Identification of novel Xanthomonas euvesicatoria type III effector proteins by a machine-learning approach

DORON TEPER1,†, DAVID BURSTEIN2, †,‡, DOR SALOMON1,§, MICHAEL GERSHOVITZ2, TAL PUPKO2, * AND GUIDO SESSA1, *

1Department of Molecular Biology and Ecology of Plants, Tel Aviv University, Tel Aviv 69978, Israel
2Department of Cell Research and Immunology, Tel Aviv University, Tel Aviv 69978, Israel

SUMMARY

The Gram-negative bacterium Xanthomonas euvesicatoria (Xcv) is the causal agent of bacterial spot disease in pepper and tomato. Xcv pathogenicity depends on a type III secretion (T3S) system that delivers effector proteins into host cells to suppress plant immunity and promote disease. The pool of known Xcv effectors includes approximately 30 proteins, most identified in the 85-10 strain by various experimental and computational techniques. To identify additional Xcv 85-10 effectors, we applied a genome-wide machine-learning approach, in which all open reading frames (ORFs) were scored according to their propensity to encode effectors. Scoring was based on a large set of features, including genomic organization, taxonomic dispersion, hypersensitive response and pathogenicity (hrp)-dependent expression, 5’ regulatory sequences, amino acid composition bias and GC content. Thirty-six predicted effectors were tested for translocation into plant cells using the hypersensitive response (HR)-inducing domain of AvrBs2 as a reporter. Seven proteins (XopAU, XopAV, XopAW, XopAP, XopAX, XopAK and XopAD) harboured a functional translocation signal and their translocation relied on the HrpF translocon, indicating that they are bona fide T3S effectors. Remarkably, four belong to novel effector families. Inactivation of the xopAP gene reduced the severity of disease symptoms in infected plants. A decrease in cell death and chlorophyll content was observed in pepper leaves inoculated with the xopAP mutant when compared with the wild-type strain. However, populations of the xopAP mutant in infected leaves were similar in size to those of wild-type bacteria, suggesting that the reduction in virulence was not caused by impaired bacterial growth.

Keywords: effector, machine learning, pathogenomics, pepper, type III secretion system, Xanthomonas euvesicatoria.

INTRODUCTION

Many Gram-negative plant-pathogenic bacteria utilize a type III secretion (T3S) system to deliver effector proteins into eukaryotic host cells (Galan et al., 2014). T3S effectors, which are selectively delivered by the secretion apparatus, modulate host cellular processes and contribute to bacterial virulence. They can subvert host signalling pathways, suppress immune responses, modify cytoskeleton structure and interfere with cellular trafficking (Dou and Zhou, 2012; Lindeberg et al., 2014).

Most known plant T3S effectors belong to conserved protein families that are represented in evolutionary distant bacteria encoding the T3S system (McCann and Gutman, 2008). However, a given bacterial pathogen species or pathovar may encode unique T3S effectors, which emerged during the evolution of its pathogenicity. The transcriptional activation of T3S effectors is co-regulated with that of structural components of the T3S system by plant inducible cis-acting regulatory elements present in their promoters (Fenselau and Bonas, 1995; Xiao and Hutcheson, 1994). At the protein level, the molecular architecture of most T3S effectors is modular and includes an N-terminal translocation signal that conveys the effector to the T3S apparatus and then into the host cell, and one or more functional domains (Dean, 2011). N-terminal translocation signals of T3S effectors are very diverse in their primary sequence, but they share common features, such as the frequency of amino acids and residues with certain physicochemical properties (Arnold et al., 2009; Petnicki-Ocwieja et al., 2002). The translocation of T3S effectors is also facilitated by the binding of accessory chaperone proteins which are involved in the stabilization of the effectors and prevention of their premature interaction with other proteins (Lohou et al., 2013). T3S effectors often display eukaryotic features, such as post-translational modifications, intracellular localization signals and enzymatic activities, that enable them to function within host cells (Dean, 2011).

Xanthomonas is a large genus of Gammaproteobacteria that cause disease in hundreds of plant hosts (Ryan et al., 2011). Xanthomonas species and pathovars display high specificity to their hosts and cause significant damage in economically important crops, such as rice, citrus, banana, cabbage, tomato, pepper and bean. With few exceptions (e.g. Xanthomonas albovectors, Xanthomonas sacchari and Xanthomonas cannabis), all...
Xanthomonas pathogenic strains require a functional T3S system to multiply in their host and cause disease. So far, sequence mining of Xanthomonas genomes has identified 52 different families of T3S effectors and a core set of nine effector genes (i.e. xopR, avrBs2, xopK, xopL, xopN, xopP, xopQ, xopX and xopZ) (Ryan et al., 2011; White et al., 2009). Nevertheless, each strain or pathovar encodes a different combination of effectors, which dictates the nature of the interaction with the host and its specificity.

Xanthomonas euvesicatoria (Xcv) is the causal agent of bacterial spot in tomato and pepper plants (Jones et al., 1998). This disease is characterized by the appearance of necrotic lesions on fruits and leaves, and causes significant economic loss in field-grown crops in warm, temperate and tropical areas of the world (Stall et al., 2009). The repertoire of known Xcv T3S effectors includes approximately 30 proteins, most originally identified in strain 85-10 (Thieme et al., 2005). Cellular processes affected by several Xcv 85-10 effectors have been identified. For example, XopD has been implicated in transcriptional regulation (Kim et al., 2013); XopX, XopN, XopB and XopQ have been shown to suppress host immune responses (Kim et al., 2009; Metz et al., 2005; Sonnewald et al., 2012; Teper et al., 2014); and XopJ has been found to interfere with protein secretion and protein degradation (Bartetzko et al., 2009; Ustun et al., 2013). The identification of the entire pool of Xcv 85-10 T3S effectors and an understanding of their mode of action will shed light on the virulence mechanisms of the bacterium and on host defence signalling.

