during the solarization period. Its high cost per surface unit makes this method not readily applicable at large scale (Joel, 2000). In addition, this technique generates a considerable amount of plastic waste but the emergence of new materials at low-cost, of biological origin and biodegradable may in the future reduce earth pollution with plastic debris derived from agriculture practices (Fernandez and Ingber, 2013).

#### Chemical Control of Seed Bank

Soil fumigation with methyl bromide has been proved one of the most effective methods to eradicate broomrape seed bank, but this chemical has been banned from use due to its toxic effects on the environment (Joel, 2000; Hershenhorn et al., 2009). Use of other soil sterilants such as metham sodium, dazomet, and 1,3-dichloropropene have shown different degrees of efficacy but their high cost, complex application and negative environmental effects have prevented their widespread use by farmers (Foy et al., 1989; Goldwasser et al., 1995; Hershenhorn et al., 2009) or conducted to the withdrawal of authorization, at least in some countries.

Soil herbigation (saturating the soil with herbicides such as sulfonylureas) effectively controls preattached stages of broomrapes (Hershenhorn et al., 2009) but is hardly compatible with other agricultural cropping practices as detrimental for many crop seedlings for several weeks or months. *In vitro* treatments of a large range of sulfonylurea herbicides inhibit broomrape germination and radicle elongation (Hershenhorn et al., 1998; Plakhine et al., 2001). Incorporation of sulfosulfuron and rimsulfuron directly to the soil provides successful control of preattached stages of broomrape weeds (Eizenberg et al., 2012).

Natural pesticides derived of microbial and plant origin are considered to be less harmful because they usually biodegrade

quicker, resulting in less pollution-related problems. Several toxins have been identified with inhibitory activity on broomrape parasitism by interfering with broomrape germination and radicle elongation (Vurro et al., 2009; Fernández-Aparicio et al., 2013; Cimmino et al., 2014). Special interest arises from those metabolites with a favorable pattern of broomrape-specific effect (e.g., tenuazonic acid) and no described side-effect to other biosystems (Vurro et al., 2009). However, the efficacy of these molecules has been proved only in laboratory essays. Until now, difficulties of purification at industrial scale have hampered the field experimentation with such metabolites (Vurro et al., 2009) despite their interesting potential.

Certain amino acids strongly inhibit the early development of broomrape without phytotoxic effects in the host (Vurro et al., 2006). The amino acid approach to control weeds is inspired on the concept of frenching disease where amino acid endproduct inhibits the activity of a controlling enzyme in the amino acid biosynthesis pathway (Vurro et al., 2006, 2009; Sands and Pilgeram, 2009). The effectiveness of amino acids as broomrape inhibitors has not been proved in real field conditions but field application of amino acids has been effective to manage other parasites such as plant-parasitic nematodes (Zhang et al., 2010). Among the amino acids producing the highest and most consistent inhibitory effects on broomrape germination and radicle elongation, some, such as methionine are being produced in large commercial scale as animal feed supplements. The use of those amino acids as pesticide is classified by the United States Environmental Protection Agency as innocuous to public and environment health (USEPA, 2004). Based on those conditions, methionine has the potential to be used as broomrape herbicide but it needs to be confirmed and its application adjusted to real field conditions.

Synthetic analogs of growth regulators can be successfully used to reduce parasitism by hampering the synchronization of the parasitic seed bank with the growth of the host. Inhibition of seed conditioning and subsequent germination mediated by inhibitors of GA synthesis reduces the receptivity of broomrape seeds to germination-inducing factors. In addition, inhibitors of ABA catabolism inhibit the germination-triggering effect of host-derived strigolactones. For example, soil application of uniconazole, a triazole that is commercially used for growth regulation has proved to reduce parasitism by inhibiting seed conditioning and subsequent germination (Joel, 2000; Zehhar et al., 2002; Song et al., 2005; Lechat et al., 2012).

Promotion of suicidal germination is the technique used to induce broomrape germination with synthetic molecules in the absence of a host to which broomrape can attach, a technique lethal for the parasite as the broomrape seedling is unable to acquire autotrophy. Therefore, it may be possible to achieve broomrape control by fooling the parasite with the delivering to the soil of synthetic analogs of the original host-derived germination-inducing factors such as strigolactones (Johnson et al., 1976). Direct application of strigolactones to the soil has been the subject of intense research. The first attempts to deplete parasitic weed seed bank was made by Johnson et al. (1976) by using the synthetic strigolactone analog GR7. However, instability of this compound, particularly at

pH > 7.5, and lack of optimal formulations rendered this technique not applicable (Saghir, 1986; Babiker et al., 1987, 1988). Recent advances in this research area has led to new, more stable strigolactone analogs and optimization of field application protocols and formulations (Bhattacharya et al., 2009; Zwanenburg et al., 2009; Mwakaboko and Zwanenburg, 2011). Another strategy to induce suicidal germination of broomrape seed bank could be the use of gibberellin agonists. They elicit GA-like germination activity in dormant seeds of several autotrophic plant species (Suttle and Schreiner, 1982; Metzger, 1983), constituting a cheap alternative to natural bioregulators for weed seed bank control (Suttle, 1983). When they are applied in vitro to seeds of P. ramosa and O. minor, they bypass the effect of germination-inducing factors, promoting broomrape germination in absence of host or any germination stimulant (Cala et al., 2015).

Once broomrape germination has occurred, chemicals that reduce the growth of broomrape radicle reduce the chances of reaching the host and therefore parasitism. A novel metabolite, ryecyanatine-A recently isolated from rye (Secale cereale L.), presents potential for broomrape control by promoting rapid cessation of broomrape radicle growth and therefore inhibiting its ability to reach the host. In addition it promotes the development of a layer of papillae at the radicle apex in the absence of host contact, morphology that resembles the attachment organ (Joel and Losner-Goshen, 1994; Cimmino et al., 2015). Because the haustorial organ in broomrape radicle is terminal and its growth is not resumed unless it can immediately penetrate the host, cessation of radicle elongation and haustorial induction in the absence of a host is lethal to the parasite. Other interesting molecules that hamper the ability of broomrape radicle to reach the host have been recently discovered from different microbial and plant origins (Fernández-Aparicio et al., 2013; Cimmino et al., 2014).

# Biological Control of Seed Bank with Living Organisms

Biological control of broomrape is based on the use of living organisms either by killing seed bank or interfering with its host-recognition ability. Assessment of pathogenicity or damages toward non-target plants has to be carefully assessed in order to avoid environmental risks. The efficient action of the biological control agent will depend on its ability to remain active over a large range of ecological conditions (Aly, 2007).

## Phytophagous insects to prevent the build-up of broomrape seed bank

More than 40 insect herbivores from 22 families have been collected on broomrape plants but a majority of them are polyphagous without any specificity for broomrape species being some of them serious pests of important crops (Klein and Kroschel, 2002). *Phytomyza orobanchia* is reported to be broomrape-specific and its main action as biocontrol agent is by reduction of broomrape reproductive activity due to their feeding activity on ovules and young seeds. Often secondary infections by fungi cause early death of broomrape shoots or limit the development of flowers and ovules (Klein and Kroschel, 2002).

The capacity of *P. orobanchia* to reduce broomrape populations is limited by cultural practices and antagonists (Klein and Kroschel, 2002; Aly, 2007).

#### Mycoherbicides attacking broomrape seeds and radicles

The broomrape seed bank efficiency to initiate parasitism can be reduced by incorporation to the soil of several pathogens able to infect preattached broomrape stages such as Fusarium sp. or Ulocladium botrytis (Müller-Stöver, 2001; Boari and Vurro, 2004; Dor and Hershenhorn, 2009). Like most seeds, broomrape seeds are resistant to rapid microbial degradation due to phenols located in its testa (Cezard, 1973). However, hyphae of specific pathogens are able to penetrate the seed coat of broomrape dormant seeds, dissolving the endosperm cell walls and metabolizing the cytoplasm. The advantage of this approach using fungi is that it can be used in absence of host cultivation (Thomas et al., 1999). The parasitic weed radicle that emerges from germinated seed and carries the attachment organ is also targeted by those mycoherbicides (Abbasher and Sauerborn, 1992). This approach would inhibit parasitism by killing the young seedling before it attaches to the host root. The control of broomrape by mycoherbicides does not so far provide the level of control required in highly infested soils (Aly, 2007). Novel approaches can increase broomrape control by fungi. A "multiple-pathogen strategy" in which two or more pathogens are combined has been proved successful for the control of broomrape causing a synergistic effect that can lead to 100% broomrape control (Dor and Hershenhorn, 2003; Müller-Stöver et al., 2005). Refined formulations and encapsulations of fungal propagules increase efficacy in biocontrol by reducing desiccation or microbial competition (Amsellem et al., 1999; Quimby et al., 1999; Kroschel et al., 2000; Müller-Stöver, 2001; Aybeke et al., 2015). Engineering of virulence-enhanced mycoherbicides is another approach of great interest. This approach is based on the selection of naturally occurring mutants that overproduce and excrete an enhanced amount of specific amino acid with broomrape inhibition properties on seed germination and radicle growth (Vurro et al., 2006; Sands and Pilgeram, 2009).

#### Bacteria as biocontrol agents

Pseudomonas aeruginosa, P. fluorescens, Bacillus atrophaeus, B. subtilis are promising biocontrol agents targeting the growth of broomrape radicles (Barghouthi and Salman, 2010). In addition to this direct effect, ethylene-producing bacteria such as Pseudomonas syringae pv. glycinea induce ethylene-mediated suicidal germination in Striga sp. (Berner et al., 1999; Ahonsi et al., 2003), a close relative of broomrapes, however, broomrape germination is not responsive to ethylene (Joel, 2000). Azospirillum brasilense is reported to inhibit broomrape radicle growth (Dadon et al., 2004).

## Microbial interactions that interfere on broomrape ability to recognize its host

Broomrape seeds are less capable to recognize crop roots colonized by arbuscular mycorrhizal fungi, *Rhizobium leguminosarum* or *Azospirillum brasilense* due to change in the composition of the root exudates in colonized plants

(Dadon et al., 2004; Mabrouk et al., 2007a; Fernández-Aparicio et al., 2009c, 2010b; Louarn et al., 2012). A reduced content of broomrape germination-inducing factors in root exudates of mycorrhizal plants has been demonstrated (López-Ráez et al., 2011). Those interactions promote the broomrape seed bank remains dormant inhibiting the initiation of broomrape parasitism, and therefore its rates of seed bank replenishment.

