**Agrobacterium-mediated gene transfer: recent advancements and layered immunity in plants**

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**Abstract**

**Main conclusion** Plant responds to Agrobacterium via three-layered immunity that determines its susceptibility or resistance to Agrobacterium infection.

**Abstract** Agrobacterium tumefaciens is a soil-borne Gram-negative bacterium that causes crown gall disease in plants. The remarkable feat of interkingdom gene transfer has been extensively utilised in plant biotechnology to transform plant as well as non-host systems. In the past two decades, the molecular mode of the pathogenesis of A. tumefaciens has been extensively studied. Agrobacterium has also been utilised as a premier model to understand the defence response of plants during plant–Agrobacterium interaction. Nonetheless, the threat of Agrobacterium-mediated crown gall disease persists and is associated with a huge loss of plant vigour in agriculture. Understanding the molecular dialogues between these two interkingdom species might provide a cure for crown gall disease. Plants respond to A. tumefaciens by mounting a three-layered immune response, which is manipulated by Agrobacterium via its virulence effector proteins. Comparative studies on plant defence proteins versus the counter-defence of Agrobacterium have shed light on plant susceptibility and tolerance. It is possible to manipulate a plant’s immune system to overcome the crown gall disease and increase its competence via A. tumefaciens-mediated transformation. This review summarises the recent advances in the molecular mode of Agrobacterium pathogenesis as well as the three-layered immune response of plants against Agrobacterium infection.

**Keywords** Agrobacterium tumefaciens · Plant–Agrobacterium interaction · Plant immunity

**Introduction**

Agrobacterium tumefaciens is a soil-borne, Gram-negative bacterium that infects and causes tumours, called crown gall disease, in a variety of plant species. More than a century ago, Agrobacterium was isolated from a crown gall tumour by two plant pathologists Smith and Townsend (Smith and Townsend 1907). There were originally three biovars of pathogenic Agrobacteria based upon the host range and manner of pathogenic response in the host. Biovar I includes A. tumefaciens, biovar II includes A. rhizogenes and biovar III includes A. vitis (Slater et al. 2009). However, research on the taxonomic classification and position of Agrobacterium is still ongoing (Gan and Savka 2018; Ormeno-Orrillo et al. 2015). Among all biovars, biovar I, Agrobacterium (C58), was the first whose genome was sequenced and made available in the database (https://www.ncbi.nlm.nih.gov/genome), where it was renamed as A. fabrum. Furthermore, Shams et al. used another synonym for Agrobacterium, i.e. Rhizobium radiobacter (Shams et al. 2013). For the convenience of the readers, the older name, i.e. A. tumefaciens will be used in this review because it is predominantly used in ongoing research and review articles.

There are two distinct lifestyles of Agrobacterium in nature: one is free-living, saprophytic and non-pathogenic and the other one is pathogenic, based on plant tissue as its...
ecological niche instead of soil (Meyer et al. 2018). In bacteria, the change between these two lifestyles is coordinated with the change in the gene expression pattern resulting from the perception of environmental cues (Duprey et al. 2014; Valentini et al. 2018). *Agrobacterium* begins its pathogenic lifestyle when it perceives signals from wounded plant cells. There is a large tumour-inducing (Ti) plasmid in *Agrobacterium*, which confers pathogenicity. All Ti plasmids contain at least four gene clusters or operons. These operons possess different functions. For instance, the Vir operon contains all the virulence genes that get activated during the pathogenic process, the repABC operon maintains the replication and separation of the Ti plasmid, the tra operon facilitates the conjugation of DNA and the trb operon synthesis the secretion system required when transmitting pTi from one to another bacterium (Wetzel et al. 2015). The virulence protein processes the T-DNA region of the Ti plasmid in response to an environmental signal. During processing, only 25 base pair direct repeats at the left and right borders of the T-DNA are processed. The T-strand, along with several virulence proteins, enters the host cytoplasm, travels towards the nucleus and is integrated into the plant genome (Nester 2015; Gelvin 2017). As T-DNA harbours auxin (iaaM, iaaH), cytokinin (ipt) and opine synthesis genes, it causes hormonal imbalances in plants and results in malignant growth. Opine synthesis genes lead to the production of opines, which serve as a source of nutrition for *Agrobacterium* and create a new ecological niche for it (Lacroix and Citovsky 2013). Recently, *Agrobacterium* fitness gene has been identified, which constitutes 3–8% of its total genes and is important for its competitive survival in plants (Torres et al. 2022).

*Agrobacterium* ranks third among the most pathogenic bacteria, next only to *Pseudomonas syringae* pathovars and *Ralstonia solanacearum* (Mansfield et al. 2012), which severely affect plant growth and vigour during crown gall disease. *Agrobacterium* has been known for several years as a plant pathogen. Studies have reported that the native T-DNA of *Agrobacterium* can be replaced with any gene of interest and that it can transform plants without causing tumours (Fraley et al. 1983; Caplan et al. 2019).

