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Phytopathogen type III effector weaponry and their plant targets

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Summary

Phytopathogenic bacteria suppress plant innate immunity and promote pathogenesis by injecting proteins called type III effectors into plant cells using a type III protein secretion system. These type III effectors use at least three major strategies to alter host responses. One strategy is to alter host protein turnover, either by direct cleavage or by modulating ubiquitination and targeting to the 26S proteasome. Another strategy involves alteration of RNA metabolism by transcriptional activation or ADP-ribosylation of RNA-binding proteins. A third major strategy is to inhibit the kinases involved in plant defence signalling, either by removing phosphates or by direct inhibition. The wide array of strategies bacterial pathogens employ to suppress innate immunity suggest that circumvention of innate immunity is critical for bacterial pathogenicity of plants.

Introduction

Gram-negative bacterial plant pathogens such as *Pseudomonas syringae, Ralstonia solanacearum* and *Xanthomonas campestris* infect their hosts by entering through wounds or natural openings. Once inside they produce virulence factors that alter the host providing an environment beneficial for the bacteria and suppressing host defences. Among the most important of these virulence factors are proteins called type III effectors (T3Es) that are injected directly into host cells using a type III protein secretion system (T3SS) known as the hypersensitive reaction and pathogenicity (Hrp) T3SS system [1]. Bacteria that are compromised in the Hrp T3SS are extremely reduced in virulence, demonstrating that T3Es are critical for successful pathogenesis [2]. The virulence activity of several T3Es is thought to be due to their ability to suppress plant innate immunity.

Plant innate immunity controls the activation of innate immune responses using mitogenactivated protein kinase (MAPK) cascades and hormones such as salicylic acid (SA). These defences include the deposition of lignin and callose in the cell wall, transcription of pathogenesis related (PR) genes, and the production of antimicrobial compounds and reactive oxygen species [3]. Plant innate immunity can be divided into two branches based on the types of microbial molecules recognized.

One branch recognizes conserved molecules essential to many microbes called pathogenassociated molecular patterns (PAMPs) or microbe-associated molecular patterns (MAMPs). PAMP recognition occurs by pattern recognition receptors (PRRs) predominately localized in the extracellular matrix. In plants these PRRs are PAMP receptor-like kinases and this

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recognition, which historically has been known as basal defense, is now referred to as PAMP-triggered immunity (PTI) [4,5,6].

The other branch recognizes specific pathogen effectors (historically referred to as avirulence (Avr) proteins) using nucleotide binding site-leucine rich repeat (NB-LRR) resistance (R) proteins. This recognition response is associated with the long-standing gene-for-gene hypothesis and is now known as effector-triggered immunity (ETI) [4]. For bacterial pathogens the only proteins known to induce ETI are T3Es. ETI has overlapping signalling pathways with PTI but generally results in a faster and more intense defence response, one outcome of which is a rapid and localized programmed cell death known as the hypersensitive response (HR).

ETI, in some cases, may be caused by the direct recognition of an effector by an R protein but a current hypothesis, known as the guard hypothesis [7], is that many R proteins detect the modifications of host targets made by specific pathogen effectors. Thus, R proteins detect effector activities rather than the proteins themselves and act to guard or monitor important host proteins for signs of pathogen ingress. Bacteria must suppress or evade both PTI and ETI responses in order to be successful pathogens and T3Es appear to play a major role in this suppression. Thus, the combination of host PRRs and R proteins, and the pathogen's T3Es largely decides whether the pathogen or the plant will be successful in this molecular tête-à-tête. An important area of research in plant-pathogen interactions is the determination of how these T3Es function. These studies provide insight into how pathogen virulence and plant innate immunity work, as well as uncovering new enzyme activities.

Widespread screens to determine the effect of T3Es on plant immunity outputs demonstrate that certain T3Es can suppress the plant innate immunity [8,9,10] or up-regulate hormones that are antagonistic to SA and other defence responses [11,12,13]. This review will focus on recent progress in elucidating T3Es activities, plant targets and mechanisms of action. The impact of this new information on the guard hypothesis will also be addressed. The review is divided into sections based on the following T3E activities: protein turnover; RNA expression or stability; protein phosphorylation and dephosphorylation; and clues about T3E activities based on interactions with host proteins. Their sites of action and enzymatic activities are summarized in Fig. 1 and Table 1.

