EffectorK, a comprehensive resource to mine for *Ralstonia*, *Xanthomonas*, and other published effector interactors in the *Arabidopsis* proteome

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Abstract

Pathogens deploy effector proteins that interact with host proteins to manipulate the host physiology to the pathogen's own benefit. However, effectors can also be recognized by host immune proteins, leading to the activation of defence responses. Effectors are thus essential components in determining the outcome of plant-pathogen interactions. Despite major efforts to decipher effector functions, our current knowledge on effector biology is scattered and often limited. In this study, we conducted two systematic large-scale yeast two-hybrid screenings to detect interactions between *Arabidopsis thaliana* proteins and effectors from two vascular bacterial pathogens: *Ralstonia pseudosolanacearum* and *Xanthomonas campestris*. We then constructed an interactomic network focused on *Arabidopsis* and effector proteins from a wide variety of bacterial, oomycete, fungal, and invertebrate pathogens. This network contains our experimental data and protein–protein interactions from 2,035 peer-reviewed publications (48,200 *Arabidopsis*–*Arabidopsis* and 1,300 *Arabidopsis*–effector protein interactions). Our results show that effectors from different species interact with both common and specific *Arabidopsis* interactors, suggesting dual roles as modulators of generic and adaptive host processes. Network analyses revealed that effector interactors, particularly "effector hubs" and bacterial core effector interactors, occupy important positions for network organization, as shown by their larger number of protein interactions and centrality. These interactomic data were incorporated in EffectorK, a new graph-oriented knowledge database that allows users to navigate the network, search for homology, or find possible paths between host and/or effector proteins. EffectorK is available at www.effectork.org and allows users to submit their own interactomic data.
1 | INTRODUCTION

Plants are continuously confronted with a wide variety of pathogens, including bacteria, oomycetes, fungi, nematodes, and insects. To prevent their proliferation, plants have evolved a complex multi-layered immune system (Jones and Dangl, 2006). Plants are able to recognize highly conserved pathogen-associated molecular patterns (PAMPs) through pattern-recognition receptors triggering induced defence responses collectively known as PAMP-triggered immunity (PTI) (Zipfel, 2014). These responses are usually enough to prevent most potential invaders; however, some pathogens secrete effector proteins to subvert the defence responses and alter diverse cellular processes to ease their proliferation (Ma et al., 2018). Plants, moreover, have evolved several intracellular nucleotide-binding site-leucine-rich repeat (NBS-LRR) receptors recognizing these effectors and activating potent defence responses collectively known as effector-triggered immunity (ETI) (Cui et al., 2015).

Although the interactors and molecular functions of some effectors have been characterized (Büttner, 2016; Giron et al., 2016; Sharpee and Dean, 2016; Vieira and Gleason, 2019), for most effectors they are still unknown. The main factors complicating the large-scale identification and characterization of effector-host protein interactions are the wide diversity of pathosystems, the difficulty in identifying bona fide effector genes, the collective contribution of effector proteins, the complexity of the host responses, and the lack of robust high-throughput techniques. For the model species Arabidopsis thaliana (Ath), to our knowledge, there are only two studies in which systematic effector-host protein interactions at the effectome-scale have been identified (Mukhtar et al., 2011; Weßling et al., 2014). In these studies plant interactors of effector proteins from Pseudomonas syringae (Psy, bacterium), Hyaloperonospora arabidopsidis (Hpa, oomycete), and Glovinomyces orontii (Gor, fungus) were identified by yeast two-hybrid (Y2H) assays. They reported that the effectors of these species converged onto a limited set of Ath proteins. These studies also demonstrated that many effector interactors are important for plant immunity and showed that their importance correlates with the level of effector convergence.

Bacterial wilt, caused by Ralstonia pseudosolanacearum (Ralstonia solanacearum phytophore I, Rps), and black rot, caused by Xanthomonas campestris pv. campestris (Xcc), are listed among the top 10 scientifically and economically important plant bacterial diseases (Mansfield et al., 2012). Both Rps and Xcc are xylem-colonizing bacteria able to infect the model plant Ath (Deslandes et al., 1998; Buell, 2002). They both rely on their type III secretion system for full virulence (Arlat et al., 1991, 1992). This "molecular syringe" allows the pathogen to deliver type III effector proteins (T3Es) directly into the host cell in order to promote disease. The roles of several of their T3Es have been characterized (White et al., 2009; Coll and Valls, 2013), but most knowledge on T3E functions comes from the study of Psy, which resides on leaf surfaces and in the leaf apoplast (Lindeberg et al., 2012; Büttner, 2016). Focusing mainly on a few species offers a partial view of effector biology. It is therefore crucial to expand our studies to other species to grasp the existing diversity of effector proteins and pathogen lifestyles.

To obtain a deeper understanding of the global Ath–effector protein interactome, we conducted three systematic large-scale screenings with T3Es from Rps and Xcc, the first vascular pathogens screened in this manner. Additionally, we conducted an extensive literature survey to gather published Ath interactors of effector proteins from pathogens from four different kingdoms of life: Bacteria, Chromista, Fungi, and Animalia. Combining all these data allowed us to identify 100 new "effector hubs" (i.e., Ath proteins interacting with two or more effectors). Together with Ath–Ath protein interactions retrieved from public databases, we generated an Ath–effector protein network that captures the wide diversity of Ath pathogens. This network allowed us to detect general trends of effector interference with the host proteome. We have created a publicly available interactive knowledge database called EffectorK (for Effector Knowledge) that allows users to access and augment this network.