Various approaches have been used previously for the identification of effectors from different important plant pathogens. Initially, the identification was limited to experimental screens based on specific functional characteristics of T3S effectors, such as translocation into the host cell or the ability to stimulate strong immune responses that are often associated with a rapid and localized programmed cell death (the hypersensitive response, HR) on resistant host or non-host plants (Greenberg and Vinatzer, 2003). More recently, the availability of genome sequences for many bacterial phytopathogens has allowed us to uncover large sets of T3S effectors based on homologies to known effectors from other pathogens or the presence of eukaryotic characteristics indicative of a function within the host cell (e.g. da Silva et al., 2002; Salanoubat et al., 2002). Effectors have also been predicted by searching for conserved regulatory sequence motifs, such as hypersensitive response and pathogenicity (hrp) or plant-inducible promoter (PIP) boxes, in the promoter regions, or characteristics of the putative N-terminal translocation signal (Jiang et al., 2009; Petnicki-Ocwieja et al., 2002). These approaches have all been successful to some extent, but are limited by the fact that they are based on a single prediction criterion.

In this study, we utilized a genome-scale computational machine-learning approach to predict novel T3S effectors of Xcv 85-10. To this end, we analysed a large set of features that potentially differentiate effectors from non-effectors. By this analysis, we identified seven novel effectors and experimentally demonstrated their translocation through the T3S system into pepper cells. Follow-up characterization of the identified effectors revealed that XopAP contributes to the development of disease symptoms caused by Xcv on pepper plants. Our study is an important step in the effort to identify the entire pool of Xcv 85-10 T3S effectors and to understand the virulence mechanisms of this bacterium.

RESULTS

Prediction of T3S effectors of Xcv strain 85-10 by a machine-learning approach

To identify novel T3S effectors of Xcv strain 85-10 (Thieme et al., 2005), we used a machine-learning approach similar to that employed previously for the identification of type III secreted proteins from Pseudomonas aeruginosa (Burstein et al., 2015) and type IV effectors from Legionella pneumophila and Coxiella burnetii (Burstein et al., 2009; Lifshitz et al., 2014). The machine-learning algorithm is based on 79 features that potentially differentiate effectors from non-effectors (Table S1A, see Supporting Information). These features cover a wide range of evolutionary, genomics and biochemical characteristics, including, for example, genomic proximity to other effectors, GC content, differential conservation among phytopathogens that do or do not encode a T3S system, amino acid composition at the N-terminus and in the entire protein, T3S-dependent regulation, homology to known T3S effectors of animal- and plant-pathogenic bacteria and similarity to host proteins.

Our algorithm was first trained on known effectors (Table S1B). Specifically, the training step aims to find the features and their relative weights (i.e. the contribution of each feature to the accuracy of the classification) that best differentiate the set of effectors from the rest of the proteins encoded in the genome. Once trained, the algorithm uses the features and their weights to examine all the open reading frames (ORFs) in the genome and provides, as output, a score reflecting the prediction confidence that a certain ORF encodes an effector protein.

The features with the highest weights are listed in Table 1. The first two features in importance analysed the homology of the ORF to known effectors in other phytopathogens. Clearly, effectors are most easily predicted if they are homologous to previously validated effectors of other bacteria. The third best feature considered the similarity of the ORF to the average amino acid composition of Xcv 85-10 T3S effectors. Notably, some highly informative features were negative. For example, the fourth feature in importance examined whether the ORF has a homologue in the genomes of phytopathogenic bacteria lacking the T3S system. This is an informative negative feature because such an ORF is highly unlikely to encode an effector. The fifth feature with high perfor-
Table 1 Performance scores of the top 25 features.

<table>
<thead>
<tr>
<th>Feature</th>
<th>Feature performance*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Degree of homology to a known effector of phytopathogenic bacteria</td>
<td>0.100</td>
</tr>
<tr>
<td>Number of homologues among known effectors of phytopathogenic bacteria</td>
<td>0.051</td>
</tr>
<tr>
<td>Similarity to the average amino acid profile of effectors from Xcv 85-10</td>
<td>0.041</td>
</tr>
<tr>
<td>Degree of homology to the closest homologue in the proteomes of phytopathogenic bacteria not encoding a type III secretion (T3S) system</td>
<td>0.037</td>
</tr>
<tr>
<td>HrpG/HrpX-dependent regulation</td>
<td>0.029</td>
</tr>
<tr>
<td>Degree of homology to a known effector of Xcv 85-10</td>
<td>0.029</td>
</tr>
<tr>
<td>Number of homologues among known effectors of Xcv 85-10</td>
<td>0.025</td>
</tr>
<tr>
<td>Number of homologues in the proteomes of phytopathogenic bacteria not encoding a T3S system</td>
<td>0.024</td>
</tr>
<tr>
<td>Genomic distance from the closest effector</td>
<td>0.014</td>
</tr>
<tr>
<td>Number of effectors in the proximity of 10 open reading frames (ORFs)</td>
<td>0.011</td>
</tr>
<tr>
<td>Relative abundance of valine in the full protein</td>
<td>0.011</td>
</tr>
<tr>
<td>Degree of homology to the closest homologue in bacteria</td>
<td>0.010</td>
</tr>
<tr>
<td>Number of effectors in the proximity of five ORFs</td>
<td>0.008</td>
</tr>
<tr>
<td>Length</td>
<td>0.008</td>
</tr>
<tr>
<td>Hydrophobicity score in the N-terminus</td>
<td>0.007</td>
</tr>
<tr>
<td>Degree of homology to a known effector of Xanthomonas spp.</td>
<td>0.006</td>
</tr>
<tr>
<td>Relative abundance of serine in the full protein</td>
<td>0.005</td>
</tr>
<tr>
<td>Relative abundance of leucine in the N-terminus</td>
<td>0.005</td>
</tr>
<tr>
<td>Number of effectors in the proximity of 15 ORFs</td>
<td>0.004</td>
</tr>
<tr>
<td>GC content</td>
<td>0.004</td>
</tr>
<tr>
<td>Number of homologues among known effectors from pathogenic bacteria of mammals</td>
<td>0.003</td>
</tr>
<tr>
<td>Relative abundance of proline in the N-terminus</td>
<td>0.002</td>
</tr>
<tr>
<td>Number of homologues among known effectors of Xanthomonas spp.</td>
<td>0.002</td>
</tr>
<tr>
<td>Number of effectors in the proximity of 20 ORFs</td>
<td>0.002</td>
</tr>
<tr>
<td>Relative abundance of histidine in the full protein</td>
<td>0.002</td>
</tr>
</tbody>
</table>