The unintended plant transformation activity of *Agrobacterium* makes it not only an important plant pathogen but also a potent biotechnological tool. Recent studies have also shown that *Agrobacterium* can transform non-host plants (Song et al. 2019) as well as non-plant systems, such as yeast (Bundock et al. 1995), fungi (De Groot et al. 1998; Li et al. 2017) and even human cells (Lacroix et al. 2006; Lacroix and Citovsky 2018).

Moreover, *Agrobacterium* has been utilised for the production of pharmaceutical proteins in plants (Kopertekh and Schiemann 2019). While *Agrobacterium* is extensively used in the field of plant biotechnology and serves as a model for plant–pathogen interaction, molecular mode of pathogenesis, etc., some plants are recalcitrant to transformation. The recalcitrance depends on the type of *Agrobacterium* used, the type of plant and the explants (Tzifra and Citovsky 2006). Moreover, the extensive use of *Agrobacterium* in biotechnological industries overshadowed the adverse effect of its natural action, namely crown gall disease. Hence, it is important to understand the innate immune response of plants and the downstream defence activation during plant–*Agrobacterium* interaction.

In nature, plants and pathogens coevolved over time. Plants have evolved a complex and versatile immune response that detects the pathogen with a wide array of receptors. Simultaneously, pathogens have managed to escape from the plant immune system with the help of effector proteins. The innate immune system of a plant is triggered during pathogen attack. The plant does not recognise the whole pathogen but instead detects its signature molecular pattern. In case of pathogens, these molecular patterns are called ‘pathogen-associated molecular pattern (PAMP)’. On the contrary, in the case of non-pathogens, the molecular signals are termed ‘microbe-associated molecular pattern (MAMP)’ (Ausubel 2005). These PAMPs are a part of the general elicitor and are present in a vast group of pathogenic bacteria that are evolutionary stable and essential for the pathogenesis of the microbe. Besides PAMP, damage-associated molecular pattern (DAMP), a product of host cellular damage after pathogen invasion, also triggers the immune response. Plants are able to detect these extracellular and intracellular milieus using a cell surface receptor named pattern recognition receptor (PRR). In this way, the plant immune system serves as a surveillance system for detecting these signals during pathogen attack (Cook et al. 2015; Gust et al. 2017). The recognition of PAMP by PRR is the first level of defence and is referred to as PAMP-triggered immunity (PTI). PTI is a determinant of the plant resistance at an initial level that affects both basal and non-host resistance. Basal resistance denotes the resistance of susceptible plants after getting infected with the adapted pathogen. However, when a plant develops resistance to a non-adapted pathogen, either pathogenic or non-pathogenic, it is called non-host resistance (Couto and Zipfel 2016; Tang et al. 2017). As soon as PAMP perception and PTI activation occur, the downstream signalling event commences immediately and blocks the infection process at an early stage. The downstream signalling events include the reactive oxygen species (ROS) burst and the activation of different kinases, such as the mitogen-activated protein kinase (MAPK) cascade that phosphorylates the defence-related genes (Noman et al. 2019). To overcome PTI, the pathogen makes plant cells more susceptible to infection using numerous virulence effector proteins. This process of pathogen counteraction is called effector triggered susceptibility (ETS). A plant’s
compatibility with the pathogen is further determined by the second level of defence, which is called effector-triggered immunity (ETI). ETI causes a hypersensitive reaction involving localised cell death of the host cells to prevent infection propagation (Janda et al. 2019). This review summarises the latest research on the molecular mechanism of T-DNA transfer in plant cells, with a focus on understanding the plant defence mechanisms and Agrobacterium counter-defence during plant–Agrobacterium interaction.

Molecular mechanism of Agrobacterium-mediated pathogenesis

Agrobacterium is used as a model organism to study plant–pathogen interactions. The pathogenesis of Agrobacterium has been studied extensively at the bacterial level. Despite this study, it is unclear how plants respond to this process. Only a small number of observations have been reported regarding host factors and their importance in Agrobacterium pathogenesis. These observations confirm that Agrobacterium, apart from its virulence protein, utilises the host cellular machinery for T-DNA cytoplasmic trafficking, nuclear import and its integration into the plant genome (Gelvin 2003; Citovsky et al. 2004; Michielse et al. 2004; Li and Pan 2017; Yang et al. 2017). As a pathogen, Agrobacterium has been extensively studied. Its pathogenesis begins with the activation of its virulence genes after being stimulated by a chemical signal released by the plant cell (Gelvin 2003). The entire process is quite complex and has already been examined extensively in many articles. Therefore, this review discusses only the most recent advancements. There are four steps in the infection process: (1) release of chemical signals and the onset of pathogenic lifestyle of Agrobacterium, (2) activation and induction of the virulence gene, (3) generation of the T-complex and its cytoplasmic trafficking and nuclear import and (4) integration of T-DNA into the plant genome and its expression.

