UC Davis UC Davis Previously Published Works

Title

Proteomic study of the membrane components of signalling cascades of Botrytis cinerea controlled by phosphorylation

Permalink <https://escholarship.org/uc/item/22b757bh>

Journal Scientific Reports, 9(1)

ISSN 2045-2322

Authors

Escobar-Niño, Almudena Liñeiro, Eva Amil, Francisco [et al.](https://escholarship.org/uc/item/22b757bh#author)

Publication Date

2019

DOI

10.1038/s41598-019-46270-0

Copyright Information

This work is made available under the terms of a Creative Commons Attribution License, available at<https://creativecommons.org/licenses/by/4.0/>

Peer reviewed

SCIENTIFIC REPERTS

Received: 11 January 2019 Accepted: 26 June 2019 Published online: 08 July 2019

Proteomic study of the membrane OPENcomponents of signalling cascades of *Botrytis cinerea* **controlled by phosphorylation**

Almudena Escobar-Niño¹, Eva Liñeiro¹, FranciscoAmil², RafaelCarrasco¹, CristinaChiva 3,4, Carlos Fuentes⁵, Barbara Blanco-Ulate ⁶, Jesús M. Cantoral Fernández¹, Eduard Sabidó3,4 & Francisco Javier Fernández-Acero ¹

Protein phosphorylation and membrane proteins play an important role in the infection of plants by phytopathogenic fungi, given their involvement in signal transduction cascades. *Botrytis cinerea* **is a well-studied necrotrophic fungus taken as a model organism in fungal plant pathology, given its broad host range and adverse economic impact. To elucidate relevant events during infection, several proteomics analyses have been performed in** *B***.** *cinerea***, but they cover only 10% of the total proteins predicted in the genome database of this fungus. To increase coverage, we analysed by LC-MS/MS the frst-reported overlapped proteome in phytopathogenic fungi, the "phosphomembranome" of** *B***.** *cinerea***, combining the two most important signal transduction subproteomes. Of the 1112 membrane-associated phosphoproteins identifed, 64 and 243 were classifed as exclusively identifed or overexpressed under glucose and deproteinized tomato cell wall conditions, respectively. Seven proteins were found under both conditions, but these presented a specifc phosphorylation pattern, so they were considered as exclusively identifed or overexpressed proteins. From bioinformatics analysis, those diferences in the membrane-associated phosphoproteins composition were associated with various processes, including pyruvate metabolism, unfolded protein response, oxidative stress response, autophagy and cell death. Our results suggest these proteins play a signifcant role in the** *B***.** *cinerea* **pathogenic cycle.**

Botrytis cinerea is one of the most destructive phytopathogenic fungi^{1[,2](#page-11-1)} since it can infect a wide range of commercial crops. It has a dual role as both a parasitic and a necrotrophic fungus, and given its life cycle it has been taken as a model in fungal plant pathology. The great versatility of *B*. *cinerea* derives from its ability to adapt its infective cycle to the perceived environmental conditions, using a diverse set of virulence/pathogenicity factors to attack plant tissues and suppress plant defence^{[2,](#page-11-1)[3](#page-11-2)}.

In the post-genomic era, proteomics studies have become a crucial tool to understand the molecular mechanism involved in a wide range of biological processes. The total number of articles on fungal plant pathogen proteomics is still increasing slowl[y4–](#page-11-3)[7](#page-11-4) . Several studies have highlighted the changes that take place in the *B*. *cinerea* proteins during its infection cycle, as well as their relationship with known pathogenicity factors, and the role of "moonlight proteins", e.g. GADPH, in the fungal proteome^{[8](#page-11-5)}. However, the proteins identified in proteomics studies of *B. cinerea* only cover around 10% of the total proteins predicted in the genome database⁹. In an attempt to increase this coverage, several different subproteomes have been characterized, particularly phosphoproteins^{10,11}, and membrane proteins¹². To further investigate the protein components of transduction pathways, a study has

1 Andalusian Center for Grape and Grapevine Research (IVAGRO), Microbiology Lab, University of Cadiz, Puerto Real, 11510, Spain. ²Bioinformatics Unit, SCAI, University of Córdoba, Ramón y Cajal Building, Rabanales Campus, 14071, Córdoba, Spain. ³Proteomics Unit, Centre for Genomic Regulation (CRG), 08003, Barcelona, Spain. ⁴Proteomics Unit, Universitat Pompeu Fabra (UPF), Barcelona, 08003, Spain. ⁵Proteomics Unit, SCAI, University of Córdoba, Ramón y Cajal Building, Rabanales Campus, 14071, Córdoba, Spain. ⁶Department of Plant Sciences, University of California, Davis, CA, 95616, USA. Almudena Escobar-Niño and Eva Liñeiro contributed equally. Correspondence and requests for materials should be addressed to F.J.F.-A. (email: franciscojavier.fernandez@uca.es)

been undertaken to obtain the subproteome (phosphoproteins) of a subproteome (membrane proteins) with the object of revealing those membrane proteins that are controlled by phosphorylation. We have named this subproteome the "phosphomembranome" of *B*. *cinerea*. As far as we know, this is the frst overlapping proteome approach reported in the study of phytopathogenic fungi.

In *B. cinerea* the infection process requires efficient communication between its biochemical machinery and the surrounding environment. Tis dialogue is mediated by signal transduction cascades, such as the Small GTPase, Ca/calmodulin-dependent and MAPK signalling pathways, which transduce the perceived external con-ditions into intracellular signals. In most cases these involve phosphorylation events^{[13](#page-12-0)}. It has been established that, among the known post-translational modifcation (PTM) processes, phosphorylation is the main mechanism of signal transduction and modification of enzymatic activity in living organisms 14 . Furthermore, this PTM has been reported to have an important role in infection processes of fungal plant pathogens, including *B*. *cinerea*[13](#page-12-0),[15](#page-12-2). It is also known that membrane proteins play an important role as receptors and transporters activat-ing signalling cascades that connect the fungal response to changes in the environment^{[16](#page-12-3),[17](#page-12-4)}. In *B. cinerea*, most of the pathogenicity/virulence factors that have been described are related to phosphorylation/dephosphorylation activity, [\(http://www.phi-base.org](http://www.phi-base.org))[18](#page-12-5); these include MAP kinases, the RAS superfamily, GTPases and histidine kinases^{[19](#page-12-6)-[21](#page-12-7)}. About 30% of all virulence factors are integral membrane proteins with transmembrane alpha-helix (TMH) domains^{22,23}.

All this existing knowledge points to the importance of both the membranome and the phosphoproteome of *B*. *cinerea* for understanding key points in the regulation of the infection process. In this paper, for the frst time, we present a combined analysis of both subproteomes (now jointly named the "phosphomembranome"), with the objectives of increasing the coverage of membrane proteins identifed, and of elucidating the main components of the transduction pathways that control the infection process. Towards these goals, we have analysed the phosphorylation of membranome proteins in *B*. *cinerea* under diferent virulence conditions, induced by plant-based elicitors, including glucose (GLU) as a constitutive stage, and deproteinized tomato cell wall (TCW) as a virulence inductor. Both conditions are known to induce diferent fungal responses in the *B*. *cinerea* phosphoproteome and membranome^{[11,](#page-11-8)[12](#page-11-9)}. Interestingly, the differences identified in the composition of membrane-associated phosphoproteins in each assayed condition may indicate the relative importance of the regulation of particular factors - the pyruvate metabolism, unfolded protein response (UPR), oxidative stress response, autophagy and cell death during the infection process of *B*. *cinerea*.

Results and Discussion

Membrane phosphoprotein identifcation. Diferences in the phosphomembranome of *B*. *cinerea* were analysed using mycelium harvested under two diferent pathogenicity stages: glucose (GLU) as a constitutive stage, and deproteinized tomato cell wall (TCW) as a virulence inductor, as has been previously described^{[11,](#page-11-8)12}. Membrane proteins were isolated and phosphorylation was enriched as described in the Methods section. Afer LC-MS/MS analysis, a total of 2318 phosphopeptides (2073 diferent peptides) were identifed under the two conditions assayed (Supplementary Data Table S1), belonging to a total of 1112 membrane-associated phosphoproteins. In order to provide greater consistency to our comparative study only those proteins identifed in at least three of the four biological samples in each assayed condition were considered for further analysis. According to this, a total of 580 phosphopeptides were considered in our research. Afer qualitative analysis, 64 phosphoproteins (72 diferent phosphopeptides, 74 including specifc phosphorylation) were exclusively identifed or overexpressed under GLU conditions (Supplementary Data Table S2a) and 243 phosphoproteins (324 diferent phosphopeptides, 327 including specifc phosphorylation) exclusively identifed or overexpressed under TCW conditions (Supplementary Data Table S2b). Seven phosphoproteins found under both GLU and TCW conditions, exhibited specifc phosphorylation patterns for each condition, and therefore, they were also considered as phosphoproteins exclusively identified or overexpressed (Supplementary Data Table S2). The mass spectrometry data obtained show diferences in the composition of the phosphomembranome in *B*. *cinerea* depending on the carbon source assayed. The mass spectrometry proteomics data have been deposited in the ProteomeXchange Consortium^{[24](#page-12-10)} via the PRIDE partner repository, with the dataset identifier PXD010961.