T3Es that impact host protein turnover

One way for T3Es to suppress plant innate immunity is to remove or inactivate its components. Certain T3Es are proteases that can remove these components by degradation. For example, AvrPphB from *P. syringae* is a papain-like cysteine protease that cleaves the *Arabidopsis* protein kinase PBS1 [14]. PBS1 exists in a complex with the R protein RPS5, which recognizes its cleavage and initiates ETI [15]. This recognition makes AvrPphB an obvious liability for *P. syringae* on RPS5-containing plants. These results support the guard hypothesis since they suggest that RPS5 is guarding PBS1 and perceives its modification by AvrPphB. *Arabidopsis pbs1* mutants are not more susceptible to *P. syringae*. Thus, it is unclear whether *P. syringae* benefits from cleaving PBS1 in plants lacking RPS5. Therefore, PBS1 may be either a redundant virulence target or a plant protein that resembles a virulence target (i.e., a decoy) used by the plant to detect AvrPphB activity.

Another *P. syringae* T3E, AvrRpt2, is a staphopain-like cysteine protease that cleaves RIN4 [16], a negative regulator of PTI [17]. It seems counter intuitive that RIN4 is a virulence target since its loss de-represses PTI, which would apparently have a negative impact on pathogenic success. However, RIN4 interacts with at least two other T3Es and two R proteins suggesting that inactivating it may destabilize this complex and benefit the pathogen. One of these R proteins, RPS2, recognizes the cleavage of RIN4 by AvrRpt2 and induces ETI [18,19]. This finding was one of the first to provide molecular evidence in support of the guard hypothesis.

Two families of T3Es (XopD and YopJ) are cysteine proteases/isopeptidases that specialize in the cleavage of ubiquitin or SUMO (small ubiquitin-like modifiers) from proteins. As the addition of a single ubiquitin or SUMO to proteins can alter their activity, it is likely that the de-ubiquitinating/de-SUMOylating activity of these T3Es alters the activity or stability of host proteins. XopD is a *X. campestris* pv. *vesicatoria* (*Xcv*) T3E that is translocated to the nucleus where it is believed to mimic endogenous plant SUMO isopeptidases [22] as its activity is specific to plant SUMOs [23].

AvrRpt2 was recently shown to be capable of altering auxin physiology [12], whether this is

a direct or indirect effect remains to be determined.

YopJ is a well-studied human pathogen T3E from *Yersinia* spp. whose family members in *Xcv* include AvrXv4, AvrBsT, AvrRxv and XopJ. These T3Es appear to be SUMO isopeptidases as the expression of AvrXv4 *in planta* leads to a reduced level of SUMOylated proteins [24]. AvrXv4 is cytoplasmic [24], whereas XopJ is targeted to the plasma membrane [25] suggesting they have different targets *in vivo*. *Yersinia* YopJ has de-ubiquitinating, de-SUMOylating and acetylating activities and targets MAPK signalling pathways [26,27], therefore these YopJ family members may also possess acetyltransferase activity and their plant targets may also be involved in MAPK signalling.

Rather than possessing protease activity some T3Es use the host ubiquitination machinery and 26S proteasome to alter protein levels. For instance, *Ralstonia solanacearum* GALA T3Es have an F-Box domain and a leucine-rich repeat domain and interact with *Arabidopsis* ASK proteins [28•]. F-box and ASK proteins are part of ubiquitin E3 ligase complexes that attach ubiquitin to specific proteins and polyubiquitinated proteins are subsequently degraded by the 26S proteasome. Therefore, GALAs probably alter host protein regulation or turnover. They also increase the virulence of *R. solanacearum*, as a mutant lacking all 7 GALAs has severely reduced growth on *Arabidopsis* [28•]