Release of chemical signal and onset of pathogenic lifestyle of Agrobacterium

When plants are subjected to biotic or abiotic stresses, the very first response is oxidative burst, which is followed by the production of phenolic compounds and other secondary metabolites (Baker et al. 2020). Acetosyringone (AS), a phenolic metabolite of plants, is most effective (Guo et al. 2017). AS is synthesised via the phenylpropanoid pathway (Maury et al. 2010), and phenylpropanoids are involved in plant defence (Fraser and Chapple 2011). However, Agrobacterium somehow utilises this defence signal to initiate infection. Recent studies on Agrobacterium have identified an antibiotic-resistant RND-type efflux pump called the MexE/MexF/AmeC pump, which enhances the concentration of the inducer required for the induction of virulence genes (Binns and Zhao 2020). Agrobacterium becomes chemotactic and travels towards the wound site once it finds an appropriate signal (Guo et al. 2017). CheW proteins of Agrobacterium are implicated in bridging CheA kinase and chemoreceptor, thus forming a core chemoreceptor complex that facilitates chemotaxis (Huang et al. 2018). When the process is successful, Agrobacterium attaches to the plant cell with the help of binding and attachment proteins (encoded by ChvA, ChvB, PscA and Att) (Tzfira and Citovsky 2002; Cangelosi et al. 2007).

Activation and induction of the virulence gene

The rhizosphere must contain three factors to activate and induce the virulence system of Agrobacterium, namely, low pH, sugar and phenolic compounds (Lacroix and Citovsky 2013). Agrobacterium’s virulence system is triggered by low pH and sugar, but these factors are not necessary to induce the virulence genes. Phenolic compounds, such as hydroxyacetosyringone, induce the virulence genes. Virulence property of an Agrobacterium is determined by its chromosomal virulence genes (cvh) and Ti plasmid virulence genes (vir genes). ExoR, a periplasmic regulator, detects low pH and activates the virulence process. Under neutral pH, ExoR interacts with the chromosome-based two-component regulatory system ChvG. However, under acidic pH condition, ExoR gets cleaved by periplasmic proteases and frees ChvG, which gets autophosphorylated and transfers its phosphate group to the response regulator ChvI (Heckel et al. 2014; Wu et al. 2012; Yuan et al. 2008; Subramoni et al. 2014). ChvI, in turn, activates the other virulence genes and also T6SS (Yuan et al. 2008; Wu et al. 2012), and the Ti plasmid-based virulence protein, VirG (Li et al. 2002; Yuan et al. 2008). It is noteworthy that low pH not only activates the ExoR-ChvG-ChvI regulatory system but also suppresses the plant’s defence response. Stable low pH affects the distribution of Ca^{2+} ions and suppresses the expression of the marker defence-related genes NDR1/HIN1-like 10 (NHL10) and FLG22-induced receptor-like kinase I (FRK-I), thereby enhancing the susceptibility of Agrobacterium-mediated gene transfer (Wang et al. 2018a). Although VirG transcription is activated by the ExoR-ChvG-ChvI cascade, VirG induction requires the phosphorylation of its Asp 52 residue by VirA sensor kinase after detecting the phenolic signal. Additionally, sugar from the plant exudates acts as a signalling molecule for virulence activation and is detected by other chromosome-based virulence proteins, such as ChvE (Cangelosi et al. 1990). Interestingly,
low pH increases the affinity of ChvE to sugar (Hu et al. 2013). Additionally, low pH promotes the interaction of the ChvE protein with VirA at its periplasmic domain, which results in VirA protein activation (Gao and Lynn 2005; Nair et al. 2011). The host range of A. tumefaciens is reportedly determined by the interaction between ChvE and VirA (He et al. 2009; Hu et al. 2013). VirA/VirG is a plasmid-based regulatory system in Agrobacterium that is essential for its virulence (Lin et al. 2014; Wise and Binns 2016). In the two-component regulatory system VirA/VirG, VirA acts as sensor histidine kinase, whereas VirG acts as a response regulator. When AS is detected by VirA, it gets autophosphorylated and, in turn, phosphorylates the Asp 52 residue of VirG response regulator, which is located in the cytoplasm of Agrobacterium. In the Ti plasmid, phosphorylated VirG binds to the 12 bp Vir box region, which is located upstream of the transcriptional start site of the virulence gene (Subramoni et al. 2014). As a result, the vir genes, which are distributed in 11 operons, are activated. The vir genes include virA, virB, virC, virD, virE, virF, virG, virH, virK, virL and virM. The virB operon encodes most proteins, 11 in total, whereas the virA and virG operons always exhibit low levels of expression (Nester 2015) so that they can sense the extracellular stimuli. The model for signal perception is depicted in Fig. 1. The single-stranded T-strand is generated from the Ti plasmid via nicking at the right and left borders by VirD2 and VirD1. Additionally, the generation of multiple copies of T-strand and its conjugative transfer are maintained by VirC1 and VirC2 proteins (Atmakuri et al. 2007). These four virulence effector proteins contribute to efficient T-strand generation, and VirD2 remains associated with the right border of T-strand. The T-strand-VirD2 nucleoprotein complex, along with several virulence effector proteins, such as VirE2, VirF, VirE3 and VirD5, is then translocated to the host cell via the type 4 secretion system (T4SS). T4SS consists of the cell envelope spanning transporter and the extracellular pilus and is synthesised from VirD4 and VirB1-B11 proteins (Li and Christie 2018). Agrobacterium and plant cells are physically attached to each other with the aid of the VirB2 protein of Agrobacterium and the cell membrane proteins of the plant. To detect the membrane protein involved in attachment to the bacterium, Hwang used yeast two-hybrid assays to screen out the interacting partner of VirB2 and observed that the Arabidopsis reticulon-like (RTNL) proteins, AtRTNLB1, AtRTNLB2 and AtRTNLB4, interacted with the VirB2 protein (Hwang and Gelvin 2004). VirD4 is a part of the transmembrane domain and may use ATP to transfer the T-strand to the pilus. VirD4 contains a C-terminal glutamine-rich conserved region that enables the recognition of the ‘VirD2-T-strand’ complex as a substrate for translocation in plants (Das 2020). VirE2 enters the host cell via clathrin-mediated endocytosis (Li