*Values obtained from the second training cycle using randomForest quantify the contribution of the feature to the learning accuracy (compared with a learning in which the feature is artificially made non-informative).

mance score examined whether the expression of the ORF is dependent on HrpG and HrpX, which are known master regulators of the T3S apparatus and its effectors in Xanthomonas spp. (Wengelnik and Bonas, 1996; Wengelnik et al., 1996). To predict HrpG- and HrpX-dependent expression, we utilized the data published by Guo et al. (2011) that compared the transcriptomes of Xanthomonas axonopodis pv. citri (Xac306), which is closely related to Xcv85-10, and its mutants in either the hrpX or hrpG genes. The high performance score obtained by this feature suggests that the T3S-dependent expression profiles of effector genes are conserved between related bacterial species and have a strong predictive power for effector identification.

The overall accuracy of the algorithm performance is assessed by a cross-validation procedure. In this procedure, the algorithm is repeatedly trained on different parts of the known data (effectors and non-effectors), and its performance is measured by comparing the resulting prediction with the data that were omitted from the training. Our algorithm achieved high accuracy: the area under the precision recall curve (AUPRC; Davis and Goadrich, 2006) obtained was over 0.82.

To predict novel effectors, the analysis included learning (Table S1C) and validation phases. The ORFs with the highest scores that were not previously reported as effectors were tested for translocation into plant cells. Two of the ORFs predicted as effectors by our machine-learning approach, XCV0324 (XopS) and XCV0442 (XopM), were reported as translocated T3S effectors in parallel to our study (Schulze et al., 2012). After effector validation, we conducted an additional learning analysis (Table S1D) that included the newly identified effectors. The list of high scoring predictions represents a pool of putative additional effectors.

Seven predicted Xcv T3S effector proteins are translocated into plant cells

To examine the translocation of predicted effectors, we utilized a reporter system based on the delivery into plant cells of a truncated form of the Xcv T3S effector AvrBs2 (amino acids 62–574). AvrBs262–574 lacks a translocation signal, but is sufficient to elicit HR in plants expressing the Bs2 resistance gene (Roden et al., 2004). A DNA fragment containing 24 base pairs upstream to the start codon and the complete ORFs of each of 36 predicted effector genes (Table S2, see Supporting Information) were cloned upstream to an AvrBs262–574::haemagglutinin (HA) tag fusion. Plasmids were mobilized into the Xcv hrpGΔΔavrBs2 strain, which constitutively expresses the T3S apparatus and contains a muta-
tion in the avrBs2 gene (Roden et al., 2004). Constitutive expression of the T3S apparatus results in an earlier and stronger HR (Wengelnik et al., 1999), enhancing the sensitivity of the translocation assay. The bacterial strains obtained were tested for elicitation of the HR in the pepper line ECW20R, which encodes a functional Bs2 resistance gene. As expected, Xcv hrpG*ΔavrBs2 expressing the AvrBs262–574::HA fusion did not elicit the HR in Bs2-expressing leaves (Fig. 1A, top panel). However, Xcv hrpG*ΔavrBs2 expressing XCV1196 (XopAU), XCV1197 (XopAV), XCV3093 (XopAW), XCV3138 (XopAP), XCV3786 (XopAX), or XCV4315 (XopAD) fused to AvrBs2::HA induced the HR at 36 h after infection, similar to AvrBs262–574::HA (Fig. 1A, top panel). No HR was observed in leaf areas inoculated with Xcv strains expressing the other tested constructs (data not shown).

To confirm that the HR observed in ECW20R pepper leaves resulted from the specific recognition of the translocated AvrBs262–574 fusion protein, bacterial strains were infiltrated into the pepper line ECW30R, which lacks a functional Bs2 gene. As expected, in this background, the HR was not detected for any of the tested genes (Fig. 1A, bottom panel). To validate that translocation of the identified effectors occurred through the T3S system, the plasmids harbouring the various effector–AvrBs262–574::HA fusions were mobilized into the Xcv hrpG*ΔhrpF strain, which carries a deletion in the hrpF translocon gene, and therefore cannot translocate effector proteins into host cells (Buttner et al., 2002). The Xcv strains obtained were tested for the elicitation of HR in pepper ECW20R leaves. None of the fusion proteins elicited the HR when expressed in the Xcv hrpG*ΔhrpF mutant (Fig. 1A, top panel), indicating that the identified effectors require a functional T3S system for translocation. We assessed the expression of the fusion proteins in Xcv ΔavrBs2 and Xcv ΔhrpF strains by Western blot analysis. As shown in Fig. 1B, all the fusion proteins were expressed in both bacterial strains, with the exception of XopAK, whose expression was not detected despite the fact that the HR was observed in areas infiltrated with Xcv hrpG*ΔavrBs2 carrying...
the XopAK–AvrBs262–574::HA fusion. Together, these results indicate that seven of the 36 tested ORFs encode bona fide T3S effectors.