2 | RESULTS

2.1 | Systematic identification of Arabidopsis interactors of R. pseudosolanacearum and X. campestris effectors

Three Y2H screenings were performed to identify Ath interactors of Rps and Xcc effector proteins. In a first screening, we identified 42 Ath interactors for 21 out of 56 T3Es from Rps strain GMI1000 screened against a library of more than 8,000 full-length Ath cDNAs (8K space). The choice of the 56 Rps T3Es was guided by the available clones at the time of screening. In the second and third screenings, we identified 176 Ath interactors for 32 out of 48 T3Es from Rps strain GMI1000 and 52 Ath interactors for 18 out of 25 T3Es from Xcc strain 8,004 screened against an extended version of the previous library containing more than 12,000 Ath full-length cDNAs (12K space) (Figure S1 and Table S1). Here the choice of Rps T3Es was constrained by a pool maximum imposed by the screening method (see Materials and Methods). T3Es were picked according to their highest degree of conservation within the species complex (Peeters et al., 2013). On average, 10.7 and 5.7 Ath interactors were found per Rps and Xcc T3E. These Ath cDNA libraries had been previously used to test interactions with 57 and 32 effector proteins from Hpa and Psy, respectively, (8K space) and 46 effector proteins from Gor (12K space) (Mukhtar et al., 2011; Weßling et al., 2014). The subset of interactions of effectors from Rps, Xcc, and Gor in the 8K space was used to compare with previously published Hpa and Psy data.
In general, Rps effectors interacted on average with more Ath proteins than the other screened species; however, this difference is only statistically significant when compared to Gor effectors (one-tailed Wilcoxon signed-rank test \( p < .001 \)). These data show that effector proteins from these five different species, on average, tend to interact with a similar number of Ath proteins regardless of kingdom, life style, or effectome size.

### 2.2 Effectors converge onto a limited set of Arabidopsis proteins

We compared the Rps and Xcc effector interactors identified in our screenings with the interactors previously identified for Hpa, Psy, and Gor effector proteins (Mukhtar et al., 2011; Weßling et al., 2014). To avoid bias related to the size of the screened library, we considered only the subset of effector interactors present in the 8K space (Figure S2). At the kingdom level, Bacteria was the kingdom with the highest number of kingdom-specific interactors, with 158 exclusive interactors out of a total of 217 interactors (72.8%), followed by Chromista, with 31 out of 117 (51.7%), and Fungi, with 16 out of 45 (35.6%). In total, 235 out of 299 effector interactors (78.6%) were kingdom-specific. At the species level, when comparing all five pathogens, the percentage of species-specific interactors was 58.9% for Psy, 58.7% for Rps, 51.7% for Hpa, 48.8% for Xcc, and 35.6% for Gor. The total number of species-specific effector interactors was 221 out of 299 (73.9%). These data show that most effector interactors are kingdom- and species-specific.

To evaluate whether Rps and Xcc effectors interact randomly or converge onto a common set of Ath proteins we performed simulations rewiring effector–Ath protein interactions within the 8K space. In these simulations, each effector was assigned randomly as many Ath protein–protein interactions that had been confirmed by classic techniques such as Y2H, co-immunoprecipitation, pull-down, experimental data (Figure 2a). The number of effector interactors observed in our screenings was significantly lower than the numbers obtained in the random simulations for both Rps and Xcc. Similar results had been reported for effectors from Hpa, Psy, and Gor (Mukhtar et al., 2011; Weßling et al., 2014). This shows that, similarly to other species, both Rps and Xcc effectors also interact with a common subset of Ath proteins (i.e., intraspecific convergence).

These random rewiring simulations also allowed us to determine whether effectors from different species interact randomly or converge onto a common set of Ath proteins. For this, the number of common interactors of effectors from different species was compared with the experimental data (Figure 2b). When comparing all three kingdoms, the number of common interactors observed was significantly higher than expected by random rewiring. We then analysed all possible binary, ternary, quaternary, and quinary combinations of species and in all cases the number of common interactors observed was higher than expected randomly (Figure 2c). These differences were all statistically significant except for the common interactors of effectors from Psy and Xcc (\( p = .058 \); Figure S3). This could indicate that these two species are the most different in terms of effector targeting. However, considering that Psy and Xcc are precisely the two species with the lowest number of effectors for which interactors have been identified (Psy: 32 and Xcc: 18 effector proteins), it is likely that the high \( p \) value is caused by the limited sample size. This shows that effectors from all these five species interact with a common subset of Ath proteins (i.e., interspecific convergence).

Altogether, our data indicate that Rps and Xcc effectors converge both intra- and interspecifically onto a set of limited Ath proteins, behaving similarly to effectors from other previously screened pathogen species. This suggests the existence of a convergent set of effector interactors common to evolutionarily distant pathogens that might have a predominant role in the general modulation of the host responses.