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**Fig. 1** Model for signal perception in Agrobacterium. Three factors, low pH, sugar and phenolic compound (Acetosyringone:AS) are sensed by ChvG/ChvI, ChvE and VirA/VirG regulatory systems, respectively. Under neutral pH, ExoR remain bounded to ChvG and make them inactive which get activated under low pH by proteolytic cleavage and dissociation of ExoR. This allow the ChvG for autophosphorylation and transfer phosphate group to ChvI response regulator which then activate VirG. Sugar and low pH induces the ChvE for binding and activation of VirA. Simultaneously, AS perceived by cytoplasmic domain of VirA which allow the autophosphorylation of VirA. VirA phosphorylates VirG and induce it for activation of Vir operon for activation of other virulence proteins.
and Pan 2017); however, the rest of the mechanism is yet to be elucidated.

**Generation of the T-complex and its cytoplasmic trafficking and nuclear import**

VirE2 is a single-stranded DNA binding (SSB) protein. VirE2 binds to the T-strand and forms a right-handed, cord-like structure inside the host cell, thus forming the T-complex (Abu-Arish et al. 2004). VirE2 has also been proposed to protect the T-strand from host nucleolytic degradation (Citovsky et al. 1989). Both VirE2 and VirD2 contain nuclear localization signals (NLSs), which import the T-strand into the nuclei. The VirD2 and VirE2 proteins use the host importins and VIP1 and VIP2 to create a ‘super T-complex’ that allows T-DNA nuclear import (Gelvin 2010, 2012; Guo et al. 2009, 2019; Shi et al. 2014). Prior to the nuclear import, the T-strand must, however, be trafficked in the host cytoplasm, which is largely accomplished by the VirE2 protein. To enter the nucleus, VirE2 gets associated with the F-actin network and the endoplasmic reticulum. Furthermore, VirE2 utilises the host myosin XI-K system for trafficking (Tu et al. 2018; Yang et al. 2017).

**Integration of the T-DNA into the plant genome and its expression**

When the T-DNA enters the host nucleus, it must first be stripped of its associated Vir and host proteins so that it can be integrated into the host genome. *Agrobacterium* VirF is similar to the F-box protein of the host, which is utilised to strip off the virulence proteins. VirF employs the host ubiquitin/proteosomal activity to degrade these proteins. VirF, however, is prone to proteosomal degradation in the host cell, and another virulence protein, VirD5, protects VirF inside the cell (Wang et al. 2014, 2018b). Interestingly, the *Agrobacterium* VirF mutant strain did not alter the transformation susceptibility in *Arabidopsis* or *Nicotiana*, which suggests that some other factor is involved in this step too. The host F-box protein has been identified and named as VIP1 binding F-box protein (VBF4). This protein targets VIP1 and VirE2 for proteosomal degradation via the S-PHASE KINASE-ASSOCIATED PROTEIN1 (SKP1)-CULLIN1 (CUL1)-F-box protein (SCFVBF) pathway (Zaltsman et al. 2010, 2013). It can, therefore, be said that although *Agrobacterium* utilises the host’s cellular machinery, it also possesses a backup strategy that involves the virulence protein.

It has been suggested that transcriptionally active regions of the genome that contain a high proportion of A=T sequences are the most likely sites for the integration of T-DNAs (Bourras et al. 2015). Later, the sequence of T-DNA/plant junction was analysed in plants without antibiotic selection, and the pattern of T-DNA integration was found to be random (Shilo et al. 2017). These junction sites are typically the double-stranded DNA repair sites in the plant genome (Kleinboelting et al. 2015; Gelvin 2017). Although the T-DNA integration process resembles a DNA repair process, such as non-homologous end joining (NHEJ) and microhomology-mediated end joining (MMEJ), the integration does not entail the same proteins involved in these repair processes (van Attikum et al. 2003; Park et al. 2015).

The T-strand enters the nucleus as a single-stranded DNA, but whether the T-DNA integrates into the host genome as a single or double-stranded form is still under investigation. However, the double-stranded T-DNA integration is comparatively more favourable (Kleinboelting et al. 2015). In 2016, it was shown that the T-DNA is integrated as a double-stranded break at the microhomologous site in the genome via the annealing and repair process. In a study by van Kregten et al., it was proven that the mutation of DNA polymerase θ gene in *Arabidopsis* inhibits its stable transformation but not its transient transformation via *Agrobacterium*. The group suggested that DNA polymerase θ initiates the first step in T-DNA integration (Van Kregten et al. 2016). In contrast, Nishizawa-Yokoi et al. proposed that T-DNA integration occurs via multiple redundant pathways and that it might involve some other unknown pathway (Nishizawa-Yokoi et al. 2021).