Conservation of the novel T3S effectors among plant-associated bacteria

Four novel effectors (XopAV, XopAU, XopAX and XopAW) do not have any homologues that have been reported previously as T3S effectors in other bacteria, and therefore define novel effector families. Homologues of three novel effectors have been reported to be translocated in other bacterial species. XopAD and XopAP share homology to the *Ralstonia solanacearum* effectors RipS and RipAL (Peeters et al., 2013), respectively. In addition, XopAK is homologous to the *Pseudomonas syringae* effector HopK1 (Li et al., 2014). A search for homologues within the *Xanthomonas* genus and in other bacterial species revealed different conservation patterns for the newly identified effectors (Fig. 2 and Table 2). Both XopAV and XopAD have homologues in the genomes of most *Xanthomonas* spp. Unlike XopAK, which is conserved in several other phytopathogenic bacteria and in *Rhizobium*, XopAV is unique to *Xanthomonas* spp. XopAP and XopAK show a similar conservation pattern within the *Xanthomonas* genus and are found in all the examined strains of the *axonopodis* clade and in a few other *Xanthomonas* spp. These effectors also appear in other bacteria with different conservation patterns. XopAU has homologues in all the strains of the *axonopodis* and *oryzae* clades. In addition, it is conserved in the genomes of *X. gardneri* and *X. vesicatoria* strains which are causal agents of spot disease in tomato and pepper plants, similar to *Xcv*. Among other bacterial strains, XopAU was found only in *Acidovorax* spp. XopAX is present in *Xcv*, but not in other *Xanthomonas* spp., and only in a few other bacterial species. In this regard, it is important to note that XopAX is encoded in the *pXCV183* plasmid of *Xcv* 85-10 (Thieme et al., 2005), suggesting that this effector was recently acquired by the bacterium via horizontal gene transfer. Finally, XopAW is present in a few *Xanthomonas* spp. of the *axonopodis* clade and in several other plant-pathogenic bacteria and *Rhizobium*. The presence of these novel effector families in several bacterial species encoding the T3S system suggests that they may play important roles in the interaction with host plants and in the establishment of disease.

The genes encoding XopAV (XCV1197) and XopAD (XCV4315) in *Xcv* 85-10 share high similarity to the N-terminus of homologues from other *Xanthomonas* strains. Interestingly, the C-terminus of these homologues is highly similar to the adjacent ORFs of XopAV and XopAD (XCV1198 for XopAV; XCV4314 and XCV4313 for XopAD). Therefore, we hypothesize that the ORFs annotated as XCV1197 (XopAV) and XCV1198, and XCV4315 (XopAD), XCV4314 and XCV4313, were originally two complete...
ORFs that were later truncated by the introduction of early stop codons.

The analysis of conserved domains revealed that four of the novel T3S effectors contain a putative catalytic domain (Fig. S1, see Supporting Information; Table 2): XopAP contains a class III lipase domain, XopAU has a serine/threonine kinase domain, XopAW contains an EF-hand calcium-binding motif and XopAK has a deamidase domain.

**New insights into shared characteristics of Xcv T3S effectors**

The analysis of features that can distinguish between effectors and non-effectors identified several characteristics that are shared by Xcv T3S effectors, but not by the other ORFs in the genome. The analysis of the amino acid composition revealed that several amino acids are differentially represented in the 25 N-terminal residues and in the whole effector protein. The N-terminus of effectors was enriched in serine, but displayed a lower content of leucine as compared with the other ORFs (Fig. 3A). When inspecting the entire protein, 10 amino acids were differentially represented in Xcv effectors as compared with the other ORFs (Fig. 3B). Notably, depletion of valine and enrichment of serine were found as key distinguishing features. The enrichment of serine residues was maintained even when excluding from the analysis the 25 N-terminal amino acids. Thus, the amino acid composition, in both the 25 N-terminal residues and the entire protein, is different between effectors and non-effectors, and contributes to the accurate identification of effectors (Table 1). Additional distinguishing characteristics are the GC content and the distribution of effectors

**Fig. 3** Amino acid composition of *Xanthomonas euvesicatoria* (Xcv) type III secretion (T3S) effectors. The abundance of each amino acid was quantified for effector and non-effector open reading frames (ORFs). The results represent the quantification of amino acid abundance in the first 25 amino acids at the N-terminus (A) or in the whole protein (B). Asterisks indicate a significant difference (P < 0.01) between effectors and non-effectors in both Student’s t-test and Wilcoxon signed-rank test.
in the genome. The average GC content of effector genes (60.3%) was significantly lower than that of other genes (64.6%) \((P < 8 \times 10^{-5}, \text{two-tailed } t\)-test) (Fig. 4A). Moreover, the distribution of effector along the genome is not random \((P < 1.6 \times 10^{-27}, \text{Wald–Wolfowitz test})\) (Fig. 4B), but rather characterized by genomic islands enriched in effectors (Fig. 4C). Notably, seven effector genes \((\text{hpaG, hpaF, xopF1, hpaA, xopD, xopA and xopM})\) are clustered in the \(\text{hrp} \) gene cluster (positions 459 555 to 495 209) and several others reside in small genomic islands.