### 2.3 Manual curation of the literature to compile Arabidopsis–effector protein interactions

In order to gather more knowledge on Ath–effector protein interactions, we conducted an extensive literature search compiling data from a wider spectrum of bacterial, fungal, oomycete, and invertebrate effector proteins. We only considered published direct protein–protein interactions that had been confirmed by classic techniques such as Y2H, co-immunoprecipitation, pull-down,
FIGURE 2  Effectors converge intra- and interspecifically onto a common set of Arabidopsis thaliana (Ath) proteins. (a) Left: random and intraspecific convergent interactions of effectors (purple squares) with Ath proteins (green circles) can be distinguished by random network rewiring and simulation. Adapted from Weßling et al. (2014). Middle and right: number of Ath interactors in the 8K space of effectors from Xanthomonas campestris pv. campestris (Xcc) strain 8,004 and Ralstonia pseudosolanacearum (Rps) strain GMI1000 found in 10,000 degree-preserving simulations (grey) versus the observed number (red arrow). (b) Left: random and interspecific convergent interactions of effectors from different species (purple and orange squares) with Ath proteins (green circles) can be distinguished by random network rewiring and simulation. Right: number of common Ath interactors in the 8K space of effectors from Chromista, Bacteria, and Fungi found in 10,000 simulations (grey) versus the observed number (red arrow). (c) Scatterplot of observed versus simulated number of common Ath interactors between all binary, ternary, quaternary, and quinary combinations of species. $x = y$ regression is represented with a dashed grey line
protein-fragment complementation, fluorescence resonance energy transfer, or mass spectrometry. We compiled 287 interactions found in 80 peer-reviewed publications involving 218 Ath proteins and 72 effectors from 22 pathogen species (Table S2). Among these 22 pathogens, there were nine bacterial species, mostly proteobacteria but also a phytoplasma species; eight invertebrate species, including both nematodes and insects; four oomycete, and one fungal species. While this collection of species does not represent the full diversity of Ath pathogens, it covers the majority of pathogens for which effector interactors have been reported. We can see that, despite being one of the major pathogen classes, few studies have described fungal effector interactors. This illustrates one of the current gaps in our knowledge of effector interactors in Ath.

2.4 Identification of one hundred new “effector hubs”

To compare experimental and published data, we combined all the interactions curated from the published data together with data from our large-scale Y2H screenings. This resulted in a total of 564 different Ath proteins interacting with pathogen effectors. Our screenings on Rps and Xcc effectors identified 235 interactors. Similar published screenings on Psy, Gor, or Hpa effectors had identified 200 interactors (Mukhtar et al., 2011; Weßling et al., 2014). The literature curation allowed us to identify 218 effector interactors. From the 235 Rps and Xcc effectors interactors found in our screening, 166 were new, which represents 29.4% of the total interactors compiled in this study (Figure 3). This highlights the potential of such systematic and high-throughput large-scale screenings in identifying novel effector interactors. The average effector degree (i.e., the number of effectors interacting with a given Ath protein) was 2.3 but it was unevenly distributed among the 564 interactors, with 350 of them interacting with only one effector (62%) and 14 interacting with more than 10 effectors (2.5%) (Figure S4). The contribution of our experimental data was important in the identification of single interactors as we identified 93 out of the 266 total effectors compiled came from bacterial effectors, as could be expected considering the orthologs of three other new hubs in other plant species also described for their altered infection or other immunity-related phenotype when mutated, silenced or overexpressed. Additionally, the orthologs of three other new hubs in other plant species also produced altered infection phenotypes when silenced or overexpressed. A total of 19 out of the 100 newly identified effector hubs have already been shown to be involved in biotic stress responses. Considering that many of the remaining newly defined effector hubs have been poorly characterized (e.g., hypothetical proteins or descriptions based on homology or belonging to a protein family), it is likely that the number of effector hubs involved in immunity was underestimated. This constitutes a valuable source of novel candidates for further functional characterization.

In terms of organism of origin, most of the 564 interactors are bacterial effector interactors, as could be expected considering that 132 out of the 266 total effectors compiled came from bacteria (Figure S4). In the case of effector hubs, it is noteworthy that 133 out of the 214 hubs described in this work interact with effectors from a single kingdom while there are only 64, 16, and one hubs interacting with effectors from two, three or four different kingdoms, respectively (Table S3). Although biased by the structure of the data, this could suggest kingdom specificity of effector targeting.

Sixteen out of the 100 new effector hub genes have already been described for their altered infection or other immunity-related phenotype when mutated, silenced or overexpressed. Additionally, the orthologs of three other new hubs in other plant species also produced altered infection phenotypes when silenced or overexpressed. A total of 19 out of the 100 newly identified effector hubs have already been shown to be involved in biotic stress responses. Considering that many of the remaining newly defined effector hubs have been poorly characterized (e.g., hypothetical proteins or descriptions based on homology or belonging to a protein family), it is likely that the number of effector hubs involved in immunity was underestimated. This constitutes a valuable source of novel candidates for further functional characterization.
2.6 | Effector interactors tend to occupy key positions in the Arabidopsis–effector protein interaction network