A T3E from *P. syringae*, AvrPtoB (also known as HopAB2), contains a C-terminal E3 ligase domain [29] that ubiquitinates the tomato protein kinase Fen (originally identified as a cause of sensitivity to the pesticide fenthion [30]) leading to the degradation of Fen by the 26S proteasome [31••]. However, ETI occurs when the R protein Prf recognizes the interaction of Fen with AvrPtoB truncations that lack the E3 ligase domain. Thus, AvrPtoB has apparently evolved to evade ETI by acquiring an E3 ligase domain that marks Fen for degradation [31••]. Interestingly, some plants seem to have countered AvrPtoB's E3 ligase activity by evolving the Pto kinase, which cannot be ubiquinated by AvrPtoB and induces Prf-mediated ETI. AvrPtoB appears to have multiple plant targets since it suppresses PTI-induced MAPK pathways in *Arabidopsis* [32]. It will be important to determine whether Fen and Pto are virulence targets or virulence decoys – if they are decoys evolved to 'trick' T3Es it would be tempting to speculate that, in a manner similar to AvrPtoB.

A final example of a T3E that alters protein turnover is HopM1 (formerly HopPtoM) from *P. syringae* [33]. It is present in the endomembrane fraction of plant cells where it enhances AtMIN7 degradation by targeting it to the 26S proteasome [34••]. AtMIN7 is an adenosine diphosphate ribosylation factor guanine nucleotide exchange factor (ARF-GEF). HopM1 may target AtMIN7 for degradation in order to disrupt vesicle transport as ARF-GEFs have been implicated in vesicle trafficking. Vesicle transport is necessary for the export of defence compounds to the cell wall and apoplast, so its disruption would impair cell wall-based defences

[34••]. HopM1 has no obvious similarity to known components of the host ubiquitination machinery, so exactly how it targets AtMIN7 to the proteasome remains to be established.

Targeting host transcription or RNA stability

T3Es can also modify protein levels by altering host transcription, which can lead to increased host susceptibility. T3Es that target host transcription include AvrBs3, PthXo6, and PthXo7 from *Xanthomonas* spp., and HsvG and HsvB from *Pantoea agglomerans*. AvrBs3 alters transcription by binding to the promoter region of the pepper transcription factor *upa20* and increasing its expression. This up-regulation of *upa20* is partly responsible for the hypertrophy of mesophyll cells seen when AvrBs3 is overexpressed in plants [35••]. In resistant plants AvrBs3 also binds to the promoter and activates the transcription of its cognate *R* gene, *Bs3*, leading to ETI [36••]. The encoded product of *Bs3* is a flavin-dependent monooxygenase rather than an NB-LRR protein and it causes ETI by an unknown mechanism [36••]. The case of AvrBs3 recognition could be considered as a variation of the guard hypothesis, in that instead of guarding a protein the *R* gene is guarding a DNA sequence. This could work by the promoter of *Bs3* acting as a decoy target for AvrBs3 by mimicking the promoter of *upa20*, which is apparently a 'real' virulence target.

AvrBs3 belongs to a family of transcription activator-like T3Es. Two members of this family, PthXo6 and PthXo7 increase the expression of the rice genes OsTFX1 (a bZIP transcription factor) and $OsTFIIA\gamma1$ (a small subunit of the transcription factor IIA), respectively [37•]. Loss of virulence in *PthXo6* mutants can be rescued by over-expression of OsTFX1 but it is not yet known if *PthXo6* directly binds to the promoter of OsTFX1 [36•]. Two other host genes *Xa27* [38] and Os8N3 [39] are specifically up-regulated by the T3Es AvrXa27 and PthXo1, respectively. Rice plants silenced for Os8N3 RNA result in plants that are less susceptible to *X. oryzae* pv. *oryzae* supporting the idea that Os8N3 is a susceptibility gene [39]. Although less established than with the *Xanthomonas* T3Es similar activities are emerging for the *P. agglomerans* T3Es HsvG and HsvB [40]. Collectively, these examples suggest that the use of T3Es to up-regulate expression of genes such as transcription factors that co-regulate many genes is a common virulence strategy employed by bacterial pathogens. These T3E-induced proteins likely represent susceptibility factors that make the plant more favourable to bacterial colonization. Whether they cause the suppression of PTI or modify the cellular environment to benefit the pathogen remains to be determined.

