An ERF121 transcription factor from *Brassica oleracea* is a target for the conserved TAL-effectors from different *Xanthomonas campestris* pv. *campestris* strains

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Transcription activator-like effectors (TALEs), which induce the expression of specific plant genes to promote infection, are the main pathogenic determinants of various *Xanthomonas* bacteria. However, investigation of TALEs from *X. campestris* pv. *campestris*, which causes black rot disease of crucifers, received little attention. In this study, we used PCR-based amplification followed by SMRT amplicon sequencing to identify TALE genes in several *X. campestris* pv. *campestris* strains. Computational prediction in conjunction with quantitative reverse transcription PCR analysis was used to find their targets in the *Brassica oleracea* genome. Transcription factor ERF121, from the AP2/ERF family, was identified as target gene for the conserved TALEs from multiple *X. campestris* pv. *campestris* strains. Several members of this family from diverse plants were previously identified as targets of TALEs from different *Xanthomonas* species. We propose that TALE-dependent activation of AP2/ERF transcription factors promotes susceptibility to *Xanthomonas* through the misregulation of plant defence pathways.

**KEYWORDS**
black rot of crucifers, susceptibility gene, susceptibility hub, transcription activator-like effector, *Xanthomonas campestris* pv. *campestris*
(Streubel et al., 2013), and TALEs that activate such genes are often conserved and widely distributed among *Xanthomonas* species (Oliva et al., 2019). Some TALE-activated S genes are conserved between different plant-*Xanthomonas* systems. The best-studied examples are members of the SWEET gene family, in which TALE-dependent activation was observed in the different plants (rice, citrus, cassava, cotton, pepper) on infection with different *Xanthomonas* species and strains (Hutin et al., 2015; Pérez-Quintero et al., 2015). In some cases, this activation has been shown to be critically important for disease development (Streubel et al., 2013). Another example of conserved TALE targets involves transcription factors. For example, different TALEs from several pathovars of *Xanthomonas citri* activate the expression of the LOB1 transcription factor in citrus, and TALEs from numerous *Xanthomonas* oryzae pv. oryzae strains up-regulate the expression of the TFX1 transcription factor (Hutin et al., 2015). Conserved S genes or S gene families are referred to as susceptibility hubs or pathogenicity hubs (Hutin et al., 2015; Mücke et al., 2019- Tran et al., 2018). Their identification is important not only for improving our understanding of the infection process, but also for developing resistant plants because the modification of conserved TALE targets can lead to more durable resistance.

Because of their role as the main pathogenic determinants, TALEs have been widely studied in *Xanthomonas* pathogens of several crop species, especially rice, citrus, and pepper. *Xanthomonas campestris* pv. *campestris* (Xcc) causes black rot, the most harmful and economically important disease of vegetable brassica crops (Vicente & Holub, 2013). The presence of TALEs has been shown for many Xcc strains of different geographical origins (Denancé et al., 2018; Kay et al., 2005; Mokryakov et al., 2010). However, the role of Xcc TALEs in black rot disease development has not been investigated in sufficient depth, and TALE target genes are largely unknown.

Isolation of TALE genes is quite challenging due to their large size, high GC content, and presence of an array of quasi-identical repeats. To isolate the TALE genes from different Xcc strains, we used two-step high-temperature PCR with conserved primers that anneal far from the central repeat region, which greatly facilitates amplification according to Hommelsheim et al. (2014). In the sample of Xcc strains from the different races, all four strains belonging to Xcc race 6 carried a single TALE gene, whereas strains from races 1, 3, and 4 carried none (Figure 1), which was further confirmed by PCR amplification of shorter TALE fragments (Figure S1). Because the primers used for full-length TALE amplification anneal at the conserved regions of the TALE genes, they apparently can be used for the inexpensive and rapid identification and isolation of TALE genes from diverse *Xanthomonas* species.

MinION (Oxford Nanopore Technologies) amplicon single-molecule real-time (SMRT) sequencing showed that the TALEs from all four strains contained 14 full repeats and a half repeat. According to the TALE class assignment algorithm from the AnnoTale suite (Grau et al., 2016), they were assigned to the TalEN class and named TalEN5 (accession number MT828881, strain XY1-1), TalEN6 (MT828882, strain XY1-2), TalEN7 (MT828883, strain XY2-1), and TalEN8 (MT828884, strain XY2-2). Phylogenetic relationships between TalEN5 to -8 and previously sequenced Xcc full-length TALE genes (Denancé et al., 2018) were studied using the DistAL algorithm (Pérez-Quintero et al., 2015). On the phylogenetic tree, TalEN5 to -8 were located closest to TalEN1, TalEN2, and TalEN3 from the Chinese strains CN-12, CN-17, and CN-18, respectively (Figure 2a). These TALEs also belong to the TalEN class according to the AnnoTale suite and were referred to as members of the Tal15g group by Denancé et al. (2018).