To further investigate the potential impact of effector proteins on the plant interactome, we evaluated the importance of their interactors for the organization of the network. We focused on two main network topology parameters: “degree” and “betweenness centrality” (Figure 4). The “degree” of a protein represents the number of proteins that it interacts with. In this study we differentiated two types of degrees depending on the nature of the interacting proteins: the Ath degree of a given effector or Ath protein (i.e. the number of interacting Ath proteins) and the effector degree for a given Ath protein (i.e. the number of interacting effector proteins). The “betweenness centrality” of a protein is the fraction of all shortest paths connecting two proteins from the network that pass through it. There are two main types of key proteins in a network (Li et al., 2017): (a) proteins important for local network organization, typically showing high degree, and (b) proteins important for the global diffusion of the information through the network, characterized by high betweenness centrality. It had been previously reported in more limited networks that effectors tend to interact with host proteins with high degree and high centrality (Memišević et al., 2015; Li et al., 2017; Ahmed et al., 2018). We then analysed whether this was the case in our network comparing effector interactors with the rest of the Ath proteins (Figure 5). The fraction of proteins decreased rapidly as the Ath degree increased. This indicates that most Ath proteins present low Ath degree and only a few of them show high Ath degree values. This trend was significantly shifted towards higher Ath degree values in effector interactors compared to the rest of Ath proteins. To represent this tendency shift we estimated and compared the area under the curve values of the cumulative distribution of the Ath degree for effector interactors and the rest of Ath proteins (Table 2). Effectively, the area under the curve value of effector interactors was higher than the value of the rest of the Ath proteins. This indicates that effector interactors present generally higher Ath degree than the rest of the Ath proteins. Similarly, we compared the betweenness centrality of these two groups of proteins (Table 2 and Figure S5). Effector interactors also presented significantly higher betweenness centrality values than the rest of the Ath proteins. Altogether, these results indicate that effectors preferentially interact with Ath proteins that are more connected to other Ath proteins and that occupy more central positions in the interactomic network as reported for smaller networks (Li et al., 2017; Ahmed et al., 2018).

2.7 | Effector hubs are better connected and more central than single effector interactors in the Arabidopsis–effector interaction network

We then wanted to test if the Ath degree and betweenness centrality values differed among distinct types of effector interactors (Table 2 and Figure S5). First, we compared multipathogen and pathogen-specific interactors as previously described (Figure S2). Multipathogen effector

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**FIGURE 4** Network topology parameters. Example of a simple interactomic network of three effector proteins (purple squares) and nine Arabidopsis thaliana (Ath) proteins (green circles) to illustrate our definition of “effector hub” (i.e., Ath protein interacting with two or more effectors; highlighted in red) and the three network topology parameters analysed in this study. 1. Effector degree: number of effectors that interact with a given Ath protein; 2, Ath degree: number of Ath proteins that interact with a given effector or Ath protein; 3, Betweenness centrality: fraction of all shortest paths connecting two proteins from the network that pass through a given protein.

| Effector degree | Ath degree | Betweenness centrality |
|----------------|------------|------------------------|
| E1             | -          | 1                      | 0.0                     |
| E2             | -          | 1                      | 0.0                     |
| E3             | -          | 2                      | 0.2                     |
| A1             | 1          | 1                      | 0.2                     |
| A2             | 2          | 1                      | 0.5                     |
| A3             | 1          | 0                      | 0.0                     |
| A4             | 0          | 5                      | 0.6                     |
| Effector hub | Protein name | Effector degree | Description of observed phenotype | Reference |
|-------------|--------------|----------------|-----------------------------------|------------|
| AT1G58100   | TCP domain protein 8 (TCP8) | 13             | Triple tcp8 tcp14 tcp15 mutant showed enhanced *Pseudomonas syringae* strain DC3000 ΔavrRps4 growth | Kim et al. (2014) |
| AT1G71230   | COP9-signalosome 5B (CSN5B) | 8              | Wheat TaCSN5 mutant showed enhanced disease symptoms caused by *Puccinia triticina* | Zhang et al. (2017) |
| AT3G12920   | BOI-related gene 3 (BRG3)    | 7              | brg3 mutant showed increased *Botrytis cinerea* lesion size | Luo et al. (2010) |
| AT5G08330   | TCP domain protein 21 (TCP21) | 7              | Rice OsTCP21 silenced and overexpressing plants showed enhanced and reduced disease symptoms caused by rice rust stunt virus (RRSV), respectively | Zhang et al. (2016) |
| AT5G61010   | Exocyst subunit EXO70 family protein E2 (EXO70E2) | 6              | exo70e2 mutant showed reduced flg22-induced callose deposition. | Redditt et al. (2019) |
| AT4G00270   | STOREKEEPER-related 1 (STKR1) | 6              | STKR1 overexpressing plants showed reduced *Hyaloperonospora arabidopsidis* spore formation | Nietzsche et al. (2018) |
| AT3G01670   | SIEVE ELEMENT OCCLUSION-related 2 (SEOR2) | 4              | *Myzus persicae* feeding from seor2 mutant showed reduced progeny | Anstead et al. (2012) |
| AT5G17490   | RGA-like protein 3 (RGL3)    | 3              | rgl3 mutant showed reduced *P. syringae* growth and increased SA content upon infection | Li et al. (2019) |
| AT3G54230   | Suppressor of abi3-5 (SUA)   | 3              | sua mutant showed enhanced *P. syringae* growth and reduced chitin-induced ROS production | Zhang et al. (2014) |
| AT3G11410   | Protein phosphatase 2CA (PP2CA) | 3              | pp2ca mutant showed reduced *P. syringae* colonization and stomatal aperture. PP2CA overexpressor showed enhanced stomatal aperture | Lim et al. (2014) |
| AT2G17290   | Calcium-dependent protein kinase 6 (CPK6) | 3              | Double cpk5-cpk6 mutant showed enhanced *P. syringae* growth and reduced flg22-induced ROS production | Boudsocq et al. (2010) |
| AT5G41410   | Homeobox protein BEL1 homolog (BEL1) | 3              | Rice OsBIHD1 mutant and overexpressing plants showed increased and reduced Magnaporthe oryzae lesion area, respectively | Liu et al. (2017) |
| AT4G26750   | LYST-interacting protein 5 (LIP5) | 2              | lip5 mutant showed enhanced *P. syringae* growth and disease symptoms and reduced endosomal structure formation upon infection | Wang et al. (2014) |
| AT4G35090   | Catalase-2 (CAT2)            | 2              | cat2 mutant showed increased ROS accumulation upon infection with incompatible *P. syringae* strain | Simon et al. (2010) |
| AT3G02870   | Inositol-phosphate phosphatase (VTC4) | 2              | vtc4 mutant showed reduced *P. syringae* growth | Mukherjee et al. (2010) |
| AT5G53060   | Regulator of CBF gene expression 3 (RCF3) | 2              | rcf3 mutant showed reduced percentage of diseased plants and higher percentage of plant survival upon *Fusarium oxysporum* infection | Dagdas et al. (2016) |
| AT3G02540   | RAD23 family protein C (RAD23C) | 2              | *rad23bcd* mutant (and not *rad23bD*) did not show *Candidatus Phytoplasma*-induced flower virescence and phyllody | MacLean et al. (2014) |
| AT5G38470   | RAD23 family protein D (RAD23D) | 2              | *rad23D* mutant did not show flower virescence and phyllody upon transgenic expression of *C. phytoplasma* SAP54 effector | MacLean et al. (2014) |
| AT2G37630   | Asymmetric leaves 1 (AS1)    | 2              | *as1* mutant showed reduced lesion size caused by *B. cinerea* and *Alternaria brassicicola* and enhanced *Pseudomonas fluorescens* and *P. syringae* growth | Nurmberg et al. (2007) |