The *P. syringae* pv. *tomato* HopU1 (formerly HopPtoS2) T3E targets RNA metabolism rather than mRNA production. HopU1 is a mono-ADP-ribosyltransferase (ADP-RT) that ADP ribosylates several *Arabidopsis* RNA-binding proteins *in vitro*, including the glycine-rich RNA-binding protein GRP7 [41••]. Furthermore, this ADP-RT activity is necessary for HopU1's ability to suppress PTI and ETI and *Arabidopsis* plants lacking GRP7 are more susceptible to *P. syringae* [41••]. ADP-RTs are well-characterized toxins of animal pathogens but not previously shown to be associated with RNA-binding proteins. The ADP-ribosylation of RNA-binding proteins by HopU1 possibly decreases their ability to bind, stabilize or process RNA [41••]. It is not known how this alteration of RNA-binding leads to the suppression of innate immunity or if it alters specific or general RNA metabolism. Nevertheless, the post-transcriptional control of RNA by T3Es represents an exciting new pathogenic strategy to suppress innate immunity.

Phosphorylation or dephosphorylation of host proteins

PTI and ETI use kinase based signalling pathways, perhaps most prominently MAPK pathways. These kinases represent ideal T3E targets as multiple signal transduction pathways often converge at them and disabling them results in the suppression of many downstream

responses. As these pathways work by sequentially phosphorylating downstream components, modifying their phosphorylation states inhibits them preventing activation of host defences.

Two T3Es that seem to use this strategy are HopAI1 and HopAO1 (formerly HopPtoD2). HopAI1 from *P. syringae* has phosphothreonine lyase activity and irreversibly inactivates MAPKs [42••]. HopAI1 binds to animal MAPKs [42••] and MPK3 and MPK6 of *Arabidopsis* that are involved in defence responses to *P. syringae* [43]. HopAI1's broad specificity leaves open the possibility that its *in vivo* targets may include other MAPKs.

HopAO1 is a protein tyrosine phosphatase that contributes to virulence of *P. syringae* by suppressing PTI and ETI [44,45,46]. Because MAPKs are phosphorylated on tyrosine residues these are attractive putative targets for HopAO1. However, HopAO1 does not appear to act on MPK3 and MPK6 [46]. Therefore, its direct target(s) remain to be identified.

A slightly different strategy for modifying phosphorylation is used by the *P. syringae* T3E AvrPto, which interacts with the PAMP receptor kinases FLS2 and EFR, inhibiting their ability to autophosphorylate and activate MAPK signalling cascades [47••]. Similarly to AvrPtoB discussed above, AvrPto induces Pto-Prf-mediated ETI [48]. AvrPto inhibits the kinase activity of Pto [49•]. However, its induction of Prf-mediated ETI appears to be independent of this inhibition [49•]. Currently, the virulence target of AvrPto appears to be PAMP receptor kinases even though it also inhibits the kinase activity of Pto. This suggests that Pto may be an R protein-guarded decoy and that other T3Es that interact with it, such as AvrPtoB, may also target PAMP receptor kinases.

Clues about T3E activities based on interactions with host proteins

Host proteins that interact with T3Es have been identified through a variety of methods. Sometimes these interactors have provided clues to the T3E function. In other cases the proteinprotein interaction itself is vital to the activity of the T3E by either stabilizing or destabilizing host protein complexes. This class of T3Es includes AvrB, AvrRpm1 and HopI1 from *P. syringae*. In pioneering research, AvrB and AvrRpm1 was found to interact with RIN4 leading to its hyperphosphorylation, which apparently allows it to be recognised by the R protein RPM1 inducing ETI [50,51]. AvrB has also been found to interact *in vivo* with RAR1 [52]. Moreover, RAR1 was identified in a genetic screen for *Arabidopsis* mutants insensitive to AvrB [52]. RAR1 is an attractive candidate target for an AvrB target because, together with SGT1 and HSP90, it stabilizes R proteins and, therefore, is a primary component of ETI.