Comparing the 1112 membrane-associated phosphoproteins identifed in this work with those proteins identifed in our previous proteomics analysis, the *B*. *cinerea* phosphoproteome and membranome[11](#page-11-8),[12](#page-11-9), a total of 436 (39.21%) new proteins have been identifed (Fig. [1](#page-3-0)). Tese new proteins mean a 2.6% increase in the 10% of the total proteins predicted in the genome database⁹ covered by the previous proteomic analyses. Focusing our attention only on those proteins exclusively identifed or overexpressed under each assayed condition, the comparative analysis showed 7 (10.94%) and 123 (50.62%) proteins identifed under GLU and TCW, respectively, which were not identifed in the phosphoproteome or the membranome (Supplementary Data Table S3). Between the proteins exclusively identifed or overexpressed under each condition only 7 were among the pathogenicity/virulence factors described at (<http://www.phi-base.org>): (i) 2 of the proteins exclusively identifed or overexpressed under the GLU condition (Bcatg3 and BcSAK1); (ii) 5 of the proteins exclusively identifed or overexpressed under the TCW condition (BcCLA4, BcSAK1, BcCRZ1, Bos5 and Bcbrn1). Putting all these results together, it is evident that the phosphomembranome has been a useful approach to identify new proteins potentially implicated in the pathogeny/virulence of *Botrytis cinerea*.

Topology analysis of membrane-associated phosphoproteins. The phosphoproteins identified were classifed based on topology prediction algorithms to determine the protein's role as a membrane component. Integral membrane proteins were identifed by predicting the presence of transmembrane α-helix domains (TMHs). Proteins without predicted trans-membrane domains were further analysed to fnd other relationships with the membrane, such as the presence of secretory signal peptides or other covalent associations (palmitoylation, prenylation, myristoylation and GPI anchor). Proteins predicted to include these modifcations were

considered to be potentially associated membrane proteins, as has been previously described¹². The proteins not assigned to any of the previously-mentioned categories were classifed as peripheral and/or cytosolic proteins.

Based only on the presence of predicted proteins with TMHs, the topological prediction analysis showed that integral membrane proteins with TMHs account for 42.2% and 14.4% of the proteins exclusively identifed or overexpressed under GLU and TCW, respectively (Fig. [2a,b\)](#page-4-0). Because this kind of approach is new in fungal proteomics, the comparison of our data with other fungal species is not possible. A previous reported analysis using human microsomes from K562 cells showed that only around 30% of the membrane proteins isolated have transmembrane regions; this could imply that a large number of non-integral proteins are associated in some way with membranes, suggesting a signifcant role for peripheral proteins and lipidated proteins in the biology of organisms[25](#page-12-11). Further, of exclusive or overexpressed proteins, 7.8% and 14.4% were classifed as secreted proteins under each condition, GLU and TCW, respectively; 37.5% and 40.3% of exclusive or overexpressed proteins were classifed as proteins potentially associated with the membrane by lipidation, under each condition, GLU and TCW, respectively; and the remaining exclusive or overexpressed proteins, 12.5% and 30.9%, were classifed as peripheral/cytosolic proteins under GLU and TCW, respectively (Fig. [2a,b](#page-4-0)). Of the predicted proteins potentially associated with the membrane under each condition, the analysis showed that palmitoylation is the most typical lipidation in both conditions (Fig. [2a,b\)](#page-4-0); this fnding agrees with our previous analysis of the membranome of *B*. *cinerea*[12.](#page-11-9) Taking all the identifed proteins with positive signals for lipidation together with secreted proteins (since many secreted proteins can be anchored to the membrane during their secretion²⁶), the percentage of identifed membrane-associated proteins increases from 42.2% to 87.5% under the GLU, and from 14.6% to 69.1% under the TCW condition. Comparing this topological prediction to our previous results in the analysis of the membranome of *B*. *cinerea*, the percentage of predicted membrane proteins is higher, especially under GLU conditions, at 87.5% compared with 42% under GLU, and at 69.1% compared with 63% under TCW^{12} TCW^{12} TCW^{12} .

For each condition assayed, the transmembrane proteins exclusively identifed or overexpressed under each condition were classified by their number of predicted TMH domains. These numbers were then compared with the predicted profle of TMH-containing proteins present in the *B*. *cinerea* genome database[27](#page-12-13), and with our previous results regarding the membrane proteome¹². The number of proteins with TMH domains in the phosphomembranome of *B*. *cinerea*, identifed under our conditions, cover only 3.7% of the total of 1659 proteins with predicted TMH domains in *B*. *cinerea*, compared with 28% of the proteins with TMHs identifed in our previous analysis of the *B. cinerea* membranome¹². In spite of the low coverage, these results show a similar distribution in terms of number of TMHs per protein, compared with proteins present in the *B*. *cinerea* genome database, which span the range from 1 to 14 TMHs (Supplementary Information Fig. S1); this fnding shows that our membrane protein isolation and phosphopeptide enrichment protocols do not produce any artifcial increase in the number of transmembrane domains. Tis result also shows that there is no special relationship between the number of TMHs and protein phosphorylation.

Analysis of phosphorylation site variation depending on the carbon source. The differences in the number of phosphopeptides and in the phosphorylated site (P-Site) pattern between the assayed conditions was studied. Under the GLU condition, 58 of the 64 exclusive or overexpressed phosphoproteins identifed presented a unique P-site, of which 44 (76%) were found in serine, 13 (22%) in threonine and 1 (2%) in tyrosine residues. Similar results were obtained under the TCW condition, in which 178 of the 243 phosphoproteins

Figure 2. Topological classifcation. (**a**) Exclusive and over-expressed proteins under the GLU condition; and (**b**) Exclusive and over-expressed proteins under the TCW condition. Keys: TMH = integral membrane proteins; $Myr+Pren=myristoylation$ and prenylation sites; $Myr+Palm=myristoylation$ and palmitoylation sites; $Palm + Pren = palmitoylation$ and prenylation sites; $Myr + Palm + Pren = palmitoylation$, myristoylation and prenylation sites; $Palm + GPI = GPI$ and palmitoylation sites; Myristoylation = myristoylation sites; Prenylation=prenylation sites; Palmitoylation=palmitoylation sites.

Figure 3. Distribution of phosphorylation sites identifed. (**a**) Distribution of phosphoproteins presenting a unique P-site by the phosphorylated amino acid: (i) external circumference represents the percentage of phosphoproteins with a unique P-site in Serine (red), Treonine (blue) or Tyrosine (green) under the TCW condition; (i) internal circumference represents the percentage of phosphoproteins with a unique P-site in Serine (red), Threonine (blue) or Tyrosine (green) under the GLU condition. (**b**) Frequency distribution of phosphoproteins according to the number of phosphorylation sites identifed.

presented a unique P-site, with 153 (86%) phosphorylated in serine and 25 (21.42%) in threonine (Fig. [3a](#page-4-1)). More than 70% of the identifed phosphorylated membrane proteins contain only one P-site (Fig. [3b](#page-4-1)), and the largest number of unique phosphorylation P-sites are in serine; these results agree with our previous data on the *B. cinerea* phosphoproteome^{[11](#page-11-8)}. Our results report differences in P-site distribution depending on the available carbon source in our study conditions; the largest number of serine phosphorylated proteins were found under TCW; Proportionately more threonine phosphorylated proteins were found under GLU than under TCW; and no unique tyrosine P-sites were found under TCW conditions. Compared to previously reported result for *B*. *cinerea*, *A*. *brassicicola*[10](#page-11-7) and *C*. *albicans*[28](#page-12-14), grown in culture media with mixed carbon sources, our data showed slightly higher phosphothreonine levels under GLU conditions and lower phosphotyrosine levels under TCW conditions. Tis confrms that phosphorylation of signal cascades components plays an important role in the regulation of *B*. *cinerea* metabolism with the object of adapting to its particular environmental conditions, such as a diferent carbon source.