Differences in the nucleotide sequences between TalEN5 to -8 and Tal15g group members occurred mainly at the 3’ part of the coding region downstream of the repeats (Figure S2). Additionally, all repeats in the central part of the Tal15g TALEs were 102 nucleotides in length and encoded 34 amino acid protein repeats, while in TalEN5 to -8, the fifth repeat was 105 nucleotides in length and coded for a 35 amino acid repeat. The combination of repeats of different lengths in a single protein is generally not very common for TALEs from diverse *Xanthomonas* species but is typical for Xcc TALEs (Denancé et al., 2018).

Despite some differences in both the nucleotide and amino acid sequences between TalEN5 to -8 and TALEs from the Tal15g group, the composition and order of RVDs were identical between them (Figure 2b). The only exception was TalEN6 from the XY1-2 strain, which contains an HH RVD instead of an HD RVD in repeat 8; however, repeats with both HH and HD RVDs preferentially bind cytosine in the genomic DNA. The geographical origin of strains Harbouring such an RVD combination varies: China and Belgium.

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**FIGURE 1** TALE gene identification in the *Xanthomonas campestris* pv. *campestris* (Xcc) strains from different races. DK-1 (lane 1), Ram 3–2 (2), Ram 4–1 (3), 276NZ (4), and Tir-2 (5) – race 1; Ram 1–3 (6), Ram 2–2 (7), 306NZ (8), and B-1 (9) – race 3; Bun-1 (10) and Xn-13 (11) – race 4; XY1-1 (12), XY1-2 (13), XY2-1 (14), and XY2-2 (15) – race 6. The strains are referred to Xcc races 1, 3, 4, and 6 according to the report of Ha et al. (2014). Marker, MassRuler High-Range DNA ladder (Thermo Fisher Scientific)
TALEs from the Tal15g group were found in seven of the 22 Xcc strains that harbour any TALE genes according to Denancé et al. (2018). Also, Tal15e had very similar RVD organization (Figure 2b). This suggests that this RVD combination is common among Xcc TALEs. Identification of a larger number of Xcc strains of different geographical origins is desirable to determine whether this RVD combination arose independently. Because the RVD composition defines a set of TALE target genes, the existence of identical RVD arrays in TALEs from multiple Xcc strains of different geographical origins suggests the importance of their targets as S genes upon infection.

The PrediTALE tool (Erkes et al., 2019) was used to identify putative EBEs for TalEN5 to -8) in the genome of the Brassica oleracea line TO1000DH3, which is highly susceptible to Xcc strains XY1-1, XY2-1, and XY2-2 on vein inoculation (Figure S3). The EBE that received the highest score was located in the promoter region of the annotated B. oleracea gene GenBank XM_013739306.1, which codes for the putative ethylene-responsive transcription factor ERF121. The nucleotide sequence of this EBE is optimal for TalEN5 to -8) binding (Figure 3a). This is quite unusual because (with very rare exceptions; Mücke et al., 2019) all known natural EBEs, including those from highly induced genes, harbour single or even multiple mismatches relative to the optimal TALE-binding sequence. Because even a single mismatch can significantly reduce the extent of TALE-dependent gene up-regulation (Cohn et al., 2016; Erkes et al., 2017; Zaka et al., 2018), the perfect match between the TalEN5 to -8) RVD array and the EBE is especially suitable for the activation of ERF121.

In addition to the optimal sequence, the placement of the EBE in the
The EBE is located upstream of the coding sequence close to the putative start codon, which is typical for TALE-activated genes (Grau et al., 2013; Pereira et al., 2014; Streubel et al., 2017). Additionally, TALE EBEs often overlap with TATA boxes (Cohn et al., 2016; Grau et al., 2013; Pereira et al., 2014), and the EBE in the \textit{ERF121} promoter comprises a TATA-like sequence (TATAWA consensus; Bernard et al., 2010; Figure 3a).

Due to the presence of the optimal EBE, \textit{ERF121} was considered as a target for TalEN5 to TalEN8, and the expression of \textit{ERF121} was studied in \textit{Brassica oleracea} line TO1000DH3 on leaf inoculation with XY1-1, XY2-1 (carrying TalEN5) and XY2-1 (carrying TalEN7) strains. A substantial increase in expression of \textit{ERF121} was observed in response to both strains, while the inoculation with TALE-less \textit{Xcc} strains did not lead to \textit{ERF121} activation (Figure 3b). When plants for inoculation were grown under different environmental conditions (greenhouse instead of growth room), even higher \textit{ERF121} activation by XY1-1 and XY2-1 relative to mock was observed (Figure S4), which was not explained by the differences in the basal level of \textit{ERF121} expression between these conditions (Figure S5). One possible explanation of this finding is the unequal induction of a bacterial type III secretion system, which translocates effectors into plant cells, in the inoculated tissues of plants grown under different conditions. The type III secretion system is known to be activated or suppressed by various plant metabolites (Anderson et al., 2014; Tang et al., 2006; Wang et al., 2020; Yuan et al., 2020) whose content could be affected by plant growth conditions. Curiously, in both experiments \textit{ERF121} induction by XY1-1 was observed earlier and was higher than by XY2-1. Because the amino acid sequences are almost identical between TalEN5 and TalEN7, this effect could possibly be explained by some intrinsic physiological differences between XY1-1 and XY2-1 strains that led to different levels of synthesis and/or different effectiveness of translocation of TALEs into the host cells upon colonization of \textit{B. oleracea} plants.