Protein-protein interactions also appear to be important for HopI1 (formerly HopPmaI) a choloroplast-targeted T3E with a potential J domain [53]. J domains are often found in the cochaperones of Hsp70, which direct proteins to Hsp70 and activate the ATPase activity and protein folding ability of Hsp70 [54]. The J domain of HopI1 is necessary for its virulence activity and overexpression of HopI1 in *Arabidopsis* causes thylakoid remodelling and the suppression of SA accumulation [55]. However, how the possible Hsp70 co-chaperone activity of HopI1 leads to these outcomes remains to be established. The targets and activities of the T3Es discussed here are summarized in Fig. 1 and Table 1.

Concluding remarks

It is clear that although much remains to be understood about how T3Es interfere with plant innate immunity several underlying themes are becoming apparent. The first is that the guard hypothesis probably accurately describes the perception of T3E activities inside plant cells. The open question now seems to be whether the guarded T3E targets identified such as Pto or RIN4 represent real virulence targets or decoys. If they indeed represent decoys this information could be exploited to identify their actual targets using similar logic employed for

AvrPto's targets (i.e., PAMP receptor kinases). In the fascinating co-evolution that has occurred between bacterial pathogens and plants it seems clear that PTI evolved prior to ETI. Did the evolution of PTI suppression provide the selection pressure necessary for ETI evolution? If so, it seems plausible that other R proteins may be guarding decoys of PAMP receptor kinase complexes.

A second emerging theme is that T3Es often have multiple targets many of which alter different aspects of innate immunity. These targets seem to function at every possible level of PTI and ETI pathways including the receptors themselves, downstream signal transduction pathways, and transcriptional and post-transcriptional responses. In this manner T3Es have apparently evolved to have the highest chance of suppressing sufficient innate immune responses to allow for the success of the bacterial pathogen. T3Es likely co-evolved to target a particularly important node of regulation or component of innate immunity. Because T3Es are targeting the innate immune response at seemingly every level, studying T3E targets promises to identify important unknown components of innate immunity and represent exciting tools for exploring plant biology.

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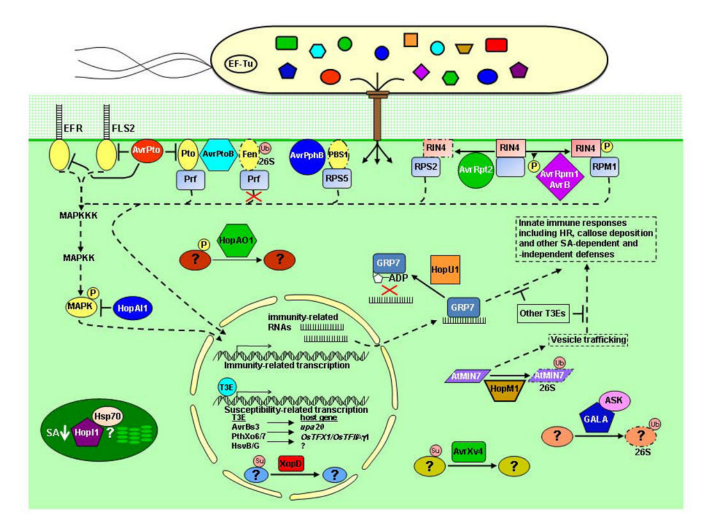


Fig. 1. Plant targets and activities of type III effectors from phytobacterial pathogens