*a* Ranked in decreasing order.

*b* Orthologous gene in other plant species, as defined by EnsemblPlants (Kersey et al., 2018), characterized for a role in immunity.
interactors presented significantly higher Ath degree and betweenness centrality compared to pathogen-specific effector interactors. We also compared effector hubs with single effector interactors. Similarly, effector hubs also showed higher betweenness centrality and Ath degree than single effector interactors. This last observation implies that an Ath protein that interacts with several effectors tends also to interact with more Ath proteins. To evaluate whether this is biologically relevant or a bias of the “stickiness” of a protein, we compared the Ath and effector degree values of all effector interactors. Our results showed that these two parameters are not correlated (Pearson correlation coefficient = 0.3221; Figure S6). This suggests that effector hubs interact with more Ath proteins than single effector interactors and that this is not due to a higher stickiness of these proteins. Altogether, these results show that the general tendencies of effector interactors (i.e. more connected to other Ath proteins and more central in the Arabidopsis–effector interaction network) are stronger among effector hubs compared to single interactors, and among multipathogen effector interactors compared to pathogen-specific interactors. This reflects the importance of interfering with key position proteins for the modulation of host–pathogen interactions.

2.8 | Bacterial core T3Es interact with more connected and central Ath proteins

Our work on Rps and Xcc together with previous work on Psy T3Es (Mukhtar et al., 2011) provided a large amount of interactomic data

![Figure 5](image.png)

**Figure 5** Ath degree of Ath proteins interacting or not with effectors. Cumulative distribution of Ath degree of Ath proteins interacting (orange) or not (purple) with effectors. The significance of the difference was validated by one-tailed Wilcoxon signed-rank test. The illustration in the upper right corner represents each compared group. Effectors are represented by squares, Ath proteins by circles and the colour code matches the cumulative distribution graph.

### Table 2 Cumulative Ath and effector degrees and betweenness centrality of different groups of effector interactors

|                          | Effector interactors | Other Ath proteins |
|--------------------------|----------------------|--------------------|
| **Ath degree**           | 2,737                | 1,010              |
| **Betweenness centrality** | 0.23                | 0.033              |

- **Multipathogen effector interactors**
  - **Pathogen-specific effector interactors**
  - **Ath degree**
    - 5,344 vs. 1,790: $p < .0001$
  - **Betweenness centrality**
    - 0.657 vs. 0.136: $p < .0001$

- **Effector hubs**
  - **Single effector interactors**
  - **Ath degree**
    - 4,067 vs. 1,810: $p < .0001$
  - **Betweenness centrality**
    - 0.407 vs. 0.118: $p < .0001$

- **Bacterial core T3Es**
  - **Rest of bacterial T3Es**
  - **Ath degree**
    - 656 vs. 712: $p = 0.4571$
  - **Betweenness centrality**
    - 0.072 vs. 0.074: $p = 0.9198$

- **Bacterial core T3E interactors**
  - **Other bacterial T3Es interactors**
  - **Ath degree**
    - 347 vs. 123: $p < .0001$
  - **Betweenness centrality**
    - 0.369 vs. 0.239: $p = 0.0007$

*Estimated area under the curve of the cumulative distribution of Ath degree, effector degree, and betweenness centrality for each group of proteins as represented in Figures 5, S5, and S7. Estimation based on numerical integration using Simpson’s rule.

*Figure illustrating the cumulative distribution graphic from which the areas under the curve compared were calculated.