The presence of the optimal EBE and the strong TALE-dependent induction indicate that \textit{ERF121} is a direct TALE target, while the presence of TALEs with this RVD combination in multiple \textit{Xcc} strains of different geographical origins suggests that \textit{ERF121} activation is a widespread \textit{Xcc} pathogenicity strategy. Transcription factors are promising TALE targets because activation of a single such gene can lead to downstream changes in the expression of numerous genes and global shifts in the cellular environment. \textit{ERF121} belongs to the large ERF group within the AP2/ERF transcription factor family (Licausi et al., 2013). TALE-dependent activation of AP2/ERF transcription factors occurs in various plants, such as rice (Pérez-Quintero et al., 2013; Tariq et al., 2019; Tran et al., 2018; Wang et al., 2017), wheat (Peng et al., 2019), and kale, on infection with diverse \textit{Xanthomonas} species. For example, expression of the rice \textit{ERF123} gene is activated by TALEs from multiple \textit{X. oryzae pv. oryzae} and \textit{X. oryzae pv. oryzicola} strains, and \textit{ERF123} is considered to be a susceptibility hub in the rice–\textit{Xanthomonas} interaction (Tran et al., 2018).

Different TALE-activated AP2/ERFs have limited homology to each other and fall into distinct groups within the AP2/ERF family (Table S1), which suggests dissimilar functions. Plants commonly
harbour more than 100 AP2/ERF transcription factors that generally have low sequence similarity and exert diverse functions; however, many of them are associated with resistance to stress factors, including numerous pathogens, which is especially true for ERFs (Licausi et al., 2013). Interestingly, rice ERF123 is up-regulated under chilling stress (Tran et al., 2018), while the expression of ERF121 relatives from Arabidopsis thaliana changes under different stress conditions and treatment with defence-associated plant hormones (Table S2) (Caarls et al., 2017; Feng et al., 2005; McGrath et al., 2005; Pierce & Rey, 2013; Postnikova et al., 2011). Curiously, in the study of Denancé et al. (2018) other defence-associated genes, which code for TGG myrosinases, were found to be the targets for TALEs from the Tal15g group. Myrosinases are well-known defence proteins that participate in the protection of cruciferous plants against herbivores and diverse pathogens (Plisecka et al., 2015; Poveda et al., 2020). According to our data, inoculation with XY1-1 and XY2-1 strains also elevated the expression of myrosinase genes in TO-1000 plants (data not shown).

TALE-induced overexpression of seemingly defence-associated genes may seem counterintuitive. However, activation of some plant defence pathways can be beneficial for pathogenic bacteria, for example through the inhibition of competing microbes in plant tissues (Wu et al., 2019) under field conditions. Also, it is well known that plant responses to biotic and abiotic stresses, and to pathogens with different lifestyles often act antagonistically to each other. Phytopathogenetic bacteria use various type III effectors to activate certain plant biotic and abiotic defence pathways, which leads to the repression of antagonistic defence responses (Kazan & Lyons, 2014; Sowden et al., 2018). Recently, TALE-mediated activation of host factors involved in abscisic acid signalling was demonstrated in different plant-Xanthomonas systems (Mücke et al., 2019; Peng et al., 2019). The objective of TALE-dependent ERF activation may be the inhibition of responses most useful on Xanthomonas attack through the misactivation of competing defence pathways. Indeed, overexpression of certain ERFs inversely regulated resistance to different pathogens and abiotic stress factors (Broekaert et al., 2006; Li et al., 2018; Lu et al., 2020; Tsutsui et al., 2009).

As long as the molecular mechanism of action of most ERFs, including B. oleracea ERF121, remains unknown, without further mechanistic studies we can only speculate about presumed defence pathways induced by TALE-activated ERFs. Generally, ERFs are the major mediators of ethylene signalling (Broekaert et al., 2006). Although ethylene plays an important role in plant defence against pathogens, its role in plant–Xanthomonas interactions is controversial (Kim et al., 2013; Shen et al., 2011; Van Loon et al., 2006). TALE-dependent up-regulation of some ERFs can activate branches of the ethylene response that contribute more to susceptibility than to defence on Xanthomonas attack. We believe that future high-throughput studies of effector-activated ERF regulons could reveal the molecular mechanisms underlying ERF-mediated plant susceptibility. It is likely that the clues can be found at the intersections of the regulons of different ERFs activated on plant-Xanthomonas interactions.

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CONFLICT OF INTEREST
The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

DATA AVAILABILITY STATEMENT
Sequences of TALE genes from this study were deposited in GenBank at https://www.ncbi.nlm.nih.gov/genbank/ under accession numbers MT828881, MT828882, MT828883, and MT828884.

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