A T-DNA sequence with eukaryotic promoter elements, such as the TATA box, CAAT box and polyadenylation signal, is expressed in eukaryotic cells (Zhang et al. 2015). This feature indicates the eukaryotic origin of the T-DNA fragment. T-DNA also exhibits microhomologies with eukaryotic promoter core elements at its 3′ end, which signifies its likelihood of integration at promoter sites (Bourras et al. 2012). The model for *Agrobacterium*-mediated T-DNA transfer is illustrated in Fig. 2A.

**Defence response of the plant during plant–*Agrobacterium* interaction**

Similar to the innate immune system of the animals, plants also have their defence system to fight against pathogen attack. When a wound occurs, plants secrete metabolites, such as H ions, phenolics and carbohydrates, to heal the cell damage at the wound site (Lacroix and Citovsky 2013). During wound formation, plant exudates are the first level of check to prevent *Agrobacterium* infection. For instance, unlike the crown gall-susceptible dicot plants, maize seedlings secrete chemicals that block the growth of *Agrobacterium*. Chemicals such as DIMBOA (2,4-dihydroxy-7-methoxy-2H-1,4-benzoxazin-3(4H)-one) and MDIBOA (2-hydroxy-4,7-dimethoxybenzoazin-3-one) inhibit the
growth of *Agrobacterium* (Zhang et al. 2000). Hence, the non-natural host of *Agrobacterium* is incapable of inducing virulence, thereby protecting it from crown gall disease. This review discusses the defence response of plants against *Agrobacterium*, which starts with the perception of pathogen signals, and then moves on to disease development.

**Pathogen signal perception and innate plant immunity**

Plants respond to elicitors, such as flagellin, elongation factor-thermo unstable (EF-Tu) and lipopolysaccharide, which trigger innate immune response during pathogen attack (Janda et al. 2019). The plant PRRs comprise two types of membrane receptors, namely, receptor-like kinases (RLKs) and receptor-like proteins (RLPs) (Couto and Zipfel 2016; Boutrot and Zipfel 2017). Both RLKs and RLPs possess extracellular and transmembrane domains; however, while RLKs contain an intracellular kinase domain, RLPs do not (Saijo et al. 2018). An *Arabidopsis thaliana* PRR that has been well characterised so far is flagellin sensing 2 (FLS2), which possesses leucine-rich repeats (LRRs) as a ligand-binding motif and belongs to the RLK family (Chinchilla et al. 2007). FLS2 recognises the 22-amino-acid long peptide of flagellin (monomer of flagella), flg22. The perception of flg22 via FLS2 stimulates plant immune response by modulating protein activity and increasing reactive oxygen species (ROS) accumulation and phytohormone synthesis, such as salicylic acid (SA) and jasmonic acid (JA) (Bigeard et al. 2015). This step is followed by the activation of defence-related genes as well as callose deposition at the cell wall to strengthen the wall composition and prevent pathogen ingress into the plant (Muthamilarasan and Prasad 2013; Janda et al. 2019).