#### **Low-Inducers Crop Genotypes**

Breeding for broomrape resistance stands out as the most economic, easy to adopt and environmentally friendly practice. Because parasitic weeds require host encoded molecules to stimulate the initiation of parasitism both at the level of seed germination and haustorium initiation, breeding for low-inducers genotypes of those processes are obvious targets for resistance (Yoder and Scholes, 2010). Sources of natural resistance based on low exudation of germination-inducing factors exist in legumes and sunflower and are highly effective in inhibiting broomrape weed parasitism (Labrousse et al., 2001, 2004; Rubiales et al., 2003b, 2009a; Pérez-de-Luque et al., 2005; Sillero et al., 2005; Abbes et al., 2010; Fernández-Aparicio et al., 2012b, 2014).

Sources of natural resistance based on reduced release of haustorium-inducing factors is a doubly interesting strategy to inhibit broomrape parasitism because not only it prevents broomrape parasitism in the current crop, but also it promotes the demise of the seed bank by promoting suicidal germination. Sources of low-inducers genotypes exist in crops species attacked by the close related parasitic weed *Striga* (Rich et al., 2004). It remains unknown whether host factors are required by broomrape radicle to initiate haustorium and consequently this strategy has not been fully explored.

# **Control Strategies Targeting Host Penetration**

The major strategy that specifically impedes the broomrape capacity to penetrate the host once the radicle has made contact with host root, is based on the use of host resistance, either genetic resistance obtained by breeding (Pérez-de-Luque et al., 2009; Yoder and Scholes, 2010), or induced resistance by abiotic or biotic agents (Gonsior et al., 2004; Kusumoto et al., 2007).

#### Resistant Crops to Broomrape Invasion

Depending on the genetic background of the resistant host, the intrusive cells of broomrape seedling can be stopped at three different levels in their way of penetration through the root layers to achieve connection with the host vascular system. The first barriers are imposed at the cortex level with reinforced cell walls mediated by either protein cross-linking or with the deposition of metabolites such as suberin, or callose. In addition, accumulation of toxic phenolic compounds at the infection point can be observed in some resistant varieties. Resistance that occurs in the endodermis is mediated by lignification of endodermal and pericycle cell walls. Resistance that occurs in the central cylinder is related

with accumulation of phenolic compounds in the surrounding tissues and nearby xylem vessels inducing a toxic release near the parasite impeding vascular connection (Pérez-de-Luque et al., 2009). However, selecting for high phenolic varieties is likely to induce many other side changes altering agronomic performance.

Lack of knowledge in the molecular regulation of the host-parasite interaction during crop invasion has impeded the development of varieties carrying transgenes with capacity to inhibit broomrape penetration. As alternative, transgenic resistant crops have been engineered with broomrape-inducible expression of toxins specifically targeting the penetrating broomrape seedling. This strategy to abort broomrape invasion requires regulating the toxin production with promoters specifically induced around the site of Orobanche penetration such as the HMG2 promoter, ensuring correct delivery of the toxic effect to the broomrape penetrating seedling and overall low concentration of the toxin in the rhizosphere. As a consequence the crop is protected from broomrape invasion (Joel and Portnoy, 1998; Westwood et al., 1998; Hamamouch et al., 2005; Aly et al., 2006).

#### Abiotic Inducers of Resistance

Induced disease resistance mediated by endogenous salicylic acid (SA) also described as systemic acquired resistance (SAR) induces hypersensitive responses in many plant species against fungal, bacterial and viral diseases. SA promotes resistance to broomrape. Abiotic inducers of SAR thus represent an innovative approach to control broomrape parasitism. Benzo-(1,2,3)-thiadiazole-7-carbothioic acid S-methyl ester (BTH) acts as a functional analog of SA and activates defense responses in susceptible hosts leading to lignification of the endodermis and a consequent inhibition to up to 98% broomrape parasitism (Gonsior et al., 2004; Pérez-de-Luque et al., 2004; Kusumoto et al., 2007). Commercially available as Bion®, field doses of 0.8 kg ha<sup>-1</sup> are recommended to inhibit P. ramosa parasitism in hemp and tobacco (Gonsior et al., 2004), crops for which resistant varieties are not available. In other pathosystems, amino acids such as D-L-β-amino*n*-butyric acid or L-methionine induce resistance in crop plants against pathogen attack. This resistance is coordinated with the expression of genes encoding for pathogenesisrelated proteins (Sarosh et al., 2005; Hasabi et al., 2014). The ability of L-methionine to stop the entrance of broomrape intrusive cells into the host-root layers has not been studied. However, when Vurro et al. (2006) applied L-methionine in pots to tomato roots the number of broomrape seedlings that successfully developed parasitism was highly reduced. Though, the effect of L-methionine on internal crop resistance was not studied and requires further investigation. If this effect is confirmed, L-methionine use to elicit resistance to broomrape in susceptible crops could be a straightforward strategy either by direct applications of this amino acid in the soil as explained in Section "Control Strategies Targeting Host Penetration" or delivered by overproducing and excreting microorganisms as explained in Section

"Strategies to Control Underground Broomrapes Acting after Establishment."

#### **Biotic Inducers of Resistance**

Biotic inducers of systemic resistance have also proved being successful against broomrape parasitism under experimental conditions. Rhizobium leguminosarum induces defense mechanisms based on elevated induction of the phenylpropanoid pathway conferring mechanical and chemical barriers to the parasite penetration (Mabrouk et al., 2007a,b,c, 2010). Although the effect of jasmonic-acid-dependent induced systemic resistance (ISR) against parasitic plants is less clear (Kusumoto et al., 2007; Hiraoka et al., 2009; Yoder and Scholes, 2010), strains of Pseudomonas sp. inducers of ISR (Gozzo, 2003) and commercially available as Proradix® can reduce broomrape parasitism by 80% in susceptible cultivars of hemp and tobacco without phytotoxic effect on the crop (Gonsior et al., 2004). Based on the results obtained in their greenhouse experiments. these authors recommended field doses of 1.6 kg ha<sup>-1</sup> for crop densities of 32,000 tobacco plants ha<sup>-1</sup>. Unfortunately this technique represents another example of highly promising broomrape control strategy that has never been validated in field experiments.

## Strategies to Control Underground Broomrapes Acting after Establishment

Once broomrape has established connection with the vascular system of its hosts, broomrape management should be performed quickly to abort at earlier stages the strong parasitic sink for nutrients and water. Major feasible strategies for controlling broomrape and gain productivity in the current crop are those based on cultural practices that promote host scape to parasitic damage by improving host sink competitiveness, selective chemical control of the parasite via the haustorium, and host resistance based in physical, chemical barriers and physiological incompatibility. Some of the strategies discussed in previous sections such as biological control maintain their control action at post-attachment stages and will not be discussed again in this section.

#### **Cultural Control of Post-attached Parasites**

Delayed sowing date is a traditional method that can show high degree of success on inhibiting parasitism if implemented correctly (López-Granados and García-Torres, 1996; Rubiales et al., 2003a; Pérez-de-Luque et al., 2004; Grenz et al., 2005). This technique promotes the host plant to fulfill its required thermal time to flower in a shorter number of days, making the grain filling period shorter. The host reproductive sinks compete earlier and stronger against the parasitic sink and in consequence less nutritive resources are allocated to the parasite (Manschadi et al., 1996). However, the overall productivity of the host-parasite system is also reduced due to the shorter growing period being detrimental for crop yield. Therefore, decisions on the date of sowing has to be well-adjusted in order to balance the loss of productivity due to shorter growing period with gain of productivity due to reduced parasitism. Delaying sowing date has, however, a general drawback by reducing yield potential

under normal development so that plant breeding program tend generally to favor long lasting cultivars with early sowing dates.

Besides date of sowing, nutrient management can promote both tolerance and increased resistance in crops to broomrape parasitism (Parker, 2009; Labrousse et al., 2010). Broomrape attack is more severe on crops growing in low fertility soils. Besides the effects of fertilization management on pre-attached broomrape stages described in previous sections, high soil fertility can induce crops to endure broomrape parasitism by helping the host to maintain a favorable osmotic potential that reduces the parasitic sink strength (Gworgwor and Weber, 1991).

#### Chemical Control of Post-attached Parasites

Four broomrape features define the post-attachment herbicidal strategy in comparison with non-parasitic weeds. First, broomrape weeds are achlorophyllous and therefore those herbicides that target photosynthetic process, e.g., triazines or substituted urease [C group in the Herbicide Resistance Action Committee (HRAC) classification], will have only limited effect on broomrapes. Second, broomrape weed exerts their damage underground right after attachment and therefore, contact herbicides applied after broomrape emergence, e.g., 2,4-D, had no effect on limiting yield loss in the current crop. Third, broomrape underground attachments do not take herbicides from the soil but only systemically from the host and therefore, this strategy is limited to systemic herbicides applied to herbicide-resistant crop varieties that do not metabolize the herbicide into inactive forms. And four, despite reports on broomrape inefficient machinery for nitrogen assimilation, and on amino acid fluxes from the host phloem to the parasite, herbicides inhibiting amino acid biosynthesis in the parasite via suppressive action on broomrape-encoded acetolactate synthase (ALS) and enolpyruvylshikimate phosphate synthase (EPSPS) enzymes are able to kill broomrape.

Current chemical control of post-attached broomrape life stages is mainly achieved with foliar applications of systemic herbicides inhibiting ALS (imidazolinones, sulfonylureas) or EPSPS (glyphosate) to the leaves of crop varieties carrying target-site resistances to those herbicides to avoid direct injury of their metabolism. The target-site herbicide-resistance is based on a modification of the enzyme in such a way that it binds to its normal substrate in the amino acid biosynthesis pathway but not to the herbicide. This kind of resistance is more interesting than other mechanisms of resistance that usually involve translocation and enhanced metabolism, resulting in lower herbicide concentration in the sap of the host plant. With target-site resistance, the herbicide translocates unmetabolised to the underground broomrape via the haustorium inflicting its suppressive action in the parasite (Gressel, 2009).

Target-site resistances have been successfully developed in crops either by classical breeding such as sunflower, by screening mutagenized crop populations such as the case of oilseed rape or by transgenic techniques such as tomato, tobacco, carrots, and oilseed rape (Joel et al., 1995; Aviv et al., 2002; Slavov et al., 2005; Tan et al., 2005). Such target-site resistance is

also available in other broomrape-susceptible crops but remains to be tested and registered to control broomrape. Engineered host crops harboring herbicide-resistance transgenes have not yet been commercialized for broomrape management (Gressel, 2009<sup>2</sup>).

An alternative to the selective use of herbicides when targetsite resistance is not available for a specific crop is the touchy use of repeated applications of non-selective herbicidal doses to promote sublethal effects for the crop but lethal effects to the initial stages of post-attached parasitism (Foy et al., 1989). This strategy requires a careful calibration of doses and timing depending on the host crop and underground phenology of broomrape determined by local conditions and crop (Hershenhorn et al., 1998, 2009; Eizenberg et al., 2006). Systemic translocation of nanoencapsulated herbicides could improve this herbicidal approach (Pérez-de-Luque and Rubiales, 2009).