Among the exclusive or overexpressed phosphoproteins identified, 7 proteins were common to both assayed conditions but presented diferent patterns in phosphorylation, with diferent phosphopeptides and/ or diferent phosphorylation sites found in the same phosphopeptide depending on the assayed carbon source (Supplementary Data Table S2). Tese variations in the pattern of phosphorylation could afect the activity of proteins, and suggests specifc regulation of the biological processes in which these proteins are involved via post-translational modifcations. Four of these proteins were identifed as: (i) stress-activated mitogen-activated protein kinase SAK1; (ii) acetyl-CoA carboxylase; (iii) putative major protein of the Woronin body; and (iv) pyruvate carboxylase. It is notable that two proteins, acetyl-CoA carboxylase and pyruvate carboxylase, involved in pyruvate metabolism are present. Tis fnding, taken with the results of the subsequent analysis of clusters (MCODE), the GO and KEGG annotation, could indicate that pyruvate metabolism regulation plays an important role in the adaptation of *B*. *cinerea* to changes in environmental conditions, and therefore in the overall pathogenesis process.

Gene ontology classification. To determine the biological relevance of the identified phosphorylated membrane proteins of *B*. *cinerea* that were afected by the changes in the carbon source used, exclusive or over-expressed proteins identifed under each condition were categorized according to their specifc gene ontology (GO) annotations, by Biological Process (BP) and Molecular Function (MF) (Fig. [4](#page-6-0)). The GO classifcation by BP (Fig. [4a\)](#page-6-0) showed 13 diferent categories present in only one of the carbon sources; one of these (autophagy) appeared only under the GLU condition, and the remaining 12 categories, including cell death, appeared only under the TCW condition. There were other categories that appeared under both conditions, but with several diferent subcategories under each of the two conditions assayed (Supplementary Data Table S4); among these categories are: (i) small molecule metabolic process; (ii) sulphur compound metabolic process; and (iii) response to stress. Under the GLU conditions the category "small molecule metabolic process" included "acetyl-CoA biosynthetic process from pyruvate"; in contrast, under TCW this category included "sulphur amino acid biosynthetic process". The category "sulphur compound metabolic process" under the GLU condition presented the subcategory "acetyl-CoA biosynthetic process from pyruvate"; conversely, under TCW this category showed the three subcategories "sulphur amino acid biosynthetic process", "sulphate assimilation" and "hydrogen sulphide biosynthetic process". Finally, the category "response to stress" presented phosphoproteins related to response to oxidative stress only under the TCW condition. The presence of membrane-associated phosphoproteins related to autophagy under the GLU condition agrees with previous reports, which illustrate the importance of the regulation of this catabolic process in the progress of infection in pathogenic fung[i29](#page-12-15). Additionally, under GLU the existence of the subcategory "acetyl-CoA biosynthetic process from pyruvate", derived from the presence of the phosphoprotein pyruvate dehydrogenase (PDH), suggests that PDH plays a pivotal role in the regulation of the pyruvate metabolism, and therefore in pathogeny, coinciding with previous results reported in *Fusarium graminearum*[30.](#page-12-16) Similarly, the presence of phosphorylated proteins identifed under the TCW condition, related to "cell death", "sulphur amino acid biosynthetic process" and "hydrogen sulphide biosynthetic process", suggests that the regulation of these processes could be essential in the progress of the infection, agreeing with previous reports, in which the association of these processes with infection in phytopathogenic fungi has been shown^{28,[31–](#page-12-17)} 33 . According to its Molecular Function (Fig. [4b\)](#page-6-0), the presence of TCW as a sole carbon source seems to increase the phosphorylation of those membrane proteins associated with the endoplasmic reticulum (ER) misfolded or unfolded protein accumulation response. These proteins are included into 4 Molecular Function categories (Unfolded protein binding, ubiquitin-like protein binding, mRNA binding and peptidase activity) and into 1 category from Biological Process classification (protein binding). These Molecular functions and the Biological Process mentioned are related to the unfolded protein response (UPR), the endoplasmic reticulum-associated degradation pathway (ERAD) and ubiquitin-dependent autophagy elimination of proteins aggregates, which are known to be connected[34–](#page-12-19)[36](#page-12-20). UPR, ERAD and its cooperation are known to be essential for infection processes in several fungi pathogenic to human and plants^{37–39}, but need to be elucidated in *B. cinerea*.

Protein interaction analysis. In order to understand better the processes differentially affected by the type of carbon source, we performed a protein interaction analysis using the STRING database and Cytoscape^{[40](#page-12-23),41}. Performing this analysis we searched our identifed proteins against *B*. *cinerea* proteins in the STRING database and predicted their physical and functional interactions using seven types of evidence. By using the exclusive and over-expressed proteins in each condition, the analysis returned a network of 108 nodes and 262 connecting edges for TCW (Supplementary Information Fig. S2a) and a network of 14 interacting nodes and 11 connecting edges for GLU (Supplementary Information Fig. S2b). Tese networks were used to perform a deeper analysis in order to identify highly related interactive proteins (clusters) by using MCODE^{[42](#page-12-25)}. Only one cluster was identified for GLU (Table [1](#page-6-1)), which contains proteins involved in the pyruvate metabolism; this fnding coincides with the

Figure 4. Gene ontology (GO) classification of membrane-associated phosphoproteins exclusively identified or over-expressed in *B*. *cinerea* under GLU and TCW condition, as sole carbon source. (**a**) Biological process classifcation, and (**b**) Molecular function classifcation. Relative abundance, on the y-axis, refers to the percentage of membrane-associated phosphoproteins identifed in each category relative to the total number of phosphoproteins GO annotations.

Table 1. Components of the single protein cluster discovered under the GLU condition using MCODE. *The information presented in the column "GO/KEGG function" is obtained from GO and KEGG analysis.

results from the GO annotation and diferential pattern of phosphorylation in common proteins. Tis result reinforces the importance of the pyruvate metabolism regulation in pathogeny. Under the TCW condition 7 clusters were identifed (Table [2](#page-7-0)); one of these was related to the proteasome, agreeing with the GO results described above, that showed under the TCW condition phosphoproteins related to the UPR or ERAD, which are known to be essential for infection processes in several pathogenic fungi^{[37](#page-12-21),[38](#page-12-26)}.

Pathways analysis. With the object of revealing the differences between the two conditions assayed in the biological pathways expressed, a pathway analysis of proteins identifed (exclusive and overexpressed) was

Table 2. Components of the 7 protein clusters discovered under the TCW condition using MCODE. *The information presented in the column "GO/KEGG function" is obtained from GO and KEGG analysis.

performed using KEGG⁴³. The KO numbers assigned by BlastKOALA 2.1 were mapped to the network datasets KEGG pathway, Reconstruct Pathway and Search&Color Pathway; the latter was used to search against all pathway maps in the "bfu organism (*B. cinerea*)" category. The results from using the Reconstruct pathway tool were divided into 2 categories. The category of "Genetic information" (Supplementary Information, Fig. S3) corroborated the presence of phosphorylated membrane proteins involved in functions related to the proteasome, ubiquitin-mediated proteolysis and protein processing in endoplasmic reticulum under the TCW condition, matching the previous GO annotation and MCODE results. The importance of the UPR and ERAD in the pathogeny of *Aspergillus fumigatus*, *Cryptococcus neoformans* and *Alternaria brassicola* has been described previously^{37–39}. However, the role in *B. cinerea* remains to be elucidated. The category "Environmental information" processing" (Supplementary Information, Fig. S4) showed the existence of membrane-associated phosphoproteins related to mTOR and cAMP signalling pathways under the GLU condition. Tese signalling pathways are known to be implicated in the regulation of autophagy^{[29](#page-12-15)}. The presence of phosphorylated membrane proteins related to this biological process under GLU has also been demonstrated in our GO results. Taken together, these results suggest that autophagy is an important route in the regulation of *B*. *cinerea* pathogeny, as previously reported[44,](#page-12-28)[45.](#page-12-29) Search&Color Pathway results (Supplementary Data Table S5) showed similar results, supporting: (i) the importance of regulation of the UPR, with the presence under the TCW condition of phosphoproteins implicated in the proteasome, ubiquitin-mediated proteolysis and protein processing in endoplasmic reticulum; and (ii) the importance of the regulation of autophagy, by the presence under the GLU condition of phosphoproteins related to autophagy.