**XopAP contributes to the development of disease symptoms caused by Xcv in pepper leaves**

To evaluate the contribution of the novel effectors to Xcv 85-10 virulence, we inactivated each of the corresponding genes, with the exception of \(\text{xopAD}\) (for technical reasons), by insertion mutagenesis. The mutant strains obtained were used to infect susceptible pepper plants which were then monitored for the development of disease symptoms and bacterial growth. Disease symptoms were estimated visually and quantified by measuring ion leakage and chlorophyll content, as parameters of cell death and leaf bleaching, respectively. Inactivation of \(\text{xopAP}\), but not that of other genes, reduced the severity of the disease symptoms (Figs 5A and S2A, see Supporting Information), whereas bacterial growth was not affected in any of the mutants (Fig. 5S, see Supporting Information). The appearance of reduced symptoms in leaves inoculated with the Xcv \(\text{xopAP}:\text{Gn}\) mutant was reflected by lower ion leakage and higher chlorophyll content when compared with plants infected with the wild-type Xcv (Fig. 5B, C). Conversely, ion leakage and chlorophyll content in leaves inoculated with the other mutants were similar to that in leaves infected with the wild-type Xcv (Fig. 5S2B, C). To confirm that the effect on Xcv virulence resulted from inactivation of the \(\text{xopAP}\) gene, the Xcv \(\text{xopAP}:\text{Gn}\) mutant was complemented with a plasmid carrying \(\text{xopAP}\) driven by the \(\text{lacZ}\) promoter. As shown in Fig. 5A–C, leaf areas inoculated with the complemented strain displayed similar disease symptoms, ion leakage and chlorophyll content as the wild-type Xcv. Together, these results suggest that XopAP acts as a virulence factor that contributes to the development of disease symptoms, but not to bacterial growth, in plants infected with Xcv.

We then examined whether XopAP can cause a detectable phenotype when overexpressed in the non-host plant \(\text{Nicotiana benthamiana}\). The benefit of this system is that the effect of the overexpressed effector can be evaluated in the absence of other Xcv bacterial proteins. Expression of \(\text{xopAP}\) in \(\text{N. benthamiana}\) leaves by an \(\text{Agrobacterium}\)-mediated transient expression assay caused a bleaching phenotype (Fig. 6A), which was apparent 3 days after \(\text{Agrobacterium}\) infiltration and was reflected by reduced chlorophyll content compared with a green fluorescent protein (GFP) control (Fig. 6B). However, this phenotype was not
Fig. 5 Development of disease symptoms in pepper plants inoculated with the Xcv 85-10 xopAP:GnR mutant. Pepper ECW30R plants were syringe infiltrated with a mock solution or with suspensions [5 × 10^6 colony-forming units (CFU)/mL] of Xcv 85–10, wild-type, mutated in xopAP (Xcv xopAP:GnR) or mutated in xopAP and complemented with a plasmid carrying the xopAP gene. Inoculated areas were monitored for the appearance of disease symptoms and photographed at 6 days post-inoculation (dpi) (A). Cell death (B) and leaf bleaching (C) in inoculated areas were quantified at different dpi by measuring electrolyte leakage and chlorophyll content, respectively. Values are the means ± standard error (SE) of five independent leaf areas. Asterisks indicate a significant difference (Student’s t-test, *P < 0.05) relative to the Xcv 85-10 wild-type treatment. The experiment was repeated three times with similar results.

Fig. 6 Transient expression of XopAP in Nicotiana benthamiana leaves. (A) Nicotiana benthamiana leaves were infiltrated with Agrobacterium expressing XopAP or a green fluorescent protein (GFP) control driven by the CaMV 35S promoter, and photographed at 5 days post-inoculation (dpi). (B) Quantification of leaf bleaching by measurement of the chlorophyll content at 1, 3 and 5 dpi. (C) Cell death was quantified by measurement of electrolyte leakage at 1, 3 and 5 dpi. Values are the means ± standard error (SE) of five independent plants. Asterisks indicate a significant difference (Student’s t-test, *P < 0.05) relative to the GFP control treatment. The experiment was repeated three times with similar results.
accompanied by a significant increase in ion leakage in the inoculated areas (Fig. 6C). Together, these results suggest that XopAP functions as a virulence determinant in Xcv bacteria, and contributes to the development of disease symptoms.

**DISCUSSION**

We utilized a machine-learning approach combined with *in planta* translocation assays to predict and validate novel T3S effectors of the bacterial pathogen Xcv strain 85-10. This analysis led to the identification of seven novel Xcv T3S effectors, four of which belong to previously unknown classes of effector families. Functional analysis of the novel effectors revealed that XopAP contributes to the development of disease symptoms.

Previous studies aimed at the identification of T3S effector proteins have been based on the analysis of a single molecular characteristic to differentiate between effectors and non-effectors. These include, for example, homology to known effectors (Buell et al., 2003; Thieme et al., 2005), the amino acid composition of the protein at the N-terminus (Petnicki-Ocwieja et al., 2002) and the presence of cis-acting elements in the promoter sequence, such as the PIP box or the *hrp* box in ORFs of *Xanthomonas* spp. and *Pseudomonas* spp., respectively (Jiang et al., 2009; Koebnik et al., 2006; Vencato et al., 2006; Xiao and Hutcheson, 1994). However, none of these characteristics is universal to all T3S effectors. For example, we found a perfect or imperfect (containing one mismatch in the TTCGC motif) PIP box only in 26% of *Xcv* 85-10 effector genes. In line with this finding, only one of the newly identified effectors, XopAU, contained a PIP box. Moreover, a PIP box was also present in 3% of non-effector genes and thus, if used as a single differentiating feature, it would have pointed to false-positive candidate effectors. The strength of our learning approach is the combination of a large set of genomic, regulatory and homology-based attributes leading to the identification of T3S effectors that were not discovered by other methods.

The selection of features that can distinguish between effectors and non-effectors sheds new light into the characteristics of *Xcv* effectors. The analysis of the amino acid composition of *Xcv* effectors revealed that several amino acids are differentially represented in the 25 N-terminal amino acids. This is in agreement with observations related to the composition of the translocation signal of T3S effectors of various pathogens (Arnold et al., 2009). Interestingly, several amino acids were also differentially represented in the entire *Xcv* effector proteins when compared with the other ORFs. This phenomenon may reflect physical characteristics that are required for the recognition and translocation of the effectors by the T3S apparatus. Alternatively, it may reflect selective constraints related to the stability or function of the effector within the host cell. An additional notable distinguishing feature identified by machine learning is the presence of homologues in phytopathogens that encode a T3S system. Phytopathogenic bacteria, which are evolutionarily related to *Xcv* but lack a T3S system, such as *Xylella fastidiosa* and *Xanthomonas albilineans*, encode homologues of *Xcv* ORFs involved in housekeeping or niche adaptation, but not effector-encoding genes. Conversely, effector-encoding genes are more likely to have homologues in more evolutionarily distant bacteria that rely on the T3S system.