*One-tailed Wilcoxon signed-rank test p value of the comparison of the Ath degree, effector degree or betweenness centrality values of all proteins from each compared group.
on bacterial pathogen species for which other resources have been generated, particularly in terms of abundance and diversity of sequenced genomes and thus curated T3E repertoires (Lindeberg et al., 2012; Guy et al., 2013; Peeters et al., 2013; Roux et al., 2015; Dillon et al., 2019; Sabbagh et al., 2019). The most conserved set of T3Es, or “core effectorome,” from each of the three bacterial species has been previously defined (Guy et al., 2013; Dillon et al., 2019; Sabbagh et al., 2019). We then tested whether these subsets of T3Es behaved differently from the rest of bacterial T3Es in terms of interaction with host proteins (Table 2 and Figure S7). Our data showed that core and variable T3Es from the three species do not differ in Ath degree nor betweenness centrality. We then tested if there were any differences between the network properties of the interactors of core T3Es and the other bacterial T3E interactors. Core T3Es interactors showed higher effector degree, Ath degree, and betweenness centrality than the rest of interactors of bacterial T3Es. This suggests that, although core T3Es in general do not have more interactors than the rest of bacterial T3Es, they do interact with more highly connected and central Ath proteins. This might imply that core T3Es have a larger potential to interfere with the host interactome, which could explain the selective pressure to maintain them in the majority of strains.

2.9 | EffectorK, an online interactive knowledge database to explore Arabidopsis–effector interactomic data

In order to facilitate the access and exploration of all the data presented in this work, we have generated EffectorK (for “Effector Knowledge”), an interactive web-based knowledge database freely available at www.effectork.org. The latest version (2 October 2019) contains 49,875 interactions for 8,617 proteins coming from 2,035 publications. Of these, 1,300 are Ath–effector protein interactions. Searches can be done based on a wide range of supported identifiers such as different protein names, NCBI or TAIR accession numbers, PubMed identifiers, and InterPro terms. Additionally, users can also query nucleotide or amino acid sequences directly with BLAST or use accession numbers from other model and crop plants to find homologs within the database. All proteins found by query are then listed in tabular format and hyperlinked to the corresponding interactomic data, external resources, and amino acid sequences. Interactomic data for a given protein can be then explored and downloaded in tabular or graphical format. The graphical representation of the interactomic data depicts proteins interacting with other proteins as nodes interconnected by edges (Figure 6). The size of a node is proportional to the number of interacting proteins, whereas the thickness of an edge represents the confidence of the interaction (i.e. whether the interaction has been detected by one [narrow] or several independent [thick] techniques). This visual interface allows users to expand or re-centre a local subnetwork based on a given protein, get information and access to external resources linked to either a protein (node) or an interaction (edge), or modify the layout and the position of the elements for optimal visualization.

Additionally, EffectorK also allows users to find the shortest paths between two queried proteins in the network.

In order to update, expand, and further improve EffectorK, we encourage users to submit their own interactomic data by filing in and sending a dedicated template available on the site. These data will be verified by the curator team prior to their incorporation in the database. More information about usage, content, and data submission is accessible online, under the tabs “Help” and “Contribute” of the database web server. Please contact us if you have any questions or suggestions by email via contact@effectork.org.

3 | DISCUSSION

In this study we identified systematically Ath interactors of effectors from the vascular bacterial pathogens Rps and Xcc. We combined this information with other Ath interactors identified in similar experimental setups. Additionally, we conducted an extensive literature review to gather published Ath interactors of effectors from a wide variety of pathogens, including other bacterial species and also oomycete, fungal, and invertebrate pathogens. Studying this combined interactomic dataset allowed us to identify new trends of how effectors interfere with the plant proteome and evaluate whether previously described network principles were still supported on a wider scale. We showed that there are no substantial differences in terms of connectivity among the effectomes of five different pathogen species screened systematically (Figure 1). We have reinforced previously described intra- and interspecific convergence of effector

FIGURE 6 Graphical representation of interactomic data on EffectorK. Graphical representation of interactomic data from Xcc effector XopAC (AvrAC). XopAC, in purple, interacts with 36 Ath proteins, in green (only 12 shown for better visualization). The size of a protein node is proportional to its degree (e.g. CSN5B interacts with 50 proteins, BIK1 with six, and APK1A only with XopAC). The thickness of the connecting edges indicates the level of confidence: narrow edges represent physical interaction detected by only one technique, whereas thick edges indicate that the interaction has been detected by at least two independent techniques (e.g. XopAC interaction with BIK1 has been detected by co-immunoprecipitation and pulldown assays, whereas the interaction with APK1A, only by Y2H)
targeting with effectors from two new species (Mukhtar et al., 2011; Weißing et al., 2014), and showed at the same time that most effector interactors are pathogen specific (Figure 2 and S2). Our analyses also supported the previously described tendency of effectors to interact with plant proteins better connected and central in the network (Li et al., 2017; Ahmed et al., 2018), and showed that this tendency is even stronger among effector hubs, multipathogen interactors, and bacterial core T3E interactors (Table 2 and Figure S5).

3.1 | The balance between interactor specificity and convergence

Our data showed that most effector interactors were pathogen-specific (Figure S2) but at the same time effectors converge interspecifically onto a small subset of Ath proteins (Figure 2B,C). These a priori contradictory observations pose an interesting question: what is the balance between the specificity and convergence of effector interactors? At this point, it is impossible to assert whether this specificity is merely caused by the limited number of pathogens screened at the effectome-scale or if it is a reflection of the different and unique ways that each pathogen has evolved to interfere with the host physiology and immunity. This issue can only be addressed by increasing the number of pathogen effectors screened thoroughly and at a large scale. Comparing large datasets of effector interactors of a wider and more diverse set of pathogens would allow evaluating where the balance is between specificity and convergence: (a) If the interactor specificity decreased, it would mean that the effectomes from the different pathogens tend to interact similarly with the host proteome. This was the case when we compared the percentage of species-specific interactors of effectors from Hpa, Psy, and Gor that passed from being 73.9%, 64.9%, and 46.7% in previous works (Mukhtar et al., 2011; Weißing et al., 2014), to 51.7%, 58.9%, and 35.6%, respectively, in the present study (Figure S2). Nevertheless, a total of five screened species is probably not powerful enough to sustain this claim. (b) If, in contrast, the interactor specificity increased with the number of screened species, it would mean that the different pathogens have evolved unique ways to modulate the interaction with the host. If this were the case, deeper analyses comparing related pathogens (e.g. species with similar lifestyle or from the same kingdom) could allow identifying trait-specific interactors (e.g. effector interactors exclusive among vascular pathogen effectors). In any case, to better understand the similarities and particularities on how effectors modulate host processes, it is essential to increase the number of pathogen species screened for effector interactors at the effectome-scale.