Proteome Mining. *Signal transduction*. The signalling cascade is essential in the pathogeny because it links the fungal response with the changes in the surrounding environment. Qualitative analyses of proteins identifed in each assayed condition suggested possible alterations in fve of the most important signalling networks of *B*. *cinerea*, with the strongest evidence of changes in the Small GTPase, MAPK cascade and Ca2+ mediated signal. The total number of signalling phosphoproteins exclusively identified or overexpressed under TCW and GLU was 25 (10%) and 10 (15.6%), respectively. The stress-activated mitogen protein kinase SAK1 (membranome; phosphoproteome; potentially associated to membrane) was identifed commonly in both conditions assayed, agreeing with our previous analysis in *B*. *cinerea*[11,](#page-11-8)[12,](#page-11-9) but it presented a diferent phosphorylation pattern depending on the carbon source. Sak1 is a well-known protein of the MAPK cascades that is phosphorylated in response to osmotic and oxidative stress by the MAPKK Bos5¹³. A protein similar to Bos5 (novel; peripheral/cytosolic) was identifed only under the TCW condition. Bos5 was not previously identifed in our analysis of *Botrytis cinerea*[11,](#page-11-8)[12,](#page-11-9) but was reported among the *B*. *cinerea* phosphoproteins by Davanture, M. *et al.*¹⁰. In addition, the presence of a type 2 C Ser/Thr phosphatase similar to PTC3 (membranome; potentially associated to membrane) under the TCW condition, but not under the GLU condition, may indicate a variation in the MAPK signalling pathway depending on the carbon source used. In relation to the Small GTPase signalling pathway under the TCW condition we found a larger number of phosphorylated membrane proteins related to this signalling network, including the p21-activated protein Cla4 (membranome; phosphoproteome; potentially associated to membrane), and two proteins that could be implicated in the regulation of Rho1: a protein similar to SAC7 (membranome; potentially associated to membrane); and a protein similar to RhoGEF (XP_001561237.2; phosphoproteome; potentially associated to membrane). In yeast, Rho1p is required for regulating the organization of the actin cytoskeleton, the activity of the (1,3)-β-D-glucan synthase and the cell wall integrity⁴⁶. In addition, a different ArfGEF (Guanine nucleotide exchange factor of the ADP ribosylation factor) was identifed under both conditions assayed; a protein similar to SYT1 (Bcsyt1) (novel; potentially associated to membrane) was identifed under TCW; and a putative guanyl-nucleotide exchange factor protein similar to the yeast SEC. 7(Bcsec. 7) (novel; potentially associated to membrane) was identified under GLU. SEC. 7 is implicated in yeast autophagosome phagophore formation⁴⁷. The last signalling system that presented more identifed phosphoproteins under TCW was the Ca2+ mediated signal pathway, which presented, among others, the Calcineurin-responsive zinc fnger transcription factor CRZ1 of Botrytis (BcCrz1)[48](#page-12-32) (novel; secreted protein), which is implicated in growth on poor media, hyphal morphology, stress response, cell wall and membrane integrity, and virulence¹³. BcCRZ1, the orthologue of the S. cerevisiae CRZ1 transcription factor, is known to be regulated by phosphorylation/dephosphorylation, being transported to the nucleus to perform its function when is dephosphorylated by calcineurin (CN) phos-phatase^{[48](#page-12-32)}. Additionally, there were two other phosphoproteins related to the $Ca2+/CN$ -signalling cascade under TCW: (i) BcCMK1 (calcium/calmodulin-dependent protein kinase) (membranome; phosphoproteome; potentially associated to membrane), the orthologue of the yeast CMK1, which is known to be phosphorylated and activated in response to Ca^{2+} counteracting the CN function⁴⁹; (ii) CND8 (novel; potentially associated to membrane), a calcineurin dependent (CND) gene that is upregulated by inhibition of CN⁵⁰. So, the presence of the phosphorylated form of BcCRZ1, BcCMK1 and CND8 indicates a negative regulation feedback of the Ca2⁺/CN-signalling cascade under the TCW condition. It has been described that the inhibition of CN and the deletion of BcCRZ1 resulted in decreased expression levels of botrydial biosynthetic genes^{48[,51](#page-12-35)}. Therefore, the negative regulation of BcCRZ1 and CN is consistent with the previously described inhibition of botrydial production under the TCW condition⁵².

All these diferences between the signal transduction phosphoproteins identifed under each condition suggest that these phosphoproteins play a vital role in the pathogeny through the process that activates them or are activated by them. A set of proteins similar to protein kinase or hypothetical kinases have been identifed under TCW condition, which have not been identified in our previous works^{[11,](#page-11-8)12}. These proteins could have important function in the adaptation of the fungus to the environmental conditions, constituting an interesting starting point for further studies.

Autophagy. Identified phosphorylated membrane proteins involved in autophagy have special relevance because it is known that induction, by nutrient starvation, of macroautophagy (hereafter autophagy) plays an important role in the formation of infection structures (appressorium) in plant pathogenic fungi, and therefore in determining the outcome of the penetration and pathogeny^{[29](#page-12-15)}. Under GLU, as a constitutive stage, as sole carbon source and according to the GO and/or KEGG annotation, 8 phosphorylated membrane-associated proteins (exclusive and overexpressed) related to autophagy were identified. Three of them (Atg3, Atg17 and Atg22) were described as ATG proteins, and the remaining five were identified as Avt3/4 (SLC36A), Kcs1, Vps41, ArfGEF and PDK1 (or PDPK1). The Atg3 protein is part of the ubiquitin-like Atg8 conjugation system that allows efficient vesicle expansion dur-ing autophagosome formation, and the Cvt pathway^{[29](#page-12-15),[53](#page-12-37)}. In filamentous fungi, Atg17 is specific and required for autophagy induction[54.](#page-12-38) In yeast, the Atg17-Atg31-Atg29 complex is present as a stable complex under both vegetative and starvation conditions, but phosphorylations of Atg29 and Atg31 generate an active Atg17-Atg31-Atg[29](#page-12-15) complex upon starvation conditions in autophagy induction^{29,55}. Atg22, Avt3 and Avt4, are vacuole permeases involved in the transport of amino acids produced in the final stage of autophagy[29,](#page-12-15)[56.](#page-13-0) Another vacuole protein identified in GLU and related to autophagy is Vps41, which is involved in docking and fusion of the double membrane vesicle with the vacuolar membrane during autophagy and the Cvt pathway^{[29](#page-12-15)[,57](#page-13-1)}. Thus the presence of the phosphorylated forms of Atg3, ATG17, Atg22, Avt3/4 and VPS 41 could be due to: (i) the constitutive activity of the Cvt pathway, because Atg3, Atg22, Avt3/4 and VPS 41 are required for both the Cvt pathway and autophagy, and the constitutive presence of the Atg17-Atg31-Atg29 complex; or (ii) the phosphorylation of Atg3, Atg22, Avt3/4 and the autophagy-specific protein Atg 17 could imply the inactivation of autophagy under nutrient-rich conditions (GLU)^{[29](#page-12-15),[55,](#page-12-39)56}. Finally, the other phosphoproteins identified under GLU: KCS1, the SEC. 7 superfamily member ArfGEF, and PDK1, are associated with the regulation of autophagy. KCS1 is an IP6K (inositol-hexakisphosphate 5-kinase) that converts Inositol hexakisphosphate (IP6) to 5-Diphosphoinositol pentakisphosphate (5-IP7), and in yeast has been shown to play a role in macroautophagy⁵⁸. The union of 5-IP7 to AKT1 causes the inhibition of this protein⁵⁹, activating autophagy by the inhibition of mTORC1 in the mTOR signalling pathway. The SEC. 7 superfamily ArfGEF is a GTP-exchange factor that catalyses the activation of the ADP-ribosylation factor family of small GTPases (ARFs). In the yeast *Saccharomyces cere*visiae three proteins containing SEC. 7 domains are required for autophagy by the activation of Arf1 and Arf2 GTPases^{47,60}. In mammalian cells, two ArfGEF members of the SEC. 7 superfamily are inhibited by phosphorylation⁶¹. The last identified phosphoprotein was mammal PDK1-like (phosphoinositide-dependent protein kinase). In some eukaryotes phosphorylated PDK1 is a positive regulator of the TOR pathway, activating mTORC1 and inhibiting autophagy^{[59,](#page-13-3)[62](#page-13-6)[,63](#page-13-7)}. These data suggest that, under GLU as sole carbon source, phosphorylated ArfGEF (inactive form) and PDK1 (active form), and possibly also an inactive phosphorylated Kcs1, are inhibiting autophagy in *B*. *cinerea*, indicating a possible role for these proteins in the regulation of autophagy, and therefore in the pathogeny of this fungus. Additionally, the glucose condition is the condition of maximum toxin production⁵², showing that the autophagy route and toxin synthesis seem to be controlled by different routes.