Several lines of evidence indicate that at least some of the newly discovered effectors have undergone horizontal gene transfer. For example, XopAU and XopAV are located in the same genomic island; XopAX is located on a plasmid and is present exclusively in *Xcv* 85-10 and not in other *Xanthomonas* strains. In addition, the irregular phyletic pattern of other newly identified effectors (Fig. 2) provides a strong indication that these effectors are readily transferred among different *Xanthomonas* strains or lost by certain strains during evolution. Finally, the average GC content of the *Xcv* effectors (60.3%) is significantly (*t*-test *P* = 5.56 × 10⁻⁷) lower than the average of non-effectors (64.6%). On the basis of these findings, it is expected that less-studied strains of *Xanthomonas* may encode novel effectors that are yet to be discovered.

Functional analysis of the novel effectors by insertion mutagenesis revealed that XopAP is a virulence factor in *Xcv* 85-10 that contributes to the development of symptoms in infected pepper plants, but not to bacterial growth. Interestingly, XopAP encodes a putative class III lipase and shares homology to the plant protein DAD1 (Fig. S4, see Supporting Information), which is involved in the biosynthesis of the plant hormone jasmonic acid (JA) (Ishiguro et al., 2001). JA regulates signalling pathways that mediate resistance against necrotrophic pathogens and increases susceptibility to biotrophic pathogens through its antagonistic effect on salicylic acid (SA) defence pathways. It is possible that XopAP manipulates host lipid biosynthesis to enhance JA production, and thereby activates JA-induced defences whilst suppressing SA-induced defences. This hypothesis is supported by recent findings showing that JA signalling components are targeted by several bacterial virulence determinants, including toxins and T3S effectors (Gimenez-Ibanez et al., 2014; Jiang et al., 2013; Uppalapati et al., 2007). The effectors HopX1 and HopZ1, and the coronatine toxin from different *P. syringae* pathovars, directly or indirectly degrade JAZ transcriptional repressors leading to the transcription of JA-dependent genes and the repression of SA-dependent defence responses against bacteria.

Four novel effectors contain conserved eukaryotic domains which may participate in their molecular function. XopAU contains a putative serine/threonine kinase domain at the C-terminus. This is the first identified T3S effector of phytopathogenic bacteria that displays a domain of this kind. Effectors that are protein kinases have been reported previously in bacterial pathogens of mammals. For example, the *Yersinia* effector YopO has been
found to interfere with host G-protein signalling by direct phosphorylation of the Gq α subunit (Navarro et al., 2007). XopAW contains a canonical EF-hand calcium-binding motif and can potentially interfere with host calcium signalling. A homologous EF-hand containing protein, CasA, from Rhizobium etli has been reported to be required for bacterioid development and symbiotic nitrogen fixation in bean (Xi et al., 2000). XopAD appears in Xcv 85-10 as a truncated protein when compared with other Xanthomonas homologues. In several Xanthomonas spp. the effector size spans from 2200 to 2950 amino acids and contains two conserved domains. The N-terminal domain, which shares homology with Xcv XopAD, consists of multiple SKW repeats of semi-conserved 42 amino acids. The C-terminal domain, which is absent in Xcv 85-10 XopAD, encodes a putative RelA-like nucleotide transferase domain, which has been found to be involved in the metabolism of the ppGpp nucleotide in Escherichia coli bacteria (Cashel et al., 1996). It is yet to be determined whether XopAD is still functional in Xcv 85-10 or whether the loss of the RelA-like domain confers a selective benefit to the bacterium. XopAK is predicted to contain a deaminase catalytic domain at the C-terminus. HopK1, the P. syringae homologue of XopAK, has been reported recently to suppress host immune responses and to localize to the chloroplast (Li et al., 2014). However, the involvement of deaminase activity in the function of HopK1 and XopAK remains to be investigated. It is interesting to note that the toxin PMT/ToxA of the bacterium Pasteurella multocida activates G-protein signalling in the host by deamination of the Gαi2 protein (Orth et al., 2009).

In summary, by employing an efficient machine-learning approach based on multiple features, we extended the repertoire of known T3S effectors and paved the way to explore further the molecular strategies used by this pathogen to cause disease in host plants.

**EXPERIMENTAL PROCEDURES**

**Bacterial strains and plant material**

The bacterial strains and plasmids used in this study are listed in Table S3 (see Supporting Information). Escherichia coli and Agrobacterium tumefaciens were grown in Luria–Bertani (LB) medium at 37 and 28 °C, respectively. Xcv was grown in nutrient yeast glycerol medium (Daniels et al., 1984) at 28 °C. The plant cultivars used were as follows: pepper (Capsicum annuum) ECW20R (Kearney and Staskawicz, 1990) and ECW30R (Minsavage et al., 1990) and Nicotiana benthamiana (Goodin et al., 2008). Plants were grown in the glasshouse at 25 °C and kept in long-day conditions (16 h light, 8 h dark).