3.2 | Large-scale screenings fill the gap in the identification of effector interactors

Including manually curated data from literature has allowed us to broaden significantly the diversity of plant pathogen species compared to similar studies. However, 346 out the 564 described Arabidopsis effector interactors have been identified exclusively through large-scale Y2H screenings against partial libraries of Ath cDNAs. As with any other large-scale screening, the technical limitations together with the incompleteness of the library might have led to an underestimation of the plant–effector interactome of the five screened species (Brückner et al., 2009). The relatively small overlap between the large-scale Y2H screenings and manually curated literature data sets might be a consequence of this limitation (Figure 3). This small overlap illustrates the current knowledge gap in the characterization of the full plant interactome of pathogen effectors. Extensive work will be required to characterize further effector–host protein interactions in other pathosystems. As one of the simplest yet powerful high-throughput techniques for protein–protein interaction detection, our work, like others before, highlights the potential of such large-scale Y2H screenings in the identification of novel effector interactors in an easy, cheap, and systematic manner.

3.3 | EffectorK, an entry point to explore and make sense of plant–effector interactomics

To conclude, our work also provides valuable resources for the plant–pathogen interaction community. We described 540 new Ath–Rps and Ath–Xcc effector protein interactions that allowed us to identify 166 new effector interactors (Table S1). We also manually curated several publications to assemble a collection of 287 Ath–effector protein interactions from a wide variety of pathogens (Table S2). All this allowed us to identify 100 novel effector hubs (Table S3). The contribution to plant immunity of these effector hubs has been described for 19 of them, but remains untested for the majority (Table 1). This constitutes a list of promising candidates for further functional characterization. All these data were integrated in EffectorK, a knowledge database where users can have easy access to the Ath–effector protein interactions and explore the resulting interactomic network visually and interactively. While major efforts were made to capture the maximal diversity on the pathogen side, we limited our work to the Arabidopsis plant model. Thanks to the built-in homology search tools available, users can also use their own data as query regardless of the species studied. It is therefore feasible to use EffectorK as a starting point to build on and extend to crop plant–effector protein interactomics. In the long term, these data could be exploited to better understand how pathogens interact with these crops with the prospect of selecting breeding candidates for improved tolerance or resistance against pathogens.

4 | EXPERIMENTAL PROCEDURES

4.1 | Cloning of Rps and Xcc T3E genes

All the cloning of the T3E genes from Rps and Xcc was performed by BP gateway BP or TOPO cloning (Thermo Fisher Scientific,
Waltham, MA, USA) to generate pENTRY plasmids, which were later transferred into the appropriate Y2H plasmids (Mukhtar et al., 2011) using the LR gateway reaction (Thermo Fisher Scientific). Table S5 contains all the PCR primers and final plasmid identities describing the collection of plasmids used in this study. Gene sequence information from Rps strain GM10000 (GenBank accessions: NC_003295 and NC_003296) (Salanoubat et al., 2002) can be obtained from www.ralstonia-T3E.org (Sabbagh et al., 2019) and from the published genome of Xcc strain 8,004 (NC_007086) (Qian, 2005).

4.2 | Y2H screenings

The Y2H screening was performed in semi-liquid (“8K space” screening) and liquid (“12K space” screening) media as recently reported (Monachello et al., 2019), which is an adaptation of a previously developed Y2H-solid pipeline (Dreze et al., 2010). In both protocols the same low copy number yeast expression vectors and the two yeast strains, Saccharomyces cerevisiae Y8930 and Y8800, were used. The expression of the GAL1-HIS3 reporter gene was tested with 1 mM 3AT (3-amino-1,2,4-triazole, a competitive inhibitor of the HIS3 gene product) unless described otherwise. Prior to Y2H screening, DB-X strains were tested for auto-activation of the GAL1-HIS3 reporter gene in the absence of AD-Y plasmid. In case of auto-activation, DB-X were physically removed from the collection of baits and screened against the (DB)-Ath-cDNA collections using their AD-X constructs. Briefly, DB-X baits expressing yeasts were individually grown (30 °C for 72 hr) in 50-ml polypropylene conical tubes containing 5 ml of fresh selective media (Sc−leucine, Sc−Leu). Pools were created by mixing a maximum of 72 and 50 individual bait yeast strains for the “8K space” and “12K space”, respectively. Subsequently, 120 and 50 µl of these individual pools were plated into 96-well and 384-well low-profile microplates for Ath-cDNA “8K space” and “12K space” collections, respectively. Glycerol stocks of the (AD)-Ath-cDNA “8K space” and “12K space” collections were thawed, replicated by hand-picking or using a colony picker Qpix2 XT into 96-well and 384-well plates filled with 120 and 50 µl of fresh selective media (Sc−tryptophan, Sc−Trp), respectively, and incubated at 30 °C for 72 hr. Culture plates corresponding to the DB-baits pools and AD-collection were replicated into mating plates filled with YEPD media and incubated at 30 °C for 72 hr. In liquid Y2H case (“12K space” screening), mating plates were then replicated into screening plates filled with 50 µl of fresh Sc−Leu−Trp−histidine + 1 mM 3AT media and incubated at 30 °C for 5 days. In order to identify primary positives, the OD_{600} of the 384-well screening plates was measured using a microplate reader Tecan Infinite M200 PRO (Tecan, Männedorf, Switzerland). In semi-liquid Y2H case (“8K space” screening), mated yeast were spotted onto Sc−Leu−Trp−histidine + 1 mM 3AT media agar plates, and incubated at 30 °C for 3 days. Protein pairs were identified by de-pooling of DB-baits in a similar targeted matricial liquid or semi-liquid assays in which all the DB-baits were individually tested against all the previously identified AD-proteins. Identified pairs were picked and checked by PCR and DNA sequencing.