#### **Resistant and Tolerant Cultivars**

Death of the young broomrape tubercles shortly after nutritive flow initiation has been observed in cultivars carrying post-haustorial resistance in the form of growth arrest and necrosis of young tubercles. Several mechanisms underlying this resistance have been described at this stage such as production of gel-like substances within host vessels blocking the transfer of nutrients, host-encoded toxic-compounds delivered into the parasitic tissue though the vascular system and hormonal incompatibility that leads to abnormal haustorial maturation with scarce vascular connections (Fernández-Aparicio et al., 2008c; Pérez-de-Luque et al., 2008, 2009). Those mechanisms kill the broomrape either by inducing toxic effects or by starving the parasite.

Tolerant varieties are able to endure infection with minor losses on productivity. Crops that reach their seed filling period earlier than broomrape initiates its underground bud development are able to restrict parasitic sink and endure parasitic damage (Manschadi et al., 1996; Grenz et al., 2005; Fernández-Aparicio et al., 2009a, 2012a). In addition, some modifications of host biochemistry have been described in tolerant crops inducing low performance of the parasite when attached. High osmotic potential in roots and drop in amino acid levels in the phloem has been reported in tolerant varieties of faba bean in response to broomrape parasitism. The consequent reduced flux of water and nutrients toward the parasite, low utilization of host-derived sucrose and lower levels of soluble proteins limits the parasitic sink strength and yield losses due to broomrape parasitism (Abbes et al., 2009).

#### CONCLUSION

If left uncontrolled during one or a few seasons, broomrape weeds build a hardly destructible seed bank in agricultural soils that further renovates at a rate of millions of seeds per ha each year a susceptible crop is infested. In addition, their mixed traits of weed and underground pathogen, make their control tricky. Even the easiest method of control, herbicides, requires broomrape specific-optimization for each cropping system to target the most vulnerable broomrape life stage, the young attachments while preserving the crop. It seems more and more obvious that a single strategy has low probability to control broomrapes. Instead an integrated control program including a battery of broomrape-specific measurements is preferable.

We have seen that several opportunities to stop the cycle of the parasite have been explored. The development of the solutions has usually not been conducted to their end so that many potential ways of controlling broomrape are not on the market. What we have often seen is that the solution has to propose a modification that makes the parasitic life cycle unfit to that of the crop. Still, as the parasite is synchronized on the crop development this means in some cases that the change disfavoring the parasite could also limit the maximum potential yield for the crop. As a consequence, except when deeply infested, the farmer (and thus the market) will not retain a solution that has economical negative drawbacks. This may well-explain why some several decades of parasitic plant research have not end up with satisfying and largely available tools for controlling this parasitic plant.

The advances yielded as intense research made connects the major critical steps of the life cycle of Orobanche, the external factors influencing it either through molecular dialog between the parasite and the crop or the soil and climatic environmental conditions naturally opens the way toward the potential effect of the cropping system in limiting broomrape parasitism: choice of the crop, timing, plant protection, soil perturbation, fertilization, etc. This lead us to build the list of the major feasible components that a model designed to quantify the effects of cropping systems on pest dynamics should include for specific broomrape control. Such a model would be a valuable tool to synthesize knowledge on broomrape life-cycle, to design and test management strategies and better predict the variability in effects observed for a given environment and set of agricultural practices. One future development would be to evaluate what could be the emerging risk at cultivating different crops, one of which may stimulate germination while the other offers opportunities for haustorium fixation. This would open the work on parasitism toward more community ecology and what can be considered the realistic nature of parasitism.

#### **AUTHOR CONTRIBUTIONS**

MF-A wrote the paper. XR and SG-L additional text, editing, and comments.

#### **ACKNOWLEDGMENTS**

This work was cofunded by the European Union and INRA, in the framework of the Marie-Curie FP7 COFUND People Program, through the award of an AgreenSkills' fellowship (under grant agreement n° PCOFUND-GA-2010-267196) to MF-A with additional support by the INRA Division "Santé des Plantes et Environnement."

<sup>&</sup>lt;sup>2</sup>http://www.isb.vt.edu

#### **REFERENCES**

- Abbasher, A. A., and Sauerborn, J. (1992). Fusarium nygamai a potential bioherbicide for Striga hermonthica control in sorghum. Biol. Control 2, 291–296. doi: 10.1016/1049-9644(92)90021-5
- Abbes, Z., Kharrat, M., Delavault, P., Chaïbi, W., and Simier, P. (2009). Nitrogen and carbon relationships between the parasitic weed *Orobanche foetida* and susceptible and tolerant faba bean lines. *Plant Physiol. Biochem.* 47, 153–159. doi: 10.1016/j.plaphy.2008.10.004
- Abbes, Z., Kharrat, M., Pouvreau, J. B., Delavault, P., Chaibi, W., and Simier, P. (2010). The dynamics of faba bean (*Vicia faba L.*) parasitism by *Orobanche foetida*. *Phytopathol. Mediterr*. 49, 239–248.
- Aber, M., Fer, A., and Salle, G. (1983). Transfer of organic substances from the host plant *Vicia faba* to the parasite *Orobanche crenata* Forsk. *Z. Planzenphysiol.* 112, 297–308. doi: 10.1016/S0044-328X(83)80047-6
- Abu-Irmaileh, B. E. (1994). Nitrogen reduces branched broomrape (*Orobanche ramosa*) seed germination. Weed Sci. 42, 57–60.
- Abu-Irmaileh, B. E., and Labrada, R. (2009). *The Problem of Orobanche spp. in Africa and Near East.* Available at: www.fao.org/ag/AGP/AGPP/IPM/Weeds/Issues/orobanche.htm
- Acharya, B. D., Khattri, B. G., Chettri, M. K., and Srivastava, X. (2002). Effect of *Brassica campestris* var. toria as a catch crop on *Orobanche aegyptiaca* seed bank. *Crop Prot.* 21, 533–537. doi: 10.1016/S0261-2194(01) 00137-5
- Ahonsi, M. O., Berner, D. K., Emechebe, A. M., Lagoke, S. T., and Sangina, N. (2003). Potential of ethylene-producing pseudomonads in combination with effective N2-fixing bradyrhizobial strains as supplements to legume rotation for Striga hermonthica control. Biol. Control 28, 1–10. doi: 10.1016/S1049-9644(03)00051-3
- Akiyama, K., Matsuzaki, K. I., and Hayashi, H. (2005). Plant sesquiterpenes induce hyphal branching in arbuscular mycorrhizal fungi. *Nature* 435, 824–827. doi: 10.1038/nature03608
- Albrecht, H., Yoder, J. I., and Phillips, D. A. (1999). Flavonoids promote haustoria formation in the root parasite *Triphysaria versicolor*. *Plant Physiol*. 119, 585– 591. doi: 10.1104/pp.119.2.585
- Aly, R. (2007). Conventional and biotechnological approaches for control of parasitic weeds. In Vitro Cell. Dev. Biol. Plant 43, 304–317. doi: 10.1007/s11627-007-9054-5
- Aly, R., Plakhin, D., and Achdari, G. (2006). Expression of sarcotoxin IA gene via a root-specific tob promoter enhanced host resistance against parasitic weeds in tomato plants. *Plant Cell Rep.* 25, 297–303. doi: 10.1007/s00299-005-0052-y
- Amsellem, Z., Zidack, N. K., Quimby, Jr P. C, and Gressel, J. (1999). Long term dry preservation of active mycelia of two mycoherbicidal organisms. *Crop Prot.* 18, 643–649. doi: 10.1016/S0261-2194(99)00070-8
- Antonova, T. S., and Ter Borg, S. J. (1996). The role of peroxidase in the resistance of sunflower against O. cumana in Russia. Weed Res. 36, 113–121. doi:10.1111/j.1365-3180.1996.tb01807.x
- Atsatt, P. R. (1977). The insect herbivore as a predictive model in parasitic seed plant biology. Am. Nat. 111, 579–586. doi: 10.1086/ 283185
- Auger, B., Pouvreau, J. B., Pouponneau, K., Yoneyama, K., Montiel, G., Le Bizec, B., et al. (2012). Germination stimulants of *Phelipanche ramosa* in the rhizosphere of *Brassica napus* are derived from the glucosinolate pathway. *Mol. Plant Microbe Interact.* 25, 993–1004. doi: 10.1094/MPMI-01-12-0006-R
- Aviv, D., Amsellem, Z., and Gressel, J. (2002). Transformation of carrots with mutant acetolactate synthase for *Orobanche* (broomrape) control. *Pest Manag. Sci.* 58, 1187–1193. doi: 10.1002/ps.567
- Aybeke, M., Şen, B., and Ökten, S. (2015). Pesta granule trials with Aspergillus alliaceus for the biocontrol of Orobanche spp. Biocontrol Sci. Technol. 25, 803–813. doi: 10.1080/09583157.2015.1018813
- Babiker, A. G. T. (2008). Orobanche crenata in Sudan: history, distribution and management. Haustorium 53, 1–3.
- Babiker, A. G. T., Ahmed, E. A., Dawoud, D. A., and Abdrella, N. K. (2007). Orobanche species in Sudan: history, distribution and management. Sudan J. Agric. Res. 10, 107–114.