Cell death and UPR. The GO and KEGG annotation of our predicted proteins showed 3 proteins related to cell death exclusively identifed under TCW conditions. Tese proteins were predicted to be homologous of *Saccharomyces cerevisiae* proteins Lot6p, Bxi1p and Ire1p, that are known to be related to activation of apoptotic or autophagic cell death in *Saccharomyces cerevisiae*. Ire1p was the only one of these proteins that had not been previously identified in our previous proteomics analysis^{[11](#page-11-8),[12](#page-11-9)}. Phosphorylated Ire1p is the active form of the protein and is related to ER-stress-induced apoptotic cell death in response to an imbalance of unfolded proteins and chaperones (UPR) $64,65$ $64,65$. Lot6p activates apoptosis and reduces the activity of the proteasome in response to oxidative stress⁶⁶. The protein Bxi1p suppresses the activation of apoptotic programmed cell death by the UPR under ER stress and prevents Ire1p phosphorylation⁶⁷. In contrast, overexpression of Bxi1p in plants induces autophagic cell death⁶⁸. The UPR pathway has been reported to be necessary for virulence and during penetration of the plant^{[37,](#page-12-21)38}, as well as a possible central regulator of fungal pathogenesis⁶⁹. The presence of these phosphorylated proteins under the TCW condition, which simulates the frst contact of the fungus with the host plant, may indicate an activation of cell death in the fungus during penetration; this fnding agrees with previous reports that describe the induction of conidia cell death preceding the generation of appressorium during progression of penetration by the fungus[70](#page-13-14),[71](#page-13-15). In addition, it is known that in eukaryotic cells the balance between autophagy and apoptosis is essential for its survival and for pathogeny⁴⁵. These results highlight the significant role of phosphorylated Lot6p, Bxi1p, Ire1p in the regulation of pathogeny by activation of cell death.

Secondary metabolism: PDH. The central carbon metabolism is very important in fungal pathogenesis, so its regulation may be a critical element in the evolution of the infection process. The pyruvate dehydrogenase complex (PDC) converts pyruvate into acetyl-CoA and its regulation is based on the phosphorylation (inactive form) and dephosphorylation (active form) of its pyruvate dehydrogenase (PDH, E1) component. In previous reports it has been shown that, in *Fusarium graminearum*, the deletion of pyruvate dehydrogenase kinase (FgPDK1) resulted in an increase of PDH activity, reduction of the production of sesquiterpene deoxynivalenol (one of most frequently studied mycotoxins in *F. graminearum*), and elimination of pathogenicity^{[30,](#page-12-16)72}. Those results demonstrate that pyruvate metabolism plays a vital role in the regulation of the growth, toxin production, and pathogenicity in fungi. In our results phosphorylated PDH is overexpressed under the GLU but not under the TCW condition; taken together with our previous report, that shows that the botrydial biosynthesis pathway is inhibited when TCW is present as a sole carbon source, these fndings suggest that the production of this toxin is regulated by phosphorylation of PDH^{52} . In addition, the presence of the phosphorylated form of BcCRZ1 (inactive) is consistent with the idea that this transcription factor is not the sole regulator of the gene cluster responsible for the biosynthesis of botrydial^{[48](#page-12-32)}. Therefore, it is possible that at least two regulation pathways are independently inhibiting the production of botrydial under the TCW condition, which implicate the phosphorylation of BcCRZ1 and PDH.

Secondary metabolism: Catalase. It is known that hydrogen peroxide (H₂O₂) can affect the tissue of the plant during the infection process, but it also plays an important role in plant defence by inhibiting pathogen infection. So H2O2 may be destroyed rapidly by plant and fungal catalases (CATs) and peroxidases (POs). Schouten *et al*. (2002) demonstrated the early induction of H_2O_2 in tomato tissue infected by *B. cinerea* and the elimination of $H₂O₂$ by an extracellular CAT during penetration of the cell wall in the leaf. Subsequent reports have demonstrated that *B*. *cinerea* responds rapidly and efficiently to high levels of H_2O_2 added exogenously to the leaves^{73,74}.

Our results showed a predicted intracellular phosphorylated catalase exclusively identifed under the TCW condition, which may indicate an early response to some plant-based elicitor present in TCW. Such a response could represent the initial environmental conditions during penetration of the fungus that prepare the fungal organism for a subsequent increase of H_2O_2 levels.

Conclusion

In this study the phosphomembranome of *B*. *cinerea* has been evaluated in two diferent pathogenicity states, under the GLU (constitutive stage) and the TCW (virulence inductor) condition. Tis evaluation revealed a variation in phospho-membranome composition depending on the used carbon source. A total of 1112 proteins have been identifed of which 39.21% were not previously identifed in the phosphoproteome or the membranome analysis. Among the proteins exclusively identifed or overexpressed under each condition, 10.94% and 50.62% under GLU and TCW, respectively, were new proteins not identifed in our previous subproteomic analysis. All these new membrane-associated phosphoproteins identifed in the present analysis could provide to the research community a new set of targets to the design of novel strategies against *B*. *cinerea*. In addition, these results highlight the usefulness of using the analysis of a combination of two subproteomes to uncover new proteins that were covered up when the subproteome analysis were performed separately. It is interesting to highlight some process which have been showed to be diferentially regulated depending on the pathogenicity state in this work: ER stress response (UPR, ERAD, ubiquitin-dependent autophagy elimination of proteins aggregates), cell death; autophagy; pyruvate metabolism (PDH); and response to oxidative stress (catalase). It is known that UPR, ERAD, cell death and autophagy are related to ER stress and their cooperation is essential in the pathogeny/virulence of some fungi, but remains largely unexplored in *Botrytis*. Therefore, the regulation and connection between these biological processes in *B*. *cinerea* must be a key point in the understanding of its infection progress and how to stop it. Finally, a possible regulation of botrydial production by phosphorylation/desphosphorylation of PDH is presented as potential regulation pathway of this toxin, in addition to the regulation performed by CN and BcCRZ1.

Methods

Fungal strains and culture conditions. *B*. *cinerea* B05.10 strain (kindly provided by Dr. Paul Tudzunski of the University of Münster, Germany) was used for this study. Conidial stock suspensions were prepared and maintained as previously reported¹¹. Two different carbon sources were used: glucose (GLU) (Panreac, Spain) as the constitutive stage; and deproteinized tomato cell walls (TCW) as the virulence inductor, as previously described by Fernandez-Acero *et al*. [75.](#page-13-19) In brief, fasks of 500mL, each containing 250mL of minimal salt medium (MSM) (50 mM NH4Cl, 7.3 mM KH2PO4, 4.2 mM MgSO4, 6.7 mM KCl, 0.07 mM FeSO₄) supplemented with 1% of the carbon source assayed, were inoculated with *B. cinerea* conidia to a final concentration of 5×10^4 conidia/mL. Four independent replicas were assayed per culture condition. Replicates were incubated in parallel at 180 rpm at 22 °C under alternating 12-h light/dark cycles for 5 days. Afer 5 days, Phosstop phosphatase inhibitor cocktail (Roche, Basiela, Switzerland) was added to the culture according to the manufacturer's instructions. Then mycelia were collected by filtration in a 30-µm nylon filter (Sefar Nytal, Switzerland) and stored at -80 °C until use for membrane protein extraction.

Isolation of phosphorylated membrane proteins. The membrane protein fraction was isolated by the temperature-dependent partition method⁷⁶ with minor modifications^{[12](#page-11-9)}. For each biological replica, mycelium was ground into a fne powder with a mortar and pestle using liquid nitrogen. Membrane protein extraction was carried out from 100 mg of powdered mycelium using the "ReadyPrep Protein Extraction Kit (membrane I)" (Bio-Rad, USA) according to the manufacturer's specifcations. A total of fve membrane protein extractions were carried out for each biological sample. Afer protein extraction, only detergent-rich fractions were used. All the replicas from the same biological sample were pooled, precipitated with acetone and quantifed using the Qubit 2.0 Fluorometer system (Invitrogen, USA).

Each membrane fraction of 500 μg was dissolved in 6 M urea, 0.2 M NH₄HCO₃, then reduced with dithiothreitol (DTT) (10 mM, 37 °C, 60 min), and alkylated with iodoacetamide (20 mM, 25 °C, 30 min). Samples were diluted with up to 2M urea, digested overnight with Lys-C at 37 °C, and then diluted again 2-fold and digested overnight with trypsin at 37°C. Peptide mixtures were desalted using a MacroSpin C18 column (The Nest Group, Inc., Southborough, MA). The digested fractions obtained $(250\,\mu$ g) were enriched in phosphopeptides using titanium dioxide (TiO₂), as previously described¹¹. TiO₂ micro-columns were prepared in gel loading tips (0.5 mg) and were equilibrated with loading buffer (80% ACN in water $+6%$ TFA). Loading buffer was added to samples and the columns were washed with 80% ACN in water $+0.1\%$ TFA. Finally, phosphopeptides were eluted with 30 μL of elution bufer (5% NH3 in water) into a 1.5-mL tube containing 30 μL of 20% of formic acid in water. Crude phosphopeptide mixtures were desalted using a MacroSpin C18 column (The Nest Group, Inc., Southborough, MA).