**Machine-learning scheme**

A machine-learning approach was employed as described previously (Burstein et al., 2009; Lifshitz et al., 2014). The following classification algorithms were used: naïve Bayes (Langley et al., 1992; Morrison, 1990), Bayesian networks (Heckerman et al., 1995), support vector machine (SVM) (Burges, 1998; Vapnik, 1999) and random forest (Breiman, 2001). The ORFs in the genome of Xcv 85-10 were divided into effector ORFs (Table S1B—newly identified effectors were added to the list in the second learning cycle) and non-effector ORFs (after filtering highly similar ORFs). This training set was used to train the four classification algorithms for a predetermined set of features (Table S1A). The ‘Wrapper’ procedure for feature selection (Kohavi and John, 1997) was carried out for each classifier, excluding random forest, which performs feature selection internally. During the training procedure, the performance estimate of each classifier was estimated using the mean AUPRC over 10-fold cross-validation, as described in Lifshitz et al. (2014). The final classification score of an ORF was a weighted mean of its scores from the four classifiers trained on the full training set. The performance estimate of each classifier was used as the classifier’s weight. For random forest, we used the randomForest R package (Liaw and Wiener, 2002), based on the original Fortran implementation described by Breiman (2001). For the other classifiers, the WEKA Java library was used with default parameters. Feature importance was estimated by random forest. The AUPRC values were calculated using AUCCalculator 0.2 (Davis and Goadrich, 2006).

**Features and feature selection**

For each ORF in the Xcv 85-10 genome, 79 features measuring different characteristics were analysed (Table S1A). The features were divided into five groups.

1. Homology features. The protein sequence of each ORF was compared with tailored datasets of protein sequences (Tables S1E–G) and the complete proteome of selected bacteria (Table S1F) using BLAST-P (Altschul et al., 1997). For sequence similarity, two measures were used: (i) the BLAST bit score of the most similar BLAST hit; (ii) the number of BLAST hits with E < 0.01.

2. Physical features. Each ORF was scored for protein length, GC content, amino acid composition in the full protein sequence or in the first 25 amino acids at the N-terminus. The presence of a secretion signal peptide was predicted using the SignalP 4.1 server (http://www.cbs.dtu.dk/services/SignalP/). The hydrophobic/hydrophilic/amphipathic scores were calculated according to the AAlindex of the first 25 amino acids at the N-terminus of the protein (Kawashima et al., 2008).

3. Genome organization features. The genomic distance between each ORF and the nearest known effector was quantified by the number of ORFs present between them. In addition, the number of effector ORFs in the proximity of 5, 10, 15, 20, 25 and 30 ORFs upstream and downstream to the selected ORF was measured.

4. Taxonomic grouping information was obtained using BLink (http://www.ncbi.nlm.nih.gov/Web/Newsltr/Spring04/blink.html).

5. Regulatory features. The 300-bp region upstream of the annotated start codon of each ORF was analysed for the presence of the TATAAT consensus motif, the perfect PiP box motif (TTCGC-N15-TTCGC) and a PiP box motif containing one nucleotide mismatch. In addition, we utilized the list of differentially regulated genes in the hrgG or hrpX deletion mutants, obtained from transcriptomic data of
X. axonopodis pv. citri 306 (Xac) (Guo et al., 2011). Specifically, for each ORF, we examined whether its Xac homologue was differentially regulated.

**Phylogenetic reconstruction**

The phylogenetic tree of Xanthomonas was reconstructed on the basis of a concatenated alignment of three genes: atpD, gyrB and rpoB. The nucleic acid sequences of these genes were globally aligned using Mafft with the 'ginsi' scheme (Katoh and Standley, 2013). The maximum likelihood tree and 100 bootstrap resampling were reconstructed using RAxML (Stamatakis, 2014) under the GTRGAMMA evolutionary model.

**Translocation assay**

For construction of the pAvrBs2-HR plasmid, the HR domain of avrBs2 (amino acids 62–574), fused to an haemagglutinin (HA) tag, was polymerase chain reaction (PCR) amplified from the pMDD1(avnBs2-3Xc:HA) plasmid (Rodent et al., 2004) and cloned into the pB8R1MCS2 plasmid (Kovach et al., 1995). Candidate effector genes and avrBs2-HA were amplified from Xcv 85-10 using gene-specific primers (Table S4, see Supporting information) and cloned into pAvrBs2-HR. The plasmids obtained were mobilized into the Xcv hrgP* ΔavrBs2 (Rodent et al., 2004) and Xcv hrgP* ΔhrpF (Casper-Lindley et al., 2002) strains by triparental mating (Rott et al., 1996).

For translocation assays, overnight bacterial cultures were suspended in 10 mM MgCl2 at an optical density at 600 nm (OD600) of 0.1 and infiltrated into the leaves of 7-week-old pepper plants using a needleless syringe. The elicitation of HR was monitored at 36 h post-inoculation. For quantification of ion leakage, three 1.5-cm-diameter leaf discs were sampled from infiltrated areas of at least five plants, and floated in 10 mL tubes containing 5 mL of double-distilled water for 4 h at 25 °C with shaking. Conductivity was measured using a DDS-12DW conductivity meter (BANTE Instruments, Shanghai, China).

**Mutant construction and complementation**

For the generation of insertion mutants by double crossover in Xcv1196 (xopAU), Xcv1197 (xopAV), Xcv3093 (xopAW), Xcv3138 (xopAP) and XcvId0086 (xopAX), the gene of interest and flanking regions were PCR amplified and subcloned into pBluescript KSII. The gentamycin-3-acetyltransferase gene (Gn*), was then inserted into the gene ORF and the construct obtained was cloned into the pVIK165 suicide vector (Frederick et al., 1998) and transformed into Agrobacterium GV2260. For transient expression, an overnight culture of Agrobacterium was pelleted, resuspended in induction medium (10 mM MgCl2, 10 mM 2-(N-morpholino)ethane-sulfonic acid (MES), pH 5.6, 200 mM acetosyringone) and incubated at 25 °C with shaking. Bacterial cultures were diluted to OD600 = 0.05 and infiltrated into leaves of N. benthamiana plants using a needleless syringe.