- Babiker, A. G. T., Hamdoun, A. M., Rudwan, A., Mansi, N. G., and Faki, H. H. (1987). Influence of soil moisture on activity and persistence of the strigol analogue GR 24. Weed Res. 27, 173–178. doi: 10.1111/j.1365-3180.1987.tb00751.x
- Babiker, A. G. T., Ibrahim, N. E., and Edwards, W. G. (1988). Persistence of GR7 and *Striga* germination stimulant(s) from *Euphorbia aegyptiaca* Boiss. in soils and in solutions. *Weed Res.* 28, 1–6. doi: 10.1111/j.1365-3180.1988.tb00778.x
- Bandaranayake, P. C. G., and Yoder, J. I. (2013). "Haustorium initiation and early development," in *Parasitic Orobanchaceae*, eds D. M. Joel, L. J. Musselman, and J. Gressel (Berlin: Springer), 61–74.
- Barghouthi, S., and Salman, M. (2010). Bacterial inhibition of Orobanche aegyptiaca and Orobanche cernua radical elongation. Biocontrol Sci. Technol. 20, 423–435. doi: 10.1080/09583150903340544
- Barker, E. R., Press, M. C., Scholes, J. D., and Quick, W. P. (1996). Interactions between the parasitic angiosperm *Orobanche aegyptiaca* and its tomato host: growth and biomass allocation. *New Phytol.* 133, 637–642. doi: 10.1111/j.1469-8137.1996.tb01932.x
- Barkman, T. J., McNeal, J. R., Lim, S. H., Coat, G., Croom, H. B., Young, N. D., et al. (2007). Mitochondrial DNA suggests at least 11 origins of parasitism in angiosperms and reveals genomic chimerism in parasitic plants. BMC Evol. Biol. 7:248. doi: 10.1186/1471-2148-7-248
- Bar-Nun, N., Ben-Hod, G., Lavi, E., and Mayer, A. M. (1996). Purification of pectin methylesterase from *Orobanche aegyptiaca*. *Phytochemistry* 41, 403–406. doi: 10.1016/0031-9422(95)00594-3
- Bar-Nun, N., and Mayer, A. M. (1993). Preconditioning and germination of Orobanche seeds: respiration and protein synthesis. Phytochemistry 34, 39–45. doi: 10.1016/S0031-9422(00)90779-9
- Bar-Nun, N., and Mayer, A. M. (2002). Composition of and changes in storage compounds in *Orobanche aegyptiaca* seeds during preconditioning. *Isr. J. Plant* Sci. 50, 277–279. doi: 10.1560/E2KB-FM11-X4U2-YC9J
- Bar-Nun, N., Sachs, T., and Mayer, A. M. (2008). A role for IAA in the infection of Arabidopsis thaliana by Orobanche aegyptiaca. Ann. Bot. 101, 261–265. doi: 10.1093/aob/mcm032
- Ben-Hod, G., Losner-Goshen, D., Joel, D. M., and Mayer, A. M. (1993).Pectin methylesterase in calli and germinating seeds of *Orobanche aegyptiaca*.Phytochemistry 32, 1399–1402. doi: 10.1016/0031-9422(93)85145-H
- Bennett, J. R., and Mathews, S. (2006). Phylogeny of the parasitic plant family Orobanchaceae inferred from phytochrome A. Am. J. Bot. 93, 1039–1051. doi: 10.3732/ajb.93.7.1039
- Berner, D. K., Schaad, N. W., and Volksch, B. (1999). Use of ethylene producing bacteria for stimulating of Striga spp. seed germination. Biol. Control 15, 274–282. doi: 10.1006/bcon.1999.0718
- Bhattacharya, C., Bonfante, P., Deagostino, A., Kapulnik, Y., Larini, P., Occhiato, E. G., et al. (2009). A new class of conjugated strigolactone analogues with fluorescent properties: synthesis and biological activity. *Org. Biomol. Chem.* 7, 3413–3420. doi: 10.1039/b907026e
- Boari, A., and Vurro, M. (2004). Evaluation of *Fusarium* spp. and other fungi as biological control agents of broomrape (*Orobanche ramosa*). *Biol. Control* 30, 212–219. doi: 10.1016/j.biocontrol.2003.12.003
- Cala, A., Rial, C., Fernandez-Aparicio, M., Molinillo, J. M. G., Varela, R. M., Rubiales, D., et al. (2015). "Phthalimide-lactones stimulate germination of parasitic weeds," in *Proceedings of the XXXV Biennial Meeting of the Spanish Royal Society of Chemistry*, eds J. A. S. López, E. I. Martínez, T. R. Blas, M. C. López, and J. P. Sestelo (A Coruña: Dario Prada-Rodríguez of University of A Coruña), 688.
- Cezard, R. (1973). "Quelques aspects particuliers de la biologie des Orobanches," in Proceedings of the European Weed Research Council on Parasitic Weeds, eds W. G. H. Edwards, L. Kasasian, C. Parker, A. R. Saghir, and W. van der Zweep (Malta: Royal University of Malta), 55–67.
- Chae, S. H., Yoneyama, K., Takeuchi, Y., and Joel, D. M. (2004). Fluridone and norflurazon, carotenoid-biosynthesis inhibitors, promote seed conditioning and germination of the holoparasite *Orobanche minor*. *Physiol. Plant.* 120, 328–337. doi: 10.1111/j.0031-9317.2004.0243.x
- Cimmino, A., Fernandez-Aparicio, M., Andolfi, A., Basso, S., Rubiales, D., and Evidente, A. (2014). Effect of fungal and plant metabolites on broomrapes (Orobanche and Phelipanche spp.) seed germination and radicle growth. J. Agric. Food Chem. 62, 10485–10492. doi: 10.1021/jf504609w

- Cimmino, A., Fernandez-Aparicio, M., Avolio, F., Andolfi, A., Rubiales, D., Yoneyama, K., et al. (2015). Ryecyanatines A and B and ryecarbonitrilines A and B, substituted cyanatophenol, cyanato-benzo[1,3] diole, and benzo[1,3]dioxolecarbonitriles from rye (*Secale cereale* L.) root exudates: new metabolites with allelophatic activity on *Orobanche* seed germination and radicle growth. *Phytochemistry* 109, 57–65. doi: 10.1016/j.phytochem.2014.10.034
- Conn, C. E., Bythell-Douglas, R., Neumann, D., Yoshida, S., Whittington, B., Westwood, J. H., et al. (2015). Convergent evolution of strigolactone perception enabled host detection in parasitic plants. *Science* 349, 540–543. doi: 10.1126/science.aab1140
- Dadon, T., Nun, N. B., and Mayer, A. M. (2004). A factor from Azospirillum brasilense inhibits germination and radicle growth of Orobanche aegyptiaca. Isr. J. Plant Sci. 52, 83–86. doi: 10.1560/Q3BA-8BJW-W7GH-XHPX
- Das, M., Fernández-Aparicio, M., Yang, Z. Z., Huang, K., Wickett, N. J., Alford, S., et al. (2015). Parasitic plants Striga and Phelipanche dependent upon exogenous strigolactones for germination have retained genes for strigolactone biosynthesis. Am. J. Plant Sci. 6, 1151–1166. doi: 10.4236/ajps.2015. 68120
- De Candolle, A. P. (1813). Théorie Elémentaire de la Botanique. Paris: Déterville.
- Dor, E., and Hershenhorn, J. (2003). The use of several phytopathogenic fungi for broomrape control. *Phytoparasitica* 31, 422.
- Dor, E., and Hershenhorn, J. (2009). Evaluation of the pathogenicity of microorganisms isolated from Egyptian broomrape (*Orobanche aegyptiaca*) in Israel. Weed Biol. Manage. 9, 200–208. doi: 10.1111/j.1445-6664.2009. 00340.x
- Dörr, I. (1997). How Striga parasitizes its host: a TEM and SEM study. Ann. Bot. 79, 463–472. doi: 10.1006/anbo.1996.0385
- Dörr, I., and Kollmann, R. (1995). Symplasmic sieve element continuity between Orobanche and its host. Bot. Acta 108, 47–55. doi: 10.1111/j.1438-8677.1995.tb00830.x
- Draie, R., Péron, T., Pouvreau, J.-B., Véronési, C., Jégou, S., Delavault, P., et al. (2011). Invertases involved in the development of the parasitic plant *Phelipanche ramosa*: characterization of the dominant soluble acid isoform, PrSAI1. *Mol. Plant Pathol.* 12, 638–652. doi: 10.1111/j.1364-3703.2010. 00702.x
- Ehleringer, J. R., and Marshall, J. D. (1995). "Water relations," in *Parasitic Plants*, eds M. C. Press and J. Graves (London: Chapman and Hall), 125–140.
- Eizenberg, H., Aly, R., and Cohen, Y. (2012). Technologies for smart chemical control of broomrape (*Orobanche* spp. and *Phelipanche* spp.). Weed Sci. 60, 316–323. doi: 10.1614/WS-D-11-00120.1
- Eizenberg, H., Colquhoun, J. B., and Mallory-Smith, C. A. (2006). Imazamox application timing for small broomrape (*Orobanche minor*) control in red clover. Weed Sci. 54, 923–927. doi: 10.1614/WS-05-151R.1
- Eizenberg, H., Lande, T., Achdari, G., Roichman, A., and Hershenhorn, J. (2007). Effect of Egyptian broomrape (*Orobanche aegyptiaca*) burial depth on parasitism dynamics and chemical control in tomato. Weed Sci. 51, 152–156. doi: 10.1614/WS-06-135
- Evidente, A., Cimmino, A., Fernández-Aparicio, M., Andolfi, A., Rubiales, D., and Motta, A. (2010). Polyphenols, including the new peapolyphenols A–C, from root exudates stimulate *Orobanche foetida* seed germination. *J. Agric. Food Chem.* 58, 2902–2907. doi: 10.1021/jf904247k
- Evidente, A., Cimmino, A., Fernández-Aparicio, M., Rubiales, D., Andolfi, A., and Melck, D. (2011). Soyasapogenol B and trans-22-dehydrocamposterol from common vetch (*Vicia sativa L.*) root exudates stimulate broomrape seed germination. *Pest Manag. Sci.* 67, 1015–1022. doi: 10.1002/ps.2153
- Evidente, A., Fernández-Aparicio, M., Cimmino, A., Rubiales, D., Andolfi, A., and Motta, A. (2009). Peagol and peagoldione, two new strigolactone like metabolites isolated from pea root exudates. *Tetrahedron Lett.* 50, 6955–6958. doi: 10.1016/j.tetlet.2009.09.142
- Fernandez, J., and Ingber, D. (2013). Bioinspired chitinous material solutions for environmental sustainability and medicine. Adv. Funct. Mater. 23, 4454–4466. doi: 10.1002/adfm.201300053
- Fernández-Aparicio, M., Andolfi, A., Evidente, A., Pérez-de-Luque, A., and Rubiales, D. (2008a). Fenugreek root exudates show species-specific stimulation of *Orobanche* seed germination. *Weed Res.* 48, 163–168. doi: 10.1111/j.1365-3180.2007.00609.x