Analysis and identifcation of proteins by LC‐MS/MS. 45% of each enriched phosphopeptide mixture was analysed using a LTQOrbitrap Velos Pro mass spectrometer (Thermo Fisher Scientific, San Jose, CA) coupled to a nano-LC (Proxeon, Odense, Denmark) equipped with a reversed-phase chromatography 25cm column with an inner diameter of 75 µm, packed with 1.9 µm C18 particles (Nikkyo Technos, Japan). A gradient of 3 to 35% acetonitrile in 360 min with 0.1% formic acid was used, at a flow rate of 250 nL/min. The Orbitrap Velos was operated in positive ion mode with nanospray voltage set at 2.2 kV and source temperature at 325 °C. The instrument was externally calibrated using Ultramark 1621 for the FT mass analyzer. An internal calibration was performed using the background polysiloxane ion signal at m/z 445.120025 as the calibrant. The instrument was operated in data-dependent mode. In all experiments, full MS scans were acquired over a mass range of m/z 350–2000 with detection in the Orbitrap mass analyzer at a resolution setting of 60,000. For each MS scan, the 20 most intense ions with multiple charged ions above a threshold ion count of 5000 were selected for fragmentation using collision-induced dissociation with multistage activation (activation of neutral losses of 98, 65.4, 49, 32.7) at a normalized collision energy of 35% in the LTQ linear ion trap. All data were acquired with Xcalibur v2.2 (Termo Fisher Scientifc, United States). Protein identifcation was performed with the Proteome Discoverer software suite v.1.4.0.288 (Thermo Fisher Scientific, United States) using MASCOT v2.3 (Matrix Science, United Kingdom) as search engine and a *B*. *cinerea* NCBI database including the most common contaminants (45755 entries; strains B05.10, T4 and BcDW1). Carbamidomethylation for cysteines was set as the fxed modifcation, and acetylation in protein N-terminal, phosphorylation in Ser, Thr and Tyr, and oxidation of methionine were set as variable modifications. Peptide tolerance was set at 7 ppm in MS, and at 0.5 Da in MS/MS mode. The maximum number of missed cleavages was set at 3. Peptides were fltered based on a false discovery rate (FDR) lower than 1%. Phosphosite location was assigned by the PhosphoRS algorithm. Areas under the curve for the identifed phosphopeptide were obtained with the "Precursor Ions Area Detector" algorithm from the Proteome Discoverer sofware. A fold change value and a Student's t-test for the phosphopeptides that were in at least three replicates of the four biological replicates in the two conditions was calculated to pinpoint diferentially abundant phosphopeptides (p-value $<$ 0.01). Obtained p-values were adjusted for multiple testing with the q-value method of Storey⁷⁷. Only those phosphopeptides with differences in the GLU/TCW ratio (R) of <0.66 and >1.5, with p-value<0.01, and with q-value<0.02 were considered as true diferentially abundant, and retained for further analyses. For the presence/absence analysis, a protein was considered exclusive to one phenotypical condition if it was present in at least three of the four biological replicates, and was not detected in any replicate of the other condition. No statistical analysis was performed for proteins that exhibited an absence/presence pattern among conditions.

Bioinformatics analysis. To predict protein topology several diferent algorithms were used. TMHMM Server v.2.0⁷⁸ was used to predict the transmembrane domain. To detect N-terminal secretory signal peptides, proteins were submitted to the SignalP 4.1 server⁷⁹. Predictions of non-classical protein secretion were performed using predictions for mammalian sequences in the SecretomeP 2.0 server⁸⁰. The Pred-GPI⁸¹ and big-PI Fungal predictors⁸² were used for the glycosylphosphatidylinositol (GPI) anchored protein prediction. Lastly, to detect prenylated, myristoylated and palmitoylated proteins, the GPS-lipid 1.0 (medium threshold) was used⁸³. Additionally, the PrepS-Prediction of Protein Prenylation^{[84](#page-13-28)}, NMT-The Myr predictor (fungi spe-cific) and CSS-Palm 4.0 (medium threshold)^{[85](#page-13-29)} were used to predict prenylated, myristoylated and palmitoylated proteins sites, respectively. The STRING protein interaction database version 10.5 (<https://string-db.org/>)^{[40](#page-12-23)} was used to generate a protein interaction network of the phosphoproteins identified (medium confidence $_$ 0.4). The protein-protein network obtained was then imported into Cytoscape (version $3.6.1$)⁴¹ and the clustering algorithm MCODE (version 1.5.1) was performed to identify potential functional clusters⁴². We performed a blastp search of the FASTA fles containing our identifed protein sequences against a database constructed from the complete *Botrytis cinerea* (strain B05.10) ASM83294v1 proteome from the ENSEMBL release 39 (found here: ftp://ftp.ensemblgenomes.org/pub/fungi/release-39/fasta/botrytis_cinerea/pep/). The parameters for the blastp search were: e-value = 1e-3 and max_target_seqs = 1. GO annotations of the proteins selected from the database used were assigned to their related proteins. The GO classification of proteins identified according to their involvement in a biological process (BP) and molecular function (MF) was carried out using the Agbase web server⁸⁶. KEGG annotations of all proteins identified were run with BlastKOALA 2.1⁸⁷. The KEGG annotations obtained were mapped to the network datasets KEGG pathway, using two KEGG mapper v3.1 tools of the KEGG database^{[43](#page-12-27)}: Reconstruct Pathway and Search&Color Pathway. The last tool was used to search against all pathway maps in the "bfu organism (*B*. *cinerea*)" category.

Data Availability

The datasets generated during the present study are available in the PRIDE repository, [[https://www.ebi.ac.uk/](https://www.ebi.ac.uk/pride/archive/) [pride/archive/\]](https://www.ebi.ac.uk/pride/archive/) with the dataset identifer PXD010961.