Some bacterial pathogens escape FLS2 immunodetection by modifying the flg22 epitope. In *A. tumefaciens*, for instance, flg22 is modified into flg22Atum, which is
undetectable by the FLS2 immunoreceptor (Felix et al. 1999; Trdá et al. 2014). flg22Atum differs from flg22 in that half of the 22 amino acid residues are modified. As a part of the plant–pathogen co-evolution effort, the flg22Atum receptor from the cell culture of\textit{Vitis riparia} was identified and named FLS2XL (FLS2 with eXtended Ligand recognition). FLS2XL had an additional 16-amino-acid extension, which might differ from FLS2 in the ligand recognition process (Fürst et al. 2020). In comparison with FLS2 (VrFLS2), FLS2XL can bind flg22Atum with a higher affinity than flg22. FLS2XL ligand accommodation in its extracellular domain differs from that of FLS2 in that the former accommodates the ligand with lesser steric hindrance. To understand the perception at the domain level, chimeric receptors with the extracellular LRR domain of FLS2XL and the cytoplasmic domain of VrFLS2 were generated, thus resulting in a chimeric FLS2XL (c-FLS2XL). The c-FLS2XL showed that 12–18 LRRs of FLS2XL were crucial for flg22Atum immunodetection; nevertheless, the other LRRs also provided sensitivity for flg22Atum epitope detection. \textit{Nicotiana} plant expressing c-FLS2XL was found to be resistant to crown gall disease (Fürst et al. 2020). In addition to FLS2, the EF-Tu receptor (EFR) has been identified in \textit{Arabidopsis} for its potential role in \textit{Agrobacterium} infection. EFR belongs to the LRR-RLK type of PRR because FLS2 and EF-Tu from \textit{Agrobacterium} work as eliciting ligands. EF-Tu is a highly conserved and abundant bacterial protein that plays a role in the protein translation process. EFR recognises the N-terminal 18 amino acid residues in EF-Tu, named elf18, which serves as a ligand epitope. In addition, EFR activates the downstream defence response, which is not identical but similar to FLS2 (Zipfel et al. 2006; Wan et al. 2019). Unlike FLS2 which is found in almost all higher plants, EFR is only found in the Brassicaceae family of plants. However, the transgenic expression of EFR in other groups of plants, such as rice (Lu et al. 2015; Schwessinger et al. 2015), wheat (Hj et al. 2015) tomato, \textit{Nicotiana} and Medicago (Lacombe et al. 2010; Pfeilmeier et al. 2019), provides tolerance against pathogenic bacteria. By generating efr mutant \textit{Arabidopsis} plants, it was confirmed that this pathway does not perceive elf18 and enables the infection of the plant by \textit{Agrobacterium} (Zipfel et al. 2006). The manner in which EF-Tu gets exposed to the outer membrane of \textit{Agrobacterium} is still under research, but it has been shown that EF-Tu is secreted to the outer membranes by other pathogen, such as \textit{Xanthomonas campestris} and \textit{Erwinia chrysanthemi} (Watt et al. 2005; Kazemi-Pour et al. 2004). The EF-Tu protein from \textit{Acinetobacter baumannii} is associated with outer membrane vesicle (OMV), cell surface and fibronectin (Dallo et al. 2012). Based on these findings, it appears that EF-Tu is transported to the outer membrane by vesicle trafficking, but the precise mechanism is yet to be elucidated. It is noteworthy that the downstream signalling of both EFR and FLS2 generates the same kind of defence response during \textit{Agrobacterium} infection in \textit{Arabidopsis}. Also, the combination of EF-Tu and flg22 fails to induce synergistic defence responses but exerts the same impact on MAP-kinase transduction cascade (Zipfel et al. 2006; Dafny-Yelin et al. 2008). Because the cytoplasmic fragment of FLS2XL is similar to FLS2, it can be speculated that FLS2XL is likely to exhibit a signalling akin to that of FLS2 during \textit{Agrobacterium} infection. Additionally, RTNLB4, a membrane associated protein that interacts with VirB2 in the T-pilus plays a possible role in defence during \textit{Agrobacterium} infection (Hwang and Gelvin 2004). Upon infection with the elf18 peptide of \textit{Agrobacterium}, plants with abnormal levels of RTNLB4, either overexpression or mutant lines, showed hampered immunity. Furthermore, the rtnlb4 mutant plants were more resistant to \textit{Agrobacterium} infection than the wild-type plant, suggesting its probable role in addition to defence. \textit{Agrobacterium} VirE2 utilises cytoplasmic trafficking to enter the nucleus. RTNLB4 is found on the plasma membrane and endoplasmic reticulum and is involved in intracellular trafficking. Therefore, RTNLB4 is supposedly involved in the intracellular transmission of T-DNA (Huang and Hwang 2020). Furthermore, RAB8A, 8B and 8D interact with several RTNLB proteins and participate in the \textit{Agrobacterium} infection process (Huang et al. 2021).

Because the VirB2 protein interacts with the RTNLB4 membrane protein, it was tested whether the VirB2 peptide acted as PAMP, similar to elf18 and flg22. Two peptide regions of VirB2, S111-T58 and I63-I80, were found to alleviate plant defence response, and the residues of the VirB2 peptide from Q-48 to V-101 might be involved in plant–\textit{Agrobacterium} interaction. In addition, both, elf18 and VirB2 have been shown to activate the early defence-related genes, namely, MPK3, MPK6, WRKY22, WRKY29, FRK1 and PR1, during \textit{Agrobacterium} infection (Huang and Hwang 2020).

### MAPK signalling

As a response to PAMPs, PTI activates the MAPK signalling pathway in several ways, including protein phosphorylation, ROS burst and transcriptional reprogramming of defence genes (Boutrot and Zipfel 2017). A previously held theory suggested that PTI induced an ROS burst that acted upstream of the MAPK signalling cascade, but later it became clear that PTI signalling was split into two distinct pathways, one triggering MAPK signalling and the other triggering ROS burst. The β subunit of G-protein (AGB1) associates with NADPH-oxidase and contributes to ROS generation. The
EFR-mediated PAMP perception of *Agrobacterium* during infection is associated with the AGB1-mediated ROS burst. The EFR-AGB1 downstream signalling hampers *Agrobacterium* infection in *Arabidopsis* (Xu et al. 2014; Ishikawa 2009). Somatic embryogenesis receptor kinase (SERK)3/brassinosteroid (BR)-associated kinase (BAK1) is a key coreceptor that recognises PAMP via FLS2. In plants, BAK1 activates the MAPK signal transduction cascade. BAK1 has previously been reported to be activated during brassinosteroid hormone regulation, which, in turn, activates the brassinosteroid-insensitive 1 (BRI1) receptor. During innate immunity too, BAK1 interacts with BRI1 and then activates the MAPK signalling cascade (Heese et al. 2007; Bigeard et al. 2015). This cascade is composed of three signalling modules, namely, MAPKKks, MAPKKks and MAPks. *Arabidopsis* encodes 60 MAPKks, 20 MAPKks and 10 MAPks, which suggests that a single MAPK can activate multiple MAPKks, which, in turn, can activate multiple MAPKKks. In *Arabidopsis*, four MAPKs, namely, MPK3, MPK6, MPK4 and MPK11, have been found to be responsive to pathogen infection (Bigeard et al. 2015).