- Fernández-Aparicio, M., Emeran, A. A., and Rubiales, D. (2008b). Control of Orobanche crenata in legumes intercropped with fenugreek (Trigonella foenumgraecum). Crop Prot. 27, 653–659. doi: 10.1016/j.cropro.2007.09.009
- Fernández-Aparicio, M., Pérez-de-Luque, A., Prats, E., and Rubiales, D. (2008c). Variability of interactions between barrel medic (*Medicago truncatula*) genotypes and *Orobanche* species. *Ann. Appl. Biol.* 153, 117–126. doi: 10.1111/j.1744-7348.2008.00241.x
- Fernández-Aparicio, M., Emeran, A. A., and Rubiales, D. (2010a). Inter-cropping with berseem clover (*Trifolium alexandrinum*) reduces infection by *Orobanche* crenata in legumes. Crop Prot. 29, 867–871. doi: 10.1016/j.cropro.2010.03.004
- Fernández-Aparicio, M., García-Garrido, J. M., Ocampo, J. A., and Rubiales, D. (2010b). Colonization of field pea roots by arbuscular mycorrhizal fungi reduces *Orobanche* and *Phelipanche* species seed germination. *Weed Res.* 50, 262–268. doi: 10.1111/j.1365-3180.2010.00771.x
- Fernández-Aparicio, M., Flores, F., and Rubiales, D. (2009a). Field response of *Lathyrus cicera* germplasm to crenate broomrape (*Orobanche crenata*). Field *Crops Res.* 113, 321–327. doi: 10.1016/j.fcr.2009.06.009
- Fernández-Aparicio, M., Flores, F., and Rubiales, D. (2009b). Recognition of root exudates by seeds of broomrape (*Orobanche* and *Phelipanche*) species. *Ann. Bot.* 103, 423–431. doi: 10.1093/aob/mcn236
- Fernández-Aparicio, M., Soto, M. J., Rubiales, D., Ocampo, J. A., and García-Garrido, J. M. (2009c). The potential of *Rhizobium* mutants for biological control of *Orobanche crenata*. *Haustorium* 54, 3–4.
- Fernández-Aparicio, M., Flores, F., and Rubiales, D. (2012a). Escape and true resistance to crenate broomrape (*Orobanche crenata* Forsk.) in grass pea (*Lathyrus sativus* L.) germplasm. Field Crops Res. 125, 92–97. doi: 10.1016/j.fcr.2011.09.003
- Fernández-Aparicio, M., Moral, A., Kharrat, M., and Rubiales, D. (2012b). Resistance against broomrapes (*Orobanche* and *Phelipanche* spp.) in faba bean (*Vicia faba*) based in low induction of broomrape seed germination. *Euphytica* 186, 897–905. doi: 10.1021/if5027235
- Fernández-Aparicio, M., Kisugi, T., Xie, X., Rubiales, D., and Yoneyama, K. (2014). Low strigolactone root exudation: a novel mechanism of broomrape (*Orobanche* and *Phelipanche* spp.) resistance available for faba bean breeding. *J. Agric. Food Chem.* 62, 7063–7071. doi: 10.1021/jf5027235
- Fernández-Aparicio, M., and Rubiales, D. (2012). Differential response of pea (*Pisum sativum*) to *Orobanche crenata*, *Orobanche foetida* and *Phelipanche aegyptiaca*. Crop Prot. 31, 27–30. doi: 10.1093/jxb/err246
- Fernández-Aparicio, M., Sillero, J. C., and Rubiales, D. (2007). Intercropping with cereals reduces infection by *Orobanche crenata* in legumes. *Crop Prot.* 26, 1166–1172. doi: 10.1016/j.cropro.2006.10.012
- Fernández-Aparicio, M., Yoneyama, K., and Rubiales, D. (2011). The role of strigolactones in host specificity of *Orobanche* and *Phelipanche* seed germination. *Seed Sci. Res.* 21, 55–61. doi: 10.1017/S0960258510000371
- Fernández-Aparicio, M., Cimmino, A., Evidente, A., and Rubiales, D. (2013). Inhibition of *Orobanche crenata* seed germination and radicle growth by allelochemicals identified in cereals. *J. Agric. Food Chem.* 61, 9797–9803. doi: 10.1021/jf403738p
- Finch-Savage, W. E., and Leubner-Metzger, G. (2006). Seed dormancy and the control of germination. *New Phytol.* 171, 501–523. doi: 10.1111/j.1469-8137.2006.01787.x
- Foy, C. L., Jain, R., and Jacobsohn, R. (1989). Recent approaches for chemical control of broomrape (*Orobanche* spp.) – a review. *Rev. Weed Sci.* 4, 123–152.
- Ghersa, C. M., and Martinez-Ghersa, M. A. (2000). Ecological of weed seed size and persistence in the soil under different tilling systems: implications for weed management. *Field Crops Res.* 67, 141–148. doi: 10.1016/S0378-4290(00) 00089-7
- Gibot-Leclerc, S., Abdennebi-Abdemessed, N., Reibel, C., and Colbach, N. (2013).Non-host facilitators, a new category that unexpectedly favours parasitic weeds.Agron. Sustain. Dev. 33, 787–793. doi: 10.1007/s13593-013-0153-x
- Gibot-Leclerc, S., Corbineau, F., Sallé, G., and Côme, D. (2004). Responsiveness of *Orobanche ramosa* L. seeds to GR24 as related to temperature, oxygen availability and water potential during preconditioning and subsequent germination. *Plant Growth Regul.* 43, 63–71. doi: 10.1023/B:GROW.0000038242.77309.73
- Goldwasser, Y., Kleifeld, Y., Golan, S., Bargutti, A., and Rubin, B. (1995).

  Dissipation of metham-sodium from soil and its effect on the control

- of Orobanche aegyptiaca. Weed Res. 35, 445-452. doi: 10.1111/j.1365-3180.1995.tb01641.x
- Gomez-Roldan, V., Fermas, S., Brewer, P. B., Puech-Pages, V., Dun, E. A., Pillot, J. P., et al. (2008). Strigolactone inhibition of shoot branching. *Nature* 455, 189–194. doi: 10.1038/nature07271
- Gonsior, G., Buschmann, H., Szinicz, G., Spring, O., and Sauerborn, J. (2004). Induced resistance – an innovative approach to manage branched broomrape (*Orobanche ramosa*) in hemp and tobacco. Weed Sci. 52, 1050–1053. doi: 10.1614/WS-04-088R1
- Gozzo, F. (2003). Systemic acquired resistance in crop protection: from nature to chemical approach. J. Agric. Food Chem. 51, 4487–4503. doi: 10.1021/jf0300258
- Grenz, J. H., Manschadi, A. M., Uygurc, F. N., and Sauerborn, J. (2005). Effects of environment and sowing date on the competition between faba bean (*Vicia faba*) and the parasitic weed *Orobanche crenata*. Field Crops Res. 93, 300–313. doi: 10.1016/j.fcr.2004.11.001
- Grenz, J. H., and Sauerborn, J. (2007). Mechanisms limiting the geographical range of the parasitic weed *Orobanche crenata*. Agric. Ecosyst. Environ. 122, 275–281. doi: 10.1016/j.agee.2007.01.014
- Gressel, J. (2009). Crops with target-site herbicide resistance for *Orobanche* and *Striga* control. *Pest Manag. Sci.* 65, 560–565. doi: 10.1002/ps.1738
- Gworgwor, N. A., and Weber, H. C. (1991). Effect of N-application on sorghum growth, Striga infestation and the osmotic presure of the parasite in relation to the host. J. Plant Physiol. 139, 194–198.
- Hamamouch, N., Westwood, J. H., Banner, I., Cramer, C. L., Gepstein, S., and Aly, R. (2005). A peptide from insects protects transgenic tobacco from a parasitic weed. *Transgenic Res.* 14, 227–236. doi: 10.1007/s11248-004-7546-1
- Harb, A. M., Hameed, K. M., and Shibli, R. A. (2004). Effect of triiodobenzoic acid on broomrape (*Orobanche ramosa*) infection and development in tomato plants. *Plant Pathol. J.* 20, 81–84. doi: 10.5423/PPJ.2004.20.2.081
- Hasabi, V., Askari, H., Alavi, S. M., and Zamanizadeh, H. (2014). Effect of amino acid application on induced resistance against citrus canker disease in lime plants. J. Plant Prot. Res. 54, 144–149. doi: 10.2478/jppr-2014-0023
- Hearne, S. J. (2009). Control the Striga conundrum. Pest Manag. Sci. 65, 603–614. doi: 10.1002/ps.1735
- Hershenhorn, J., Eizenberg, H., Dor, E., Kapulnik, Y., and Goldwasser, Y. (2009). *Phelipanche aegyptiaca* management in tomato. *Weed Res.* 49(Suppl. 1), 34–37. doi: 10.1111/j.1365-3180.2009.00739.x
- Hershenhorn, J., Goldwasser, Y., Plakhine, D., Lavan, Y., Blumenfeld, T., Bucsbaum, H., et al. (1998). Orobanche aegyptiaca control in tomato fields with sulfonylurea herbicides. Weed Res. 38, 343–349. doi: 10.1046/j.1365-3180.1998.00105.x
- Hibberd, J. M., Quick, W. P., Press, M. C., and Scholes, J. D. (1998). Can source-sink relations explain responses of tobacco to infection by the root holoparasitic angiosperm *Orobanche cernua? Plant Cell Environ.* 21, 333–340. doi: 10.1046/i.1365-3040.1998.00272.x
- Hibberd, J. M., Quick, W. P., Press, M. C., Scholes, J. D., and Jeschke, W. D. (1999).
  Solute fluxes from tobacco to the parasitic angiosperm *Orobanche cernua* and the influence of infection on host carbon and nitrogen relations. *Plant Cell Environ*. 22, 937–947. doi: 10.1046/j.1365-3040.1999.00462.x
- Hiraoka, Y., Ueda, H., and Sugimoto, Y. (2009). Molecular responses of Lotus japonicus to parasitism by the compatible species Orobanche aegyptiaca and the incompatible species Striga hermonthica. J. Exp. Bot. 60, 641–650. doi: 10.1093/jxb/ern316
- Jain, R., and Foy, C. L. (1992). Nutrient effects on parasitism and germination of Egyptian broomrape (Orobanche aegyptiaca). Weed Technol. 6, 269–275.
- Joel, D. M. (2000). The long-term approach to parasitic weeds control: manipulation of specific developmental mechanisms of the parasite. *Crop Prot.* 19, 753–758. doi: 10.1016/S0261-2194(00)00100-9
- Joel, D. M. (2009). The new nomenclature of Orobanche and Phelipanche. Weed Res. 49, 6–7. doi: 10.1111/j.1365-3180.2009.00748.x
- Joel, D. M. (2013). "The haustorium and the life cycles of parasitic Orobanchaceae," in *Parasitic Orobanchaceae*, eds D. M. Joel, J. Gressel, and L. J. Musselman (Heidelberg: Springer Berlin), 21–23.
- Joel, D. M., Back, A., Kleifeld, Y., and Gepstein, S. (1991). "Seed conditioning and its role in *Orobanche* seed germination: inhibition by paclobutrazol," in *Progress* in *Orobanche Research. Proceedings of the International Workshop on Orobanche Research*, eds K. Wegmann and L. J. Musselman (Obermarchtal, FRG: Eberhard Karls Universität), 147–156.