**Phylogenetic reconstruction**

The phylogenetic tree was reconstructed on the basis of a concatenated alignment of three genes: atpD, gyrB and rpoB. The nucleic acid sequences of these genes were globally aligned using Mafft with the ‘ginsi’ scheme (Katoh and Standley, 2013). The maximum likelihood tree and 100 bootstrap resampling were reconstructed using RAxML (Stamatakis, 2014) under the GTRGAMMA evolutionary model.

**Plant inoculation and measurement of bacterial growth, relative chlorophyll content and ion leakage**

For virulence assays, 7-week-old pepper plants were infiltrated with bacterial cultures (106 colony-forming units (CFU)/mL when monitoring bacterial growth; 107 CFU/mL when measuring ion leakage and relative chlorophyll content) suspended in 10 mM MgCl2, using a needleless syringe.

For measurement of bacterial growth, three 1-cm2 leaf discs were sampled from at least three plants and ground in 1 mL of 10 mM MgCl2. Bacterial numbers were determined by plating 10 μL from 10-fold serial dilutions and counting the resulting colonies.

For the quantification of leaf bleaching, the relative chlorophyll ratio was measured using a CCM-200 plus chlorophyll meter (Opti-Sciences Inc., Hudson, NH, USA) in three leaves of at least five plants. Chlorophyll readings were standardized to mock treatment.

For the quantification of ion leakage, three 1.5-cm-diameter leaf discs were sampled from infiltrated areas of at least five plants, and floated in 10-mL tubes containing 5 mL of double-distilled water for 4 h at 25 °C with shaking. Conductivity was measured using a DDS-12DW conductivity meter (BANTE Instruments, Shanghai, China).

**Agrobacterium-mediated transient expression**

The xopAP gene was cloned with a C-terminal myc epitope into the pTEX binary vector (Frederick et al., 1998) and transformed into Agrobacterium GV2260. For transient expression, an overnight culture of Agrobacterium was pelleted, resuspended in induction medium (10 mM MgCl2, 10 mM 2-(N-morpholino)ethanesulfonic acid (MES), pH 5.6, 200 mM acetosyringone) and incubated at 25 °C with shaking. Bacterial cultures were diluted to OD600 = 0.05 and infiltrated into leaves of N. benthamiana plants using a needleless syringe.

**Acknowledgements**

This research was supported by the Israel Science Foundation (ISF; grant no. 326/10 to G.S.) and by the Chief Scientist Office of the Israeli Ministry of Science and Technology (grant no. 3-8178 to G.S. and T.P.). D.B. was a fellow of the Converging Technologies Program of the Israeli Council for Higher Education. M.G., D.B. and T.P. were supported by the Edmond J. Safra Center for Bioinformatics at Tel Aviv University. This work benefited from interactions promoted by COST Action FA1208 (https://www.cost-sustain.org).

**References**


Fig. 51 Conserved domains of newly identified effectors. (A) Schematic representation of the XopAP, XopAX, XopAD and XopAK effectors. Boxes represent conserved catalytic domains in Xop proteins. (B) Amino acid conservation in the six SKW repeats (each consisting of 42 amino acids) extending from amino acid 360 to 417 in XopA, XopX, XopD and XopK as suppressors of plant immunity. New Phytol. 195, 894–911.

Fig. 52 Development of disease symptoms in pepper plants inoculated with Xanthomonas euvesicatoria (Xcv) strains carrying mutations in novel effector genes. Pepper ECW30R plants, which are susceptible to Xcv 85-10, were syring wounded with a mock solution or with suspensions of 5 × 10^8 colony-forming units (CFU)/ml of Xcv 85-10 strains, either wild-type or mutated in the indicated effector genes by insertion of a gentamycin cassette (Fig. S3). Whole plant phenotypes were assayed by measuring relative chlorophyll content. Values are means ± standard error (SE) of five independent replicate experiments. The experiment was repeated three times with similar results.

Fig. S3 Contribution of the newly identified effectors to Xanthomonas euvesicatoria (Xcv) virulence. Pepper ECW30R plants were syring wounded with suspensions of 5 × 10^8 colony-forming units of Xcv 85-10.
units (CFU/mL) of wild-type 85-10 and the following mutant strains obtained by insertion of a gentamycin cassette (Gn\(\beta\)): Xcv xopAU:Gn\(\beta\) and Xcv xopAW:Gn\(\beta\) (A); Xcv xopAP:Gn\(\beta\) and Xcv xopAX:Gn\(\beta\) (B); Xcv xopAV:Gn\(\beta\) and Xcv xopAK:Gn\(\beta\) (C). Bacterial populations were determined at the indicated days post-inoculation (dpi). Values are means ± standard error (SE) of five independent plants. The experiment was repeated three times with similar results.

**Fig. S4** Amino acid sequence alignment of Xcv XopAP (XCV3138) and Arabidopsis thaliana DAD1 (Q948R1.1). Protein sequences were aligned with the CLUSTALX1.83 program using default parameters. Conserved amino acids are shaded in black. Amino acids shaded in grey share similar biochemical properties.

**Table S1** Datasets used in the machine-learning scheme. (A) Features used in the machine-learning analysis. (B) Type III secretion (T3S) effectors of Xanthomonas euvesicatoria (Xcv) 85-10 used as the training dataset. (C) Prediction scores of Xcv 85-10 open reading frames (ORFs) in the first learning phase. (D) Prediction scores of Xcv 85-10 ORFs in the second learning phase. (E) T3S effectors of phytopathogenic bacteria. (F) T3S effectors of pathogenic bacteria of mammals. (G) T3S effectors of Xanthomonas spp. (H) Proteomes used for homology analysis.

**Table S2** Xanthomonas euvesicatoria (Xcv) 85-10 candidate effectors tested for translocation into pepper cells.

**Table S3** Bacterial strains and plasmids used in this study.

**Table S4** Primers used in this study.