References

- 1. Dean, R. *et al*. Te Top 10 fungal pathogens in molecular plant pathology. *Mol Plant Pathol* **13**, 414–430 (2012).
- 2. Elad, Y., Williamson, B., Tudzynski, P. & Delen, N. *Botrytis: Biology, Pathology and Control*. (Springer Netherlands, 2007).
- 3. Choquer, M. *et al*. Botrytis cinerea virulence factors: new insights into a necrotrophic and polyphageous pathogen. *FEMS Microbiol Lett* **277**, 1–10 (2007).
- 4. Fillinger, S. & Elad, Y. *Botrytis - the Fungus, the Pathogen and its Management in Agricultural Systems*. (Springer International Publishing, 2016).
- 5. Gorna, K., Perlikowski, D., Kosmala, A. & Stepien, L. Host extracts induce changes in the proteome of plant pathogen *Fusarium proliferatum*. *Fungal*. *Biol* **121**, 676–688 (2017).
- 6. Jefery Daim, L. D. *et al*. Comparative proteomic analysis of oil palm leaves infected with *Ganoderma boninense* revealed changes in proteins involved in photosynthesis, carbohydrate metabolism, and immunity and defense. *Electrophoresis* **36**, 1699–1710 (2015).
- 7. Ashwin, N. M. R. *et al*. Advances in proteomic technologies and their scope of application in understanding plant-pathogen interactions. *J Plant Biochem Biot* **26**, 371–386,<https://doi.org/10.1007/s13562-017-0402-1> (2017).
- 8. Jefery, C. J. Why study moonlighting proteins? *Front Genet* **6**, 211 (2015).
- 9. Liñeiro, E., Cantoral, J. M. & Fernández-Acero, F. J. In *Botrytis - the Fungus, the Pathogen and its Management in Agricultural Systems* (eds Sabine Fillinger & Yigal Elad) 315–333 (Springer International Publishing, 2016).
- 10. Davanture, M. *et al*. Phosphoproteome profles of the phytopathogenic fungi *Alternaria brassicicola* and *Botrytis cinerea* during exponential growth in axenic cultures. *Proteomics* **14**, 1639–1645 (2014).
- 11. Lineiro, E., Chiva, C., Cantoral, J. M., Sabido, E. & Fernandez-Acero, F. J. Phosphoproteome analysis of *B*. *cinerea* in response to diferent plant-based elicitors. *J Proteomics* **139**, 84–94 (2016).
- 12. Lineiro, E., Chiva, C., Cantoral, J. M., Sabido, E. & Fernandez-Acero, F. J. Modifcations of fungal membrane proteins profle under pathogenicity induction: A proteomic analysis of *Botrytis cinerea* membranome. *Proteomics* **16**, 2363–2376 (2016).
- 13. Schumacher, J. In *Botrytis - the Fungus, the Pathogen and its Management in Agricultural Systems* (eds Sabine Fillinger & Yigal Elad) 247-267 (Springer International Publishing, 2016).
- 14. Humphrey, S. J., James, D. E. & Mann, M. Protein Phosphorylation: A Major Switch Mechanism for Metabolic Regulation. *Trends Endocrinol Metab* **26**, 676–687 (2015).
- 15. Turra, D., Segorbe, D. & Di Pietro, A. Protein kinases in plant-pathogenic fungi: conserved regulators of infection. *Annu Rev Phytopathol* **52**, 267–288 (2014).
- 16. Whitelegge, J. P. Integral membrane proteins and bilayer proteomics. *Anal Chem* **85**, 2558–2568 (2013).
- 17. Vuckovic, D., Dagley, L. F., Purcell, A. W. & Emili, A. Membrane proteomics by high performance liquid chromatography-tandem mass spectrometry: Analytical approaches and challenges. *Proteomics* **13**, 404–423 (2013).
- 18. Urban, M. *et al*. Te Pathogen-Host Interactions database (PHI-base): additions and future developments. *Nucleic Acids Res* **43**, D645–655 (2015).
- 19. Zheng, L., Campbell, M., Murphy, J., Lam, S. & Xu, J. R. Te BMP1 gene is essential for pathogenicity in the gray mold fungus *Botrytis cinerea*. *Mol Plant Microbe Interact* **13**, 724–732 (2000).
- 20. Schumacher, J. et al. The cAMP-dependent signaling pathway and its role in conidial germination, growth, and virulence of the gray mold *Botrytis cinerea*. *Mol Plant Microbe Interact* **21**, 1443–1459 (2008).
- 21. Viaud, M. *et al*. A class III histidine kinase acts as a novel virulence factor in *Botrytis cinerea*. *Mol Plant Microbe Interact* **19**, 1042–1050 (2006).
- 22. Gourgues, M., Brunet-Simon, A., Lebrun, M. H. & Levis, C. The tetraspanin BcPls1 is required for appressorium-mediated penetration of *Botrytis cinerea* into host plant leaves. *Mol Microbiol* **51**, 619–629 (2004).
- 23. Schoonbeek, H., Del Sorbo, G. & De Waard, M. A. Te ABC transporter BcatrB afects the sensitivity of *Botrytis cinerea* to the phytoalexin resveratrol and the fungicide fenpiclonil. *Mol Plant Microbe Interact* **14**, 562–571 (2001).
- 24. Vizcaino, J. A. *et al*. ProteomeXchange provides globally coordinated proteomics data submission and dissemination. *Nat Biotechnol* **32**, 223–226 (2014).
- 25. Ghosh, D., Beavis, R. C. & Wilkins, J. A. Te identifcation and characterization of membranome components. *J Proteome Res* **7**, 1572–1583 (2008).
- 26. Lodish, H. *et al*. Molecular Cell Biology 4th edn, (2000).
- 27. Staats, M. & van Kan, J. A. Genome update of Botrytis cinerea strains B05.10 and T4. *Eukaryot Cell* **11**, 1413–1414 (2012).
- 28. Willger, S. D. *et al*. Analysis of the *Candida albicans* Phosphoproteome. *Eukaryot Cell* **14**, 474–485 (2015).
- 29. Liu, X. H. *et al*. Autophagy in plant pathogenic fungi. *Semin Cell Dev Biol* **57**, 128–137 (2016).
- 30. Gao, T., Chen, J. & Shi, Z. *Fusarium graminearum* pyruvate dehydrogenase kinase 1 (FgPDK1) Is Critical for Conidiation, Mycelium Growth, and Pathogenicity. *PLoS One* **11**, e0158077 (2016).
- 31. AP, G., J, H., A, D., A, V. & NL, G. Regulated Forms of Cell Death in Fungi. *Front Microbiol*. 8 (2017).
- 32. Lisjak, M., Teklic, T., Wilson, I. D., Whiteman, M. & Hancock, J. T. Hydrogen sulfde: environmental factor or signalling molecule? *Plant Cell Environ* **36**, 1607–1616 (2013).
- 33. Amich, J. *et al*. Exploration of Sulfur Assimilation of *Aspergillus fumigatus* Reveals Biosynthesis of Sulfur-Containing Amino Acids as a Virulence Determinant. *Infect Immun* **84**, 917–929 (2016).
- 34. Lu, K., Psakhye, I. & Jentsch, S. Autophagic clearance of polyQ proteins mediated by ubiquitin-Atg8 adaptors of the conserved CUET protein family. *Cell* **158**, 549–563, <https://doi.org/10.1016/j.cell.2014.05.048> (2014).
- 35. Friedlander, R., Jarosch, E., Urban, J., Volkwein, C. & Sommer, T. A regulatory link between ER-associated protein degradation and the unfolded-protein response. *Nature Cell Biology* **2**, 379,<https://doi.org/10.1038/35017001> (2000).
- 36. Hetz, C. Te unfolded protein response: controlling cell fate decisions under ER stress and beyond. *Nature reviews. Molecular cell biology* **13**, 89–102,<https://doi.org/10.1038/nrm3270>(2012).
- 37. Cheon, S. A. *et al*. Unique evolution of the UPR pathway with a novel bZIP transcription factor, Hxl1, for controlling pathogenicity of *Cryptococcus neoformans*. *PLoS Pathog* **7**, e1002177 (2011).
- 38. Joubert, A. *et al*. Impact of the unfolded protein response on the pathogenicity of the necrotrophic fungus *Alternaria brassicicola*. *Mol Microbiol* **79**, 1305–1324 (2011).
- 39. Richie, D. L. et al. The virulence of the opportunistic fungal pathogen *Aspergillus fumigatus* requires cooperation between the endoplasmic reticulum-associated degradation pathway (ERAD) and the unfolded protein response (UPR). *Virulence* **2**, 12–21, <https://doi.org/10.4161/viru.2.1.13345>(2011).
- 40. Szklarczyk, D. *et al*. Te STRING database in 2017: quality-controlled protein-protein association. *Nucleic Acids Res* **4**, D362–368, <https://doi.org/10.1093/nar/gkw937>(2017).
- 41. Shannon, P. *et al*. Cytoscape: a sofware environment for integrated models of biomolecular interaction networks. *Genome Res* **13**, 2498–2504 (2003).
- 42. Bader, G. D. & Hogue, C. W. An automated method for fnding molecular complexes in large protein interaction networks. *BMC Bioinformatics* **4**, 2 (2003).
- 43. Kanehisa, M., Sato, Y., Kawashima, M., Furumichi, M. & Tanabe, M. KEGG as a reference resource for gene and protein annotation. *Nucleic Acids Res* **44**, D457–462 (2016).
- 44. Ren, W. et al. The Autophagy Gene BcATG8 Regulates the Vegetative Differentiation and Pathogenicity of Botrytis cinerea. Appl *Environ Microbiol* **84** (2018).
- 45. Veloso, J. & van Kan, J. A. L. Many Shades of Grey in Botrytis-Host Plant Interactions. *Trends Plant Sci* **23**, 613–622 (2018).