**Defence responsive genes**

A PTI response activates the MAP-kinase signal transduction cascade, which further activates the transcription factors such as WRKY33 and VIP1, thus leading to the activation of several defence-related proteins. An in-depth analysis of the interaction between *Ageratum* and *Agrobacterium* has been performed to determine how plants respond to *Agrobacterium* infection (Ditt et al. 2001). Ditt et al. showed that plants expressed defence responsive genes after 24–48 h of *Agrobacterium* infection, which mostly include PR protein, NtPrP27 from tobacco, defence responsive proteins of phenylpropanoid pathway, cytochrome P450 monooxygenase from *Arabidopsis* and disease-resistant protein, Xa21, from rice (Ditt et al. 2001, 2006). Also, Veena et al. demonstrated that the defence responsive genes are differentially expressed during the early hours (3–6 h) of *Agrobacterium* infection in plants, which include glutathione-S-transferase, PR genes and SARs (Veena et al. 2003).

In addition to triggering the innate immune response of plants, *Agrobacterium* hijacks the host machinery during infection. *Arabidopsis* VirE2 interacting protein 1 (VIP1), a bZIP transcription factor, is activated by phosphorylation via MAPK (MPK3), which further activates the pathogenesis-related 1 (PR1) promoter containing defence genes (Djamei et al. 2007). VIP1 binds to the VIP1 responsive element (VRE; ACN\(\text{GCT}\)) of the PR1 promoter (Pitzschke et al. 2009). VIP1 is localised in the cytoplasm under non-stress conditions; however, upon infection with *Agrobacterium*, it gets phosphorylated at the Ser79 residue (via MPK3) and relocates to the nucleus to activate the defence genes (Djamei et al. 2007). VirE2, an *Agrobacterium* effector protein, utilises the nuclear import activity of VIP1 to transport T-DNA into the host nucleus (Tzfira et al. 2001). In another study, it was found that VirE2 binds to VIP1 and decreases its level inside the cell, which, in turn, lowers the level of defence response during *Agrobacterium* infection. (Shi et al. 2014). According to Lapham et al., VirE2 localised in the cytoplasm modulates plant RNAs and genes to promote transformation (Lapham et al. 2021).

Plant transcription factor VFP4 (VirF binding protein) and its downstream gene ATL31 have both been found to be activated in response to *Agrobacterium* infection, and the overexpression of the gene renders the plant resistant to infection. VFP4 differentially regulates the defence responsive genes, including antibacterial genes; thus, VFP4 provides another defence layer against *Agrobacterium*. The bacteria detects VFP4 via VirF effector protein, which is then processed for proteasomal degradation via the SCFVirF pathway (García-Cano et al. 2018). Besides *Arabidopsis* VFP1, HvVFP1 from barley has also been identified to be activated during *Agrobacterium* infection. The HvVFP1 protein contains a conserved bZIP domain and exhibits a positive correlation with barley’s PR genes, such as HvPR1, HvPR4 and HvPR10. Apart from PR1 activation, HvVFP1 also activates one of the MAP-kinase members, HvMPK1 (El Sarraf et al. 2019). It is possible that HvVFP1 confers *Agrobacterium* resistance in barley. A VIP1 from *Populus trichocarpa*, PtVIP1, also serves against pathogen invasion by activating the PR1 gene (Wang et al. 2019). Recently, a PR 10 gene from the *Agrobacterium*-recalcitrant plant *Hypericum perforatum* has been identified and named as phenolic oxidative coupling protein (Hyp-1). This defence gene has been found to hinder *Agrobacterium* infection in Tobacco (Hou et al. 2020). During infection, Hyp-1 transcripts get upregulated and play an important role in plant defence (Karpinnen et al. 2016). It has been suggested that Hyp-1 confers tolerance to *Agrobacterium* by downregulating auxin signalling pathway genes, such as *NiaTIR1* and *NiaAFR8*. During pathogen attack, Hyp-1 induces the expression of MiR160, which targets the *ARF10, ARF16* and *ARF17* transcripts of the auxin signalling cascade and downregulates their expression (Hou et al. 2020; Pinweha et al. 2015; Wójcik et al. 2017). Dunoyer found low levels of small RNAs associated with the iaaM (tryptophan 2 oxygenase) and ags (agropine synthase) genes of T-DNA in *Nicotiana tabacum* after 3 days of *Agrobacterium* infiltration. Additionally, RNAi-deficient plants were susceptible to *Agrobacterium*, suggesting that RNA silencing may provide a defence against *Agrobacterium*.