- Joel, D. M., Bar, H., Mayer, A. M., Plakhine, D., Ziadne, H., Westwood, J. H., et al. (2012). Seed ultrastructure and water absorption pathway of the root-parasitic plant *Phelipanche aegyptiaca* (Orobanchaceae). *Ann. Bot.* 109, 181–195. doi: 10.1093/aob/mcr261
- Joel, D. M., Chaudhuri, S. K., Plakhine, D., Ziadna, H., and Steffens, J. C. (2011). Dehydrocostus lactone is exuded from sunflower roots and stimulates germination of the root parasite *Orobanche cumana*. *Phytochemistry* 72, 624– 634. doi: 10.1016/j.phytochem.2011.01.037
- Joel, D. M., Hershenhorn, J., Eizenberg, H., Aly, R., Ejeta, G., Rich, P. J., et al. (2007). Biology and management of weedy root parasites. *Hortic. Rev.* 33, 267–349. doi: 10.1002/9780470168011.ch4
- Joel, D. M., Kleifeld, Y., Losner-Goshen, D., Herzlinger, G., and Gressel, J. (1995).
  Transgenic crops against parasites. *Nature* 374, 220–221. doi: 10.1038/374220a0
- Joel, D. M., and Losner-Goshen, D. (1994). The attachment organ of the parasitic angiosperms *Orobanche cumana* and *O. aegyptiaca* and its development. *Can. J. Bot.* 72, 564–574. doi: 10.1139/b94-075
- Joel, D. M., and Portnoy, V. H. (1998). The angiospermous root parasite *Orobanche* L. (Orobanchaceae) induces expression of a pathogenesis related (PR) gene in susceptible tobacco roots. *Ann. Bot.* 81, 779–781. doi: 10.1006/anbo.1998.0629
- Johnson, A. W., Rosebery, G., and Parker, C. (1976). A novel approach to Striga and Orobanche control using synthetic germination stimulants. Weed Res. 16, 223–227. doi: 10.1111/j.1365-3180.1976.tb00406.x
- Katan, J. (1981). Solar heating (solarization) control of soilborne pests. Ann. Rev. Phytopathol. 19, 211–236. doi: 10.1146/annurev.py.19.090181.001235
- Kebreab, E., and Murdoch, A. J. (1999). A quantitative model for loss of primary dormancy and induction of secondary dormancy in imbibed seeds of Orobanche spp. J. Exp. Bot. 50, 211–219. doi: 10.1093/jxb/50.331.211
- Kebreab, E., and Murdoch, A. J. (2001). Simulation of integrated control strategies for *Orobanche* spp. based on a life cycle model. *Exp. Agric.* 37, 37–51. doi: 10.1017/S001447970100401X
- Keyes, W. J., O'Malley, R. C., Kim, D., and Lynn, D. G. (2000). Signalling organogenesis in parasitic angiosperms: xenognosin generation, perception, and response. J. Plant Growth Regul. 19, 217–231.
- Keyes, W. J., Palmer, A. G., Erbil, W. K., Taylor, J. V., Apkarian, R. P., Weeks, E. R., et al. (2007). Semagenesis and the parasitic angiosperm *Striga asiatica*. *Plant J.* 51, 707–716. doi: 10.1111/j.1365-313X.2007.03171.x
- Klein, O., and Kroschel, J. (2002). Biological control of Orobanche spp. with Phytomyza orobanchia, a review. Biocontrol 47, 245–277. doi: 10.1023/A:1015654429456
- Kroschel, J., Mueller-Stoever, D., Elzein, A., and Sauerborn, J. (2000). "The development of mycoherbicides for the management of parasitic weeds of the genus Striga and Orobanche—a review and recent results," in Proceedings of the X International Symposium on Biological Control of Weeds, ed. N. R. Spencer (Bozeman, MT: Montana State University), 139.
- Kuijt, J. (1969). The Biology of Parasitic Fowering Plants. Berkeley, CA: University of California Press.
- Kusumoto, D., Goldwasser, Y., Xie, X., Yoneyama, K., and Takeuchi, Y. (2007). Resistance of red clover (*Trifolium pratense*) to the root parasitic plant *Orobanche minor* is activated by salicylate but not by jasmonate. *Ann. Bot.* 100, 537–544. doi: 10.1093/annbot/mcm148
- Labrousse, P., Arnaud, M. C., Griveau, Y., Fer, A., and Thalouarn, P. (2004). Analysis of resistance criteria of sunflower recombined inbred lines against *Orobanche cumana* Wallr. Crop Prot. 23, 407–413. doi: 10.1016/j.cropro.2003.09.013
- Labrousse, P., Arnaud, M. C., Seryes, H., Berville, A., and Thalouarn, P. (2001). Several mechanisms are involved in resistance of *Helianthus* to *Orobanche cumana* Wallr. *Ann. Bot.* 88, 859–868. doi: 10.1006/anbo.2001.1520
- Labrousse, P., Delmail, D., Arnaud, M. C., and Thalouarn, P. (2010). Mineral nutrient concentration influences sunflower infection by broomrape (Orobanche cumana). Botany 88, 839–849. doi: 10.1139/B10-057
- Lechat, M. M., Brun, G., Montiel, G., Veronesi, C., Simier, P., Thoiron, S., et al. (2015). Seed response to strigolactone is controlled by abscisic acid-independent DNA methylation in the obligate root parasitic plant, *Phelipanche ramosa* L. Pomel. *J. Exp. Bot.* 6, 3129–3140. doi: 10.1093/jxb/erv119
- Lechat, M. M., Pouvreau, J. B., Péron, T., Gauthier, M., Montiel, G., Veronesi, C., et al. (2012). PrCYP707A1, an ABA catabolic gene, is a key component of *Phelipanche ramosa* seed germination in response to the strigolactone analogue GR24. *J. Exp. Bot.* 63, 5311–5322. doi: 10.1093/jxb/ers189

- Lee, J. A., and Stewart, G. R. (1978). Ecological aspects of nitrogen assimilation. *Adv. Bot. Res.* 6, 1–43. doi: 10.1016/S0065-2296(08)60328-6
- Lieberman, M. (1979). Biosynthesis and action of ethylene. *Annu. Rev. Plant Physiol.* 30, 533–591. doi: 10.1146/annurev.pp.30.060179. 002533
- Linke, K. H., and Saxena, M. C. (1991). "Study on viability and longevity of Orobanche seed under laboratory conditions," in Proceedings of the International Workshop on Orobanche Research: Progress in Orobanche Research, eds K. Wegmann and L. J. Musselman (Obermarchtal: Eberhard-Karls Universitat), 110–114.
- Lins, R. D., Colquhoun, J. B., and Mallory-Smith, C. A. (2007). Effect of small broomrape (*Orobanche minor*) on red clover growth and dry matter partitioning. Weed Sci. 55, 517–520. doi: 10.1614/WS-07-049.1
- Liu, Q., Zhang, Y., Matusova, R., Charnikhova, T., Amini, M., Jamil, M., et al. (2014). Striga hermonthica MAX2 restores branching but not the very low fluence response in the Arabidopsis thaliana MAX2 mutant. New Phytol. 202, 531–541. doi: 10.1111/nph.12692
- Logan, D., and Stewart, G. R. (1995). Thidiazuron stimulates germination and ethylene production in *Striga hermonthica* comparison with the effects of GR24, ethylene and 1-aminocyclopropane-1-carboxylic acid. *Seed Sci. Res.* 5, 99–108. doi: 10.1017/S0960258500002671
- López-Bellido, R. J., Benítez-Vega, J., and López-Bellido, L. (2009). No-tillage improves broomrape control with glyphosate in faba-bean. *Agron. J.* 101, 1394–1399. doi: 10.2134/agronj2009.0014
- López-Granados, F., and García-Torres, L. (1996). Effects of environmental factors on dormancy and germination of crenate broomrape (*Orobanche crenata*). Weed Sci. 44, 284–289.
- López-Granados, F., and García-Torres, L. (1999). Longevity of crenate broomrape (Orobanche crenata) seed under soil and laboratory conditions. Weed Sci. 47, 161–166.
- López-Ráez, J. A., Charnikhova, T., Fernandez, I., Bouwmeester, H., and Pozo, M. J. (2011). Arbuscular mycorrhizal symbiosis decreases strigolactone production in tomato. J. Plant Physiol. 168, 294–297. doi: 10.1016/j.jplph.2010. 08.011
- Losner-Goshen, D., Portnoy, V. H., Mayer, A. M., and Joel, D. M. (1998).
  Pectolytic activity by the haustorium of the parasitic plant *Orobanche L*. (Orobanchaceae) in host roots. *Ann. Bot.* 81, 319–326. doi: 10.1006/anbo. 1997.0563
- Louarn, J., Carbonne, F., Delavault, P., Becard, G., and Rochange, S. (2012). Reduced germination of *Orobanche cumana* seeds in the presence of arbuscular mycorrhizal fungi or their exudates. *PLoS ONE* 7:e49273. doi: 10.1371/journal.pone.0049273
- Mabrouk, Y., Mejri, S., Hemissi, I., Simier, P., Delavault, P., Saidi, M., et al. (2010). Bioprotection mechanisms of pea plant by *Rhizobium leguminosarum* against Orobanche crenata. Afr. J. Microbiol. Res. 4, 2570–2575.
- Mabrouk, Y., Simier, P., Arfaoui, A., Sifi, B., Delavault, P., Zourgui, L., et al. (2007a). Induction of phenolic compounds in pea (*Pisum sativum L.*) inoculated by *Rhizobium leguminosarum* and infected with *Orobanche crenata*. *J. Phytopathol*. 155, 728–734. doi: 10.1111/j.1439-0434.2007.01307.x
- Mabrouk, Y., Simier, P., Delavault, P., Delgrange, S., Sifi, B., Zourgui, L., et al. (2007b). Molecular and biochemical mechanisms of defence induced in pea by *Rhizobium leguminosarum* against *Orobanche crenata*. Weed Res. 47, 452–460. doi: 10.1111/j.1365-3180.2007.00583.x
- Mabrouk, Y., Zourgui, L., Sifi, B., Delavault, P., Simier, P., and Belhadj, O. (2007c). Some compatible *Rhizobium leguminosarum* strains in peas decrease infections when parasitized by *Orobanche crenata*. Weed Res. 47, 44–53. doi: 10.1111/j.1365-3180.2007.00548.x
- Manschadi, A. M., Kroschel, J., and Sauerborn, J. (1996). Dry matter production and partitioning in the host-parasite association *Vicia faba–Orobanche crenata*. *J. Appl. Bot.* 70, 224–229.
- Matvienko, M., Wojtowicz, A., Wrobel, R., Jamison, D., Goldwasser, Y., and Yoder, J. I. (2001). Quinone oxidoreductase message levels are differentially regulated in parasitic and non-parasitic plants exposed to allelopathic quinones. *Plant J.* 25, 375–387. doi: 10.1046/j.1365-313x.2001.00971.x
- Mauro, R. P., Lo Monaco, A., Lombardo, S., Restuccia, A., and Mauromicale, G. (2015). Eradication of *Orobanche/Phelipanche* spp. seedbank by soil solarization and organic supplementation. *Sci. Hortic.* 193, 62–68. doi: 10.1016/j.scienta.2015.06.038