4.3 | Database content and manual curation

Binary interactions between Ath proteins with each other and with pathogen effector proteins were compiled on tabular form keeping track of the protein names and accessions, species and ecotypes/strains of origin, techniques used to detect the interactions and the reference. Ath–Ath protein interactions were compiled from the Arabidopsis Interactome (Dreze et al., 2011; Smakowska-Luzan et al., 2018) and the public databases BioGrid (www.thebiogrid.org [Stark et al., 2006], downloaded in September 2019) and IntAct (www.ebi.ac.uk/intact [Orchard et al., 2014], downloaded in September 2019). We only kept the direct interactions with the evidence codes “co-crystal structure,” “FRET” (fluorescence resonance energy transfer), “PCA” (protein-fragment complementation assay), “reconstituted complex” or “two-hybrid” on BioGrid and “physical association” on IntAct. Ath–effector protein interactions were gathered from our experimental Y2H data together with the similarly produced data on Hpa, Psy, and Gor effectors (Mukhtar et al., 2011; Weßling et al., 2014). In addition, an extensive keyword search on effector–Arabidopsis literature was done to retrieve interactions from 80 published articles. A confidence level was assigned to each interaction depending on the number of independent techniques used in a publication for validation: “1” if the interaction was detected by only one technique and “2” if the interaction was validated by at least a second technique. Some interactions lacked important information but, in order to maximize the extent of our network, several assumptions were taken instead of discarding useful data. First, gene models for Ath proteins were rarely mentioned on publications so we assumed the first zero, the value was set to p<0.0001.

4.4 | In silico analysis

4.4.1 | Computational simulations of random targeting of Ath proteins by single pathogen effectors (intraspecific convergence)

Significance of the intraspecific convergence was tested, comparing our experimental data with random simulations as previously published (Weßling et al., 2014). Briefly, for each effector of Xcc and Rps we assigned randomly the same number of Ath interactors as experimentally observed from the degree-preserved list of 8K proteins. The distribution obtained from 10,000 simulations was plotted and compared to the experimentally obtained data. The p value of the experimental data were calculated as follows: number of simulations where the number of interactors is lower than or equal to experimentally observed is divided by the number of simulations. When the number of simulations with fewer interactors than observed was zero, the p value was set to <0.0001.
4.4.2 | Computational simulations of random targeting of Ath proteins by several pathogen effectors (interspecific convergence)

The significance of the interspecific convergence was tested by comparing our experimental data and previously published data with random simulations as published (Mukhtar et al., 2011; Weßling et al., 2014). Briefly, for each effector of all compared pathogens we assigned the same number of Ath interactors as experimentally observed/published from the list of 8K proteins. The distribution obtained from 10,000 simulations was plotted and compared to experimentally and published data. The \( p \) values of the experimental data were calculated as follows: number of simulations where the number of common interactors between species was higher or equal than the experimentally observed is divided by the number of simulations. When the number of simulations with more common interactors than observed was zero, the \( p \) value was set to \( <0.0001 \).

\[
\begin{align*}
p \text{ value} & = \frac{\text{number of simulations where the number of common interactors \( \geq \) experimentally observed number of interactors}}{\text{number of simulations}} \tag{1} 
\end{align*}
\]

4.5 | Database construction

The databases were built using the software architecture recently described (Carrère et al., 2019). The files submitted by the curator team were automatically checked for typographic mistakes using ad hoc Perl scripts and loaded into a Neo4J database and indexed in an ElasticSearch search engine. Each release was rebuilt from scratch. Data were made accessible through a web interface (see Results and Discussion) built on Cytoscape.js library (Franz et al., 2016). The raw data used for the database setup are available in the “Data” section of www.effectork.org and the source code is available at https://framagit.org/LIPM-BIOINFO/KGBB.

\[
\begin{align*}
p \text{ value} & = \frac{\text{number of simulations where the number of common interactors \( \geq \) experimentally observed number of common interactors}}{\text{number of simulations}} \tag{2} 
\end{align*}
\]

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CONFLICT OF INTEREST

None of the authors has a conflict of interest to declare.

DATA AVAILABILITY STATEMENT

The data that support the findings of this study are openly available in EffectorK at www.effectork.org.

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**SUPPORTING INFORMATION**

Additional Supporting Information may be found online in the Supporting Information section.