However, small interfering RNA (siRNA) has been detected during the initial days of infection in the tissues; later, the anti-silencing state is maintained by *Agrobacterium*, which inhibits the synthesis of siRNA. Thus,
again, *Agrobacterium* takes over the plant defence system and leads to successful infection (Dunoyer et al. 2006). Recently, we have identified a tau class GST in rice, GSTU5, which interacts with the VirE2 protein of *Agrobacterium* and hinders its single-stranded DNA binding (SSB) property. The GSTU5-knockdown lines in rice were more susceptible to *Agrobacterium* infection in comparison with its overexpressing lines, thus alluding the probable role of GSTU5 as a defence gene in rice (Tiwari et al. 2022). Therefore, it can be said that plants have a three-layered immunity against *Agrobacterium*. In the first layer, PRRs, such as FLS2XL, FLS2, EFR and RTNLB4, play a role (further confirmation is required for RTNLB4). The second layer of plant immunity is related to signal integration, which involves MAP-kinase defence-related genes, especially MPK3, MPK4, MPK6 and MPK11. The last layer of immune response involves defence induction and amplification of defence-related transcription factors, such as VIP1, WRKY22 and WRKY29, which activate the defence-related genes. Thus, during plant–*Agrobacterium* interaction, disease susceptibility and resistance are dependent upon the layered immunity of plants and the virulence effector proteins of *Agrobacterium*. A model for PAMP perception and plant immune response during *Agrobacterium* infection is given in Fig. 2B.

A list of genes that act during *Agrobacterium* infection in different plants is presented in Table 1.

### Table 1 A list of plant genes act during *Agrobacterium* infection

| Plant genes | Specific role | Plant | Reference |
|-------------|---------------|-------|-----------|
| Flagellin sensing 2 (FLS2) | Perceives flg22 of *Agrobacterium* | *Arabidopsis thaliana* | Chinchilla et al. (2007) |
| FLS2 with eXtended Ligand recognition (FLS2XL) | Perceives flg22Atum of *Agrobacterium* | *Vitis riparia* | Fürst et al. (2020) |
| Elongation factor-Thermo unstable (EF-Tu) receptor (EFR) | Perceives 18 aa residues in EF-Tu named elf18 of *Agrobacterium* | *Arabidopsis thaliana* | Zipfel et al. (2006) |
| RETICULON-LIKE4 (RTNLB4) | Perceives elf18 peptide of *Agrobacterium* | *Arabidopsis thaliana* | Hwang and Gelvin (2004) |
| NtPRp27 | Defence gene | *Nicotiana tabacum* | Ditt et al. (2001); Ditt et al. (2006) |
| Cytochrome P450 monooxygenase | Defence gene | *Arabidopsis thaliana* | Ditt et al. (2001); Ditt et al. (2006) |
| Xa21 | Defence gene | *Oryza sativa* | Djamei et al. (2007) |
| VirE2 interacting protein (VIP1) | Activates defence genes | *Hordeum vulgare* | El Sarraf et al. (2019) |
| HvVIP1 | Activates *PR1* gene (HvPR1, HvPR4, and HvPR10) | | |
| PtVIP1 | Activates the *PR1* gene | *Populus trichocarpa* | Wang et al. (2019) |
| Phenolic oxidative coupling protein (Hyp-1) | Defence gene | *Nicotiana tabacum* | Hou et al. (2020); Hou et al. (2020); Pinweha et al. (2015); Wójcik et al. (2017) |
| OsGSTU5 | Defence gene | *Oryza sativa* | Tiwari et al. (2022) |
| MPK3, MPK6 | MAP-kinase defence genes | *Arabidopsis thaliana* | Huang and Hwang (2020) |
| WRKY22, WRKY29 | Defence genes | *Arabidopsis thaliana* | Huang and Hwang (2020) |
| FRK1, and PR1 | Defence genes | *Arabidopsis thaliana* | Huang and Hwang (2020) |

### Concluding remark

The remarkable abilities of *Agrobacterium* never cease to attract plant biologists. The bacterium captures immense attention because of its pathogenesis as well as its biotechnological significance. The unique action of *Agrobacterium* is determined by virulence and host proteins. Research on plant–*Agrobacterium* interaction provides an insight into the molecular communication between these two organisms. The more we learn about host–pathogen interactions, the more interested we become in the next step in their path. There is still much to be explored in terms of plant–*Agrobacterium* interaction. For instance, VirE2 was recently shown to enter the plant cells via clathrin-mediated endocytosis, however, it remains unclear how T-DNA and other virulence effector proteins enter via the cell membrane. Furthermore, the T-complex enters the nucleus via VirE2-mediated cytoplasmic trafficking, but it is unclear how other virulence proteins, such as VirF and VirD5, enter the plant nucleus. The exact mechanism of T-DNA integration is still elusive.

### Future prospects

As we gain an understanding of layered immunity during plant–*Agrobacterium* interaction, we can utilise this information in plant breeding to control crown gall. By elevating...
and stacking the extracellular immunity receptors, it might be possible to broaden the disease resistance. In addition, modulating the downstream signal transduction pathway to promote the expression of defence-related genes would enhance disease resistance. Recent genetic editing techniques, such as CRISPR/Cas9, allow the targeted modification of extracellular immune receptors and might be useful in exploring the downstream pathway in plants during Agrobacterium infection.

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Declarations

Conflict of interest The authors declare no conflict of interest.

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