- Mauromicale, G., Lo Monaco, A., and Longo, M. G. A. (2008). Effect of branched broomrape (*Orobanche ramosa*) infection on the growth and photosynthesis of tomato. *Weed Sci.* 56, 574–581. doi: 10.1614/WS-07-147.1
- Mauromicale, G., Restuccia, G., and Marchese, A. (2001). Soil solarization, a non-chemical technique for controlling *Orobanche crenata* and improving yield of faba bean. *Agronomie* 21, 757–765. doi: 10.1051/agro:2001167
- Mayer, A. M., and Bar-Nun, N. (1994). "Metabolism during preconditioning and germination of Orobanche aegyptiaca," in Proceedings of the 3rd International Workshop on Orobanche and related Striga Research: Biology and management of Orobanche, eds A. H. Pieterse, J. A. C. Verkleij, and S. C. ter Borg (Amsterdam: Royal Tropical Institute), 146–156.
- Mayer, A. M., and Bar-Nun, N. (1997). "Germination of *Orobanche* seeds: some aspects of metabolism during preconditioning," in *Basic and Applied Aspects of Seed Biology*, eds R. H. Ellis, M. Black, A. J. Murdoch, and T. D. S. Hing (Dordrecht: Kluwer Academic Publishers), 633–639.
- McNally, S. F., Orebamjo, T. O., Hirel, B., and Stewart, G. R. (1983). Glutamine synthetase isozymes of *Striga hermonthica* and other angiosperm root parasites. *J. Exp. Bot.* 34, 610–619. doi: 10.1093/jxb/34.5.610
- Metzger, J. (1983). The promotion of germination of dormant weed seeds by substituted phthalimides and gibberellic acid. Weed Sci. 31, 285–289.
- Mohamed, K. I., Papes, M., Williams, R., Benz, B. W., and Peterson, A. T. (2006). Global invasive potential of 10 parasitic witchweeds and related Orobanchaceae. Ambio 35, 281–288. doi: 10.1579/05-R-051R.1
- Müller-Stöver, D. (2001). Possibilities of biological control of Orobanche crenata and O. cumana with Ulocladium botrytis and Fusarium oxysporum f. sp. orthoceras. Agroecology 3, 174.
- Müller-Stöver, D., Buschmann, H., and Sauerborn, J. (2005). Increasing control reliability of *Orobanche cumana* through integration of a biocontrol agent with a resistance-inducing chemical. *Eur. J. Plant Pathol.* 111, 193–202. doi: 10.1007/s10658-004-2814-8
- Murdoch, A. J., and Kebreab, A. (2013). "Germination ecophysiology," in *Parasitic Orobanchaceae*, eds D. M. Joel, J. Gressel, and L. J. Musselman (Heidelberg: Springer Berlin), 195–219.
- Musselman, L. J. (1980). The biology of Striga, Orobanche and other root parasitic weeds. Annu. Rev. Phytopathol. 18, 463–489. doi: 10.1146/annurev.py.18.090180.002335
- Musselman, L. J., and Dickison, W. C. (1975). The structure and development of the haustorium in parasitic Scrophulariaceae. *Bot. J. Linn. Soc.* 70, 183–212. doi: 10.1111/j.1095-8339.1975.tb01645.x
- Mwakaboko, A. S., and Zwanenburg, B. (2011). Strigolactone analogs derived from ketones using a working model for germination stimulants as a blueprint. *Plant Cell Physiol.* 52, 699–715. doi: 10.1093/pcp/pcr031
- Nandula, V. K., Foster, J. G., and Foy, C. L. (2000). Impact of Egyptian broomrape (*Orobanche aegyptiaca* (Pers.) parasitism on amino acid composition of carrot (*Daucus carota* L). *J. Agric. Food Chem.* 48, 3930–3934. doi: 10.1021/jf991145w
- Panetta, F. D., and Lawes, R. (2005). Evaluation of weed eradication programs: the delimitation of extent. *Divers. Distrib.* 11, 435–442. doi: 10.1111/j.1366-9516.2005.00179.x
- Parker, C. (2009). Observations on the current status of *Orobanche* and *Striga* problems worldwide. *Pest Manag. Sci.* 65, 453–459. doi: 10.1002/ps.1713
- Parker, C. (2014). Orobanche crenata in UK- an update. Haustorium 65, 5-6.
- Parker, C., and Riches, C. R. (1993). Parasitic Weeds of the World: Biology and Control. Wallingford: CAB International.
- Pérez-de-Luque, A., Fondevilla, S., Pérez-Vich, B., Aly, R., Thoiron, S., Simier, P., et al. (2009). Understanding *Orobanche* and *Phelipanche*-host plant interactions and developing resistance. *Weed Res.* 49, 8–22. doi: 10.1111/j.1365-3180.2009.00738 v.
- Pérez-de-Luque, A., Jorrín, J., Cubero, J. I., and Rubiales, D. (2005). Resistance and avoidance against *Orobanche crenata* in pea (*Pisum* spp.) operate at different developmental stages of the parasite. Weed Res. 45, 379–387. doi: 10.1111/i.1365-3180.2005.00464.x
- Pérez-de-Luque, A., Jorrín, J., and Rubiales, D. (2004). Broomrape (*Orobanche crenata* Forsks.) control in pea (*Pisum sativum* L.) by foliar applications of benzothiadiazole (BTH). *Phytoparasitica* 32, 21–29. doi: 10.1007/BF02980855
- Pérez-de-Luque, A., Moreno, M. T., and Rubiales, D. (2008). Host plant resistance against broomrapes (*Orobanche* spp.): defence reactions and mechanisms of resistance. *Ann. Appl. Biol.* 152, 131–141. doi: 10.1111/j.1744-7348.2007.00212.x

- Pérez-de-Luque, A., and Rubiales, D. (2009). Nanotechnology for parasitic plant control. *Pest Manang. Sci.* 65, 540–545. doi: 10.1002/ps.1732
- Pérez-Vich, B., Velasco, L., Rich, P. J., and Ejeta, G. (2013). "Marker-assisted and physiology-based breeding for resistance to root parasitic Orobanchaceae," in *Parasitic Orobanchaceae*, eds D. M. Joel, J. Gressel, and L. J. Musselman (Heidelberg: Springer Berlin), 369–391.
- Péron, T., Véronési, C., Mortreau, E., Pouvreau, J. B., Thoiron, S., Leduc, N., et al. (2012). Role of the sucrose synthase encoding PrSus1 gene in the development of the parasitic plant *Phelipanche ramosa* L. (Pomel). *Mol. Plant Microbe Interact*. 25, 402–411. doi: 10.1094/MPMI-10-11-0260
- Plakhine, D., Eizenberg, H., Hershenhorn, J., Goldwasser, Y., and Kleifeld, Y. (2001). "Control of Orobanche aegyptiaca with sulfonylurea herbicides in tomato—polyethylene bag studies," in International Parasitic Weed Symposium, eds A. Fer, P. Thalouarn, D. M. Joel, C. Musselman, and J. A. C. Verkleij (Nantes: University of Nantes), 294–295.
- Plakhine, D., Ziadna, H., and Joel, D. M. (2009). Is seed conditioning essential for Orobanche germination? Pest Manag. Sci. 65, 492–496. doi: 10.1002/ps.1716
- Privat, G. (1960). Recherches sur les phanerogames parasites (etude d'*Orobanche* hederae Duby). *Ann. Sci. Nat. Bot.* 12, 722–865.
- Quimby, P. C. Jr., Zidack, N. K., and Boyette, C. D. (1999). A simple method for stabilizing and granulating fungi. *Biocontrol Sci. Technol.* 9, 5–8. doi: 10.1080/09583159929857
- Reda, F. (2006). Orobanche crenata in Ethiopia. Haustorium 49, 3.
- Rich, P. J., Grenier, C., and Ejeta, G. (2004). Striga resistance in the wild relatives of sorghum. Crop Sci. 44, 2221–2229. doi: 10.2135/cropsci2004.2221
- Riopel, J. L., and Timko, M. P. (1995). "Haustorial initiation and differentiation," in Parasitic Plants, eds M. C. Press and J. D. Graves (London: Chapman and Hall), 39–79.
- Rubiales, D., Alcántara, C., Pérez-de-Luque, A., Gil, J., and Sillero, J. C. (2003a). Infection of chickpea (*Cicer arietinum*) by crenate broomrape (*Orobanche crenata*) as influenced by sowing date and weather conditions. *Agronomie* 23, 359–362. doi: 10.1051/agro:2003016
- Rubiales, D., Pérez-de-Luque, A., Joel, D. M., Alcantara, C., and Sillero, J. C. (2003b). Characterization of resistance in chickpea to crenate broomrape (*Orobanche crenata*). Weed Sci. 51, 702–707. doi: 10.1614/P2002-151
- Rubiales, D., Fernández-Aparicio, M., Pérez-de-Luque, A., Castillejo, M. A., Prats, E., Sillero, J., et al. (2009a). Breeding approaches for crenate broomrape (*Orobanche crenata* Forsk.) management in pea (*Pisum sativum* L.). Pest Manag. Sci. 65, 553–559. doi: 10.1002/ps.1740
- Rubiales, D., Fernández-Aparicio, M., Wegmann, K., and Joel, D. (2009b). Revisiting strategies for reducing the seedbank of *Orobanche* and *Phelipanche* spp. Weed Res. 49, 23–33. doi: 10.1111/j.1365-3180.2009.00742.x
- Rubiales, D., Fernandez-Aparicio, M., and Rodriguez, M. J. (2008). First report of crenate broomrape (*Orobanche crenata*) on lentil (*Lens culinaris*) and common vetch (*Vicia sativa*) in Salamanca Province, Spain. *Plant Dis.* 92, 1368. doi: 10.1094/PDIS-92-9-1368B
- Saghir, A. R. (1986). "Dormancy and germination of Orobanche seeds in relation to control methods," in Proceedings of a Workshop in Wageningen: Biology and Control of Orobanche, ed. S. J. Ter Borg (Wageningen: LH/VPO), 25–34.
- Sands, D. C., and Pilgeram, A. L. (2009). Methods for selecting hypervirulent biocontrol agents of weeds: why and how? *Pest Manag. Sci.* 65, 581–587. doi: 10.1002/ps.1739
- Sarosh, B. R., Sivaramakrishnan, S., and Shetty, H. S. (2005). Elicitation of defense related enzymes and resistance by L-methionine in pearl millet against downy mildew disease caused by Sclerospora graminicola. Plant Physiol. Biochem. 43, 808–815. doi: 10.1016/j.plaphy.2005.06.009
- Sauerborn, J. (1991). "The economic importance of the phytoparasites Orobanche and Striga," in Proceedings of the Fifth Symposium on Parasitic Weeds, Nairobi, eds J. K. Ransom, L. J. Musselman, A. D. Worsham, and C. Parker (Nairobi: CIMMYT), 137–143.
- Sauerborn, J., Linke, K. H., Saxena, M. C., and Koch, W. (1989). Solarization, a physical control method for weeds and parasitic plants (*Orobanche* spp.) in Mediterranean agriculture. Weed Res. 29, 391–393. doi: 10.1111/j.1365-3180.1989.tb01310.x
- Schneeweiss, G. M. (2007). Correlated evolution of life history and host range in the nonphotosynthetic parasitic flowering plants *Orobanche* and *Phelipanche* (Orobanchaceae). *J. Evol. Biol.* 20, 471–478. doi: 10.1111/j.1420-9101.2006.01273.x