- 46. Perez, P. & Rincon, S. A. Rho GTPases: regulation of cell polarity and growth in yeasts. *Te Biochemical journal* **426**, 243–253, <https://doi.org/10.1042/BJ20091823>(2010).
- 47. van der Vaart, A., Grifth, J. & Reggiori, F. Exit from the Golgi Is Required for the Expansion of the Autophagosomal Phagophore in Yeast *Saccharomyces cerevisiae*. *Mol Biol Cell* **21**, 2270–2284 (2010).
- 48. Schumacher, J., de Larrinoa, I. F. & Tudzynski, B. Calcineurin-responsive zinc fnger transcription factor CRZ1 of *Botrytis cinerea* is required for growth, development, and full virulence on bean plants. *Eukaryot Cell* **7**, 584–601,<https://doi.org/10.1128/EC.00426-07> (2008).
- 49. Cisneros-Barroso, E. *et al*. Negative feedback regulation of calcineurin-dependent Prz1 transcription factor by the CaMKK-CaMK1 axis in fssion yeast. *Nucleic Acids Res* **42**, 9573–9587,<https://doi.org/10.1093/nar/gku684>(2014).
- 50. Harren, K., Schumacher, J. & Tudzynski, B. Te Ca2+ /calcineurin-dependent signaling pathway in the gray mold *Botrytis cinerea*: the role of calcipressin in modulating calcineurin activity. *PLoS One* **7**, e41761,<https://doi.org/10.1371/journal.pone.0041761> (2012)
- 51. Viaud, M. *et al*. and calcineurin functions investigated by gene inactivation, cyclosporin A inhibition and cDNA arrays approaches in the phytopathogenic fungus *Botrytis cinerea*. *Mol Microbiol* **50**, 1451–1465, <https://doi.org/10.1046/j.1365-2958.2003.03798.x> (2003).
- 52. Lineiro, E. *et al*. Phenotypic Efects and Inhibition of Botrydial Biosynthesis Induced by Diferent Plant-Based Elicitors in *Botrytis cinerea*. *Curr Microbiol* **75**, 431–440 (2018).
- 53. Ren, W. *et al*. Ubiquitin-like activating enzymes BcAtg3 and BcAtg7 participate in development and pathogenesis of *Botrytis cinerea*. *Current genetics* **64**, 919–930,<https://doi.org/10.1007/s00294-018-0810-3>(2018).
- 54. Pollack, J. K. & Harris, S. D. & Marten, M. R. Autophagy in flamentous fungi. *Fungal Genet Biol* **46**, 1–8 (2009).
- 55. Feng, Y., He, D., Yao, Z. & Klionsky, D. J. Te machinery of macroautophagy. *Cell Res* **24**, 24–41 (2014).
- 56. Lynch-Day, M. A. & Klionsky, D. J. Te Cvt pathway as a model for selective autophagy. *FEBS Lett* **584**, 1359–1366 (2010).
- 57. Lin, X. *et al*. RILP interacts with HOPS complex via VPS41 subunit to regulate endocytic trafcking. *Sci Rep* **4**, 7282 (2014).
	- 58. Taylor, R. Jr., Chen, P. H., Chou, C. C., Patel, J. & Jin, S. V. KCS1 deletion in *Saccharomyces cerevisiae* leads to a defect in translocation of autophagic proteins and reduces autophagosome formation. *Autophagy* **8**, 1300–1311 (2012).
	- 59. Tota, S. G. & Bhandari, R. Te emerging roles of inositol pyrophosphates in eukaryotic cell physiology. *J Biosci* **40**, 593–605 (2015). 60. Jackson, C. L. & Casanova, J. E. Turning on ARF: the Sec. 7 family of guanine-nucleotide-exchange factors. *Trends Cell Biol* **10**, 60–67 (2000)
	- 61. Kuroda, F., Moss, J. & Vaughan, M. Regulation of brefeldin A-inhibited guanine nucleotide-exchange protein 1 (BIG1) and BIG2 activity via PKA and protein phosphatase 1gamma. *Proc Natl Acad Sci USA* **104**, 3201–3206 (2007).
- 62. McManus, S. *et al*. Autophagy and 3-Phosphoinositide-Dependent Kinase 1 (PDK1)-Related Kinome in Pagetic Osteoclasts. *J Bone Miner Res* **31**, 1334–1343 (2015).
- 63. Khan, M. S. & Hemalatha, S. Autophagy: Molecular Insight and Role in Plant Programmed Cell Death and Defense Mechanism. *Int Res J Biological Sci* **4**, 78–83 (2015).
- 64. Heimel, K. *et al*. Crosstalk between the unfolded protein response and pathways that regulate pathogenic development in *Ustilago maydis*. *Plant Cell* **25**, 4262–4277 (2013).
- 65. Tabas, I. & Ron, D. Integrating the mechanisms of apoptosis induced by endoplasmic reticulum stress. *Nat Cell Biol* **13**, 184–190 (2011)
- 66. Sollner, S., Durchschlag, M., Frohlich, K. U. & Macheroux, P. Te redox-sensing quinone reductase Lot6p acts as an inducer of yeast apoptosis. *FEMS Yeast Res* **9**, 885–891 (2009).
- 67. Cebulski, J., Malouin, J., Pinches, N., Cascio, V. & Austriaco, N. Yeast Bax inhibitor, Bxi1p, is an ER-localized protein that links the unfolded protein response and programmed cell death in *Saccharomyces cerevisiae*. *PLoS One* **6**, e20882 (2011).
- 68. Xu, G. *et al*. Plant Bax Inhibitor-1 interacts with ATG6 to regulate autophagy and programmed cell death. *Autophagy* **13**, 1161–1175 (2017).
- 69. Krishnan, K. & Askew, D. S. Endoplasmic reticulum stress and fungal pathogenesis. *Fungal biology reviews* **28**, 29–35, [https://doi.](https://doi.org/10.1016/j.fbr.2014.07.001) org/10.1016/j.fbr.2014.07.001 (2014).
- 70. Ryder, L. S. & Talbot, N. J. Regulation of appressorium development in pathogenic fungi. *Curr Opin Plant Biol* **26**, 8–13 (2015).
- 71. Goncalves, A. P., Heller, J., Daskalov, A., Videira, A. & Glass, N. L. Regulated Forms of Cell Death in Fungi. *Front Microbiol* **8**, 1837 (2017).
- 72. Ene, I. V., Brunke, S., Brown, A. J. & Hube, B. Metabolism in fungal pathogenesis. *Cold Spring Harb Perspect Med* **4**, a019695 (2014). 73. Patykowski, J. Role of hydrogen peroxide and apoplastic peroxidase in tomato-*Botrytis cinerea* interaction. *Acta Physiol Plant* **28**, 589–598,<https://doi.org/10.1007/s11738-006-0054-6> (2006).
- 74. Schouten, A. *et al*. Functional analysis of an extracellular catalase of *Botrytis cinerea*. *Mol Plant Pathol* **3**, 227–238 (2002).
- 75. Fernandez-Acero, F. J. *et al*. 2-DE proteomic approach to the *Botrytis cinerea* secretome induced with diferent carbon sources and plant-based elicitors. *Proteomics* **10**, 2270–2280 (2010).
- 76. Zhao, Y., Zhang, W., Kho, Y. & Zhao, Y. Proteomic analysis of integral plasma membrane proteins. *Anal Chem* **76**, 1817–1823 (2004).
- 77. Storey, J. D. Te positive false discovery rate: a Bayesian interpretation and the q -value. *Ann Statist* **31**, 2013–2035, [https://doi.](https://doi.org/10.1214/aos/1074290335) [org/10.1214/aos/1074290335](https://doi.org/10.1214/aos/1074290335) (2003).
- 78. Krogh, A., Larsson, B., von Heijne, G. & Sonnhammer, E. L. Predicting transmembrane protein topology with a hidden Markov model: application to complete genomes. *J Mol Biol* **305**, 567–580 (2001).
- 79. Petersen, T. N., Brunak, S., von Heijne, G. & Nielsen, H. SignalP 4.0: discriminating signal peptides from transmembrane regions. *Nat Methods* **8**, 785–786 (2011).
- 80. Bendtsen, J. D., Jensen, L. J., Blom, N., Von Heijne, G. & Brunak, S. Feature-based prediction of non-classical and leaderless protein secretion. *Protein Eng Des Sel* **17**, 349–356 (2004).
- 81. Pierleoni, A., Martelli, P. L. & Casadio, R. PredGPI: a GPI-anchor predictor. *BMC Bioinformatics* **9**, 392 (2008).
- 82. Eisenhaber, B., Schneider, G., Wildpaner, M. & Eisenhaber, F. A sensitive predictor for potential GPI lipid modifcation sites in fungal protein sequences and its application to genome-wide studies for *Aspergillus nidulans*, *Candida albicans*, *Neurospora crassa*, *Saccharomyces cerevisiae* and *Schizosaccharomyces pombe*. *J Mol Biol* **337**, 243–253 (2004).
- 83. Xie, Y. *et al*. GPS-Lipid: a robust tool for the prediction of multiple lipid modifcation sites. *Sci Rep* **6**, 28249 (2014).
- 84. Maurer-Stroh, S. & Eisenhaber, F. Refnement and prediction of protein prenylation motifs. *Genome Biol* **6**, R55–R55.15 (2005).
- 85. Ren, J. *et al*. CSS-Palm 2.0: an updated sofware for palmitoylation sites prediction. *Protein Eng Des Sel* **21**, 639–644 (2008).
- 86. McCarthy, F. M. *et al*. AgBase: supporting functional modeling in agricultural organisms. *Nucleic Acids Res* **39**, D497–506 (2011).
- 87. Kanehisa, M., Sato, Y. & Morishima, K. BlastKOALA and GhostKOALA: KEGG Tools for Functional Characterization of Genome and Metagenome Sequences. *J Mol Biol* **428**, 726–731 (2016).

Acknowledgements

Tis research has been fnanced by the Spanish Government under project AGL2015-65684-C2-2-R supported by the Spanish Ministry of Economy and Competitiveness. Eva Liñeiro was supported by an FPI grant from the University of Cádiz (2010-152