Bacterial plant pathogens inject many different type III effectors (T3Es) into plant cells via the type III secretion system. The activities of T3Es can be recognized by plant resistance (R) proteins inducing effector-triggered immunity (ETI). The R protein RPM1 causes ETI by recognizing the phosphorylation (P) of RIN4 by the T3Es AvrRpm1 and AvrB, while RPS2 causes ETI upon the cleavage of RIN4 by the T3E AvrRpt2. The T3E AvrPphB degrades the PBS1 kinase inducing RPS5-dependent ETI. The R protein Prf recognizes the interaction of the Pto kinase with AvrPto or AvrPtoB eliciting ETI, however, AvrPtoB ubiquinates (Ub) the Fen kinase targeting it for degradation and preventing recognition by Prf. Plants can also use receptor kinases such as EFR or FLS2 to detect pathogen-associated molecular patterns (PAMPs). This leads to PAMP-triggered immunity (PTI), which like ETI, induces innate immune responses. AvrPto inhibits the kinase activity of Pto, FLS2, and EFR. The HopAI1 T3E is a phosphothreonine lyase that suppresses MAPKs. The HopAO1 T3E is a protein tyrosine phosphatase whose target is unknown. The HopU1 T3E is a mono-ADPribosyltransferase that modifies GRP7 glycine-rich RNA-binding protein and likely prevents it from binding RNA. The HopM1 T3E causes the ubiquination and degradation via the 26S proteasome of AtMIN7, which may be involved in vesicle trafficking. The GALA T3Es contain F-box domains and can interact with plant ASK proteins (part of an SCF-type E3 ubiquitin ligase complex). GALAs are predicted to change the ubiquitination status of host proteins. The T3Es XopD and AvrXv4, which function in different locations in the plant cell, are

isopeptidases that remove SUMO (Su) from host proteins. The chloroplast localized, J domaincontaining T3E HopI1, suppresses salicylic acid (SA) production and may associate with Hsp70. T3Es AvrBs3, PthXo6/7 and HsvG/B bind to specific promoters in the nucleus inducing the transcription of genes favoring pathogenesis. Broken lines indicate plant responses and solid lines T3E activities.

Table 1

T3E activities and plant targets

T3E	Species	Activity	Target	Reference
AvrB	<i>P. syringae</i> pv. <i>glycinea</i> race 0	Induces phosphorylation	RIN4/RAR1	[50,52]
AvrBs3	X. campestris pv. vesicatoria race	Transcription activator-like	upa20/Bs3	[35,36]
AvrPphB	P. syringae pv. phaseolicola race	Cysteine protease	PBS1	[14]
AvrPto	P. syringae pv. tomato JL1065	Kinase inhibitor	Pto/EFR/FLS2	[47,49]
AvrPtoB	P. syringae pv. tomato DC3000	E3 ubiquitin ligase	Fen	[29,31]
AvrRpm1	P. syringae pv. glycinea race 0	Induces phosphorylation	RIN4	[48]
AvrRpt2	P. syringae pv. tomato T1	Cysteine protease	RIN4	[18,19]
AvrXa27	X. oryzae pv. oryzae PXO99 ^A	Transcription activator-like	Xa27	[38]
AvrXv4	X. campestris pv. vesicatoria T3	DeSUMOylating cysteine protease	unknown	[24]
GALA	R. solanacearum GMI1000	F-box and LRR domains	ASK(s)	[28]
HopAI1	P. syringae pv. tomato DC3000	Phosphothreonine lyase	MPK3/MPK6	[42]
HopAO1	P. syringae pv. tomato DC3000	Protein tyrosine phosphatase	unknown	[44,45]
HopI1	P. syringae pv. maculicola ES4326	J-domain protein (possible Hsp70 co- chaperone)	unknown	[55]
HopM1	P. syringae pv. tomato DC3000	unknown	AtMIN7 and others	[34]
HopU1	P. syringae pv. tomato DC3000	Mono-ADP-ribosyltransferase	GRP7 and other RNA- binding proteins	[41]
HsvB	Pantoea agglomerans pv. betae 4188	Transcriptional activator-like	unknown	[40]
HsvG	Pantoea agglomerans pv. gypsophilae 824-1	Transcriptional activator-like	unknown	[40]
PthXo1	X. oryzae pv. oryzae PXO99 ^A	Transcriptional activator-like	Os8N3	[39]
PthXo6/7	X. oryzae pv. oryzae PXO99 ^A	Transcriptional activator-like	OsTFX1 OsTFIIA y1	[37]
XopD	X. campestris pv. vesicatoria 85-10	DeSUMOylating cysteine protease	unknown	[22]