- Seo, M., Nambara, E., Choi, G., and Yamaguchi, S. (2009). Interaction of light and hormone signals in germinating seeds. *Plant Mol. Biol.* 69, 463–472. doi: 10.1007/s11103-008-9429-y
- Sholmer-Ilan, A. (1993). Germinating seeds of the root parasite *Orobanche aegyptiaca* Pers. excrete enzymes with carbohydrase activity. *Symbiosis* 15, 61–70
- Sillero, J. C., Moreno, M. T., and Rubiales, D. (2005). Sources of resistance to crenate broomrape among species of Vicia. Plant Dis. 89, 23–27. doi: 10.1094/PD-89-0023
- Singh, A., and Singh, M. (1993). Cell wall-degrading enzyme in Orobanche aegyptiaca and its host Brassica campestris. Physiol. Plant. 89, 177–181. doi: 10.1111/j.1399-3054.1993.tb01802.x
- Slavov, S., Valkov, V., Batchvarova, R., Atanassova, S., Alexandrova, M., and Atanassov, A. (2005). Chlorsulfuron resistant transgenic tobacco as a tool for broomrape control. *Transgenic Res.* 14, 273–278. doi: 10.1007/s11248-004-8081-9
- Song, W. J., Zhou, W. J., Jin, Z. L., Cao, D. D., Joel, D. M., Takeuchi, Y., et al. (2005). Germination response of *Orobanche* seeds subjected to conditioning temperature, water potential and growth regulator treatments. *Weed Res.* 45, 467–476. doi: 10.1111/j.1365-3180.2005.00477.x
- Southwood, O. R. (1971). The effects of superphosphate application, 2,4-DB and grazing on broomrape (*Orobanche minor* Sm.) in a subterranean clover pasture. *Weed Res.* 11, 240–246. doi: 10.1111/j.1365-3180.1971.tb01006.x
- Stewart, G. R., and Press, M. C. (1990). The physiology and biochemistry of parasitic angiosperms. Annu. Rev. Plant Physiol. Plant Mol. Biol. 41, 127–151. doi: 10.1146/annurev.pp.41.060190.001015
- Suttle, J. C. (1983). N-substituted phthalimides as plant bioregulants. News Bull. Br. Plant Growth Regul. Group 6, 11–19.
- Suttle, J. C., and Schreiner, D. R. (1982). The biological activity of AC-94, 377 [1-(3-chlorophthalimido)-cyclohexane-arboxamide]. *J. Plant Growth Regul.* 1, 139–146.
- Takeuchi, Y., Omigawa, Y., Ogasawara, M., Yoneyama, K., Konnai, M., and Worsham, A. D. (1995). Effects of brassinosteroids on conditioning and germination of clover broomrape (*Orobanche minor*) seeds. *Plant Growth Regul.* 16, 153–160. doi: 10.1007/BF00029536
- Tan, S., Evans, R. R., Dahmer, M. L., Sing, B. K., and Shaner, D. (2005). Imidazolinone-tolerant crops: history, current status and future. *Pest Manag. Sci.* 61, 246–257. doi: 10.1002/ps.993
- Tank, D. C., Beardsley, P. M., Kelchner, S. A., and Olmstead, R. G. (2006). Review of the systematics of Scrophulariaceae s.l. and their current disposition. *Aust. Syst. Bot.* 19, 289–307. doi: 10.1071/SB05009
- Thomas, H., Heller, A., Sauerborn, J., and Müller-Stöver, D. (1999). Fusarium oxysporum f. sp. orthoceras, a potential mycoherbicide, parasitizes seeds of Orobanche cumana (Sunflower broomrape): a cytological study. Ann. Bot. 83, 453–458. doi: 10.1006/anbo.1998.0847
- Toh, S., Kamiya, Y., Kawakami, N., Nambara, E., McCourt, P., and Tsuchiya, Y. (2012). Thermoinhibition uncovers a role for strigolactones in *Arabidopsis* seed germination. *Plant Cell Physiol.* 53, 107–117. doi: 10.1093/pcp/pcr176
- Tomilov, A., Tomilova, N., Shin, D. H., Jamison, D., Torres, M., Reagan, R., et al. (2006). "Chemical signalling between plants: mechanistic similarities between phytotoxic allelopathy and host recognition by parasitic plants," in *Chemical Ecology: From Gene to Ecosystem*, eds M. Dicke and W. Takken (Dordrecht: Springer), 55–69.
- Umehara, M., Hanada, A., Yoshida, S., Akiyama, K., Arite, T., Takeda-Kamiya, N., et al. (2008). Inhibition of shoot branching by new terpenoid plant hormones. *Nature* 455, 195–200. doi: 10.1038/nature07272
- USEPA (2004). List of Inert Pesticide Ingredients List 4b. U. S. Environmental Protection Agency. Available at: www.epa.gov/opprd001/inerts/inerts\_list4Bname.pdf
- Van Delft, G. J., Graves, J. D., Fitter, A. H., and Van Ast, A. (2000). Striga seed avoidance by deep planting and no-tillage in sorghum and maize. Int. J. Pest Manag. 46, 251–256. doi: 10.1080/09670870050206019
- van Hezewijk, M. J., and Verkleij, J. A. C. (1996). The effect of nitrogenous compounds on in vitro germination of *Orobanche crenata* Forsk. *Weed Res.* 36, 395–404. doi: 10.1111/j.1365-3180.1996.tb01669.x
- Vaucher, J. P. (1823). Mémoire sur la Germination des Orobanches, Vol. 10. Paris: Mémoires du Museum d'Histoire Naturelle, 261–273.

- Veronesi, C., Bonnin, E., Benharrat, H., Fer, A., and Thalouarn, P. (2005). Are pectinolytic activities of *Orobanche cumana* seedlings related to virulence towards sunflower? *Isr. J. Plant Sci.* 53, 19–27. doi: 10.1560/ETEL-C34X-Y6MG-YT0M
- Veronesi, C., Bonnin, E., Calvez, S., Thalouarn, P., and Simier, P. (2007). Activity of secreted cell wall-modifying enzymes and expression of peroxidase-encoding gene following germination of *Orobanche ramosa*. *Biol. Plant* 51, 391–394. doi: 10.1007/s10535-007-0084-y
- Vurro, M., Boari, A., Evidente, A., Andolfi, A., and Zermane, N. (2009). Natural metabolites for parasitic weed management. *Pest Manag. Sci.* 65, 566–571. doi: 10.1002/ps.1742
- Vurro, M., Boari, A., Pilgeram, A. L., and Sands, D. C. (2006). Exogenous amino acids inhibit seed germination and tubercle formation by *Orobanche ramosa* (broomrape): potential application for management of parasitic weeds. *Biol. Control* 36, 258–265. doi: 10.1016/j.biocontrol.2005.09.017
- Westwood, J. H. (2013). "The physiology of the established parasite-host association," in *Parasitic Orobanchaceae*, eds D. M. Joel, J. Gressel, and L. J. Musselman (Berlin: Springer), 87–114.
- Westwood, J. H., dePamphilis, C. W., Das, M., Fernández-Aparicio, M., Honaas, L. A., Timko, M. P., et al. (2012). The parasitic plant genome project: new tools for understanding the biology of *Orobanche* and *Striga. Weed Sci.* 60, 295–306. doi: 10.1614/WS-D-11-00113.1
- Westwood, J. H., and Foy, C. L. (1999). Influence of nitrogen on germination and early development of broomrape (*Orobanche* spp.). *Weed Sci.* 47, 2–7.
- Westwood, J. H., Yu, X., Foy, C. L., and Cramer, C. L. (1998). Expression of a defense-related 3-hydroxy-3-methylglutaryl CoA reductase gene in response to parasitism by *Orobanche* spp. *Mol. Plant Microbe Interact.* 11, 530–536. doi: 10.1094/MPMI.1998.11.6.530
- Xie, X., Yoneyama, K., and Yoneyama, K. (2010). The strigolactone story. *Annu. Rev. Phytopathol.* 48, 93–117. doi: 10.1146/annurev-phyto-073009-114453
- Yang, Z., Wafula, E. K., Honaas, L. A., Zhang, H., Das, M., Fernandez-Aparicio, M., et al. (2015). Comparative transcriptome analyses reveal core parasitism genes and suggest gene duplication and repurposing as sources of structural novelty. *Mol. Biol. Evol.* 32, 767–790. doi: 10.1093/molbev/msu343
- Yoder, J. I., and Scholes, J. D. (2010). Host plant resistance to parasitic weeds; recent progress and bottlenecks. Curr. Opin. Plant Biol. 13, 478–484. doi: 10.1016/j.pbi.2010.04.011

- Yoneyama, K., Xie, X., Kim, H. I., Kisugi, T., Nomura, T., Sekimoto, H., et al. (2012). How do nitrogen and phosphorus deficiencies affect strigolactone production and exudation? *Planta* 235, 1197–1207. doi: 10.1007/s00425-011-1568-8
- Yoneyama, K., Xie, X., Kusumoto, D., Sekimoto, H., Sugimoto, Y., Takeuchi, Y., et al. (2007a). Nitrogen deficiency as well as phosphorus deficiency in sorghum promotes the production and exudation of 5deoxystrigol, the host recognition signal for arbuscular mycorrhizal fungi and root parasites. *Planta* 227, 125–132. doi: 10.1007/s00425-007-0600-5
- Yoneyama, K., Yoneyama, K., Takeuchi, Y., and Sekimoto, H. (2007b).
  Phosphorus deficiency in red clover promotes exudation of orobanchol, the signal for mycorrhizal symbionts and germination stimulant for root parasites. *Planta* 225, 1031–1038. doi: 10.1007/s00425-006-0410-1
- Zehhar, N., Ingouff, M., Bouya, D., and Fer, A. (2002). Possible involvement of gibberellins and ethylene in *Orobanche ramosa* germination. *Weed Res.* 42, 464–469. doi: 10.1046/j.1365-3180.2002. 00306.x
- Zhang, Y., Luc, J. E., and Crow, W. T. (2010). Evaluation of amino acids as turfgrass nematicides. J. Nematol. 42, 292–297.
- Zwanenburg, B., Mwakaboko, A. S., Reizelman, A., Anilkuma, G., and Sethumadhavan, D. (2009). Structure and function of natural and synthetic signaling molecules in parasitic weed germination. *Pest Manag. Sci.* 65, 478–491. doi: 10.1002/ps.1706
- **Conflict of Interest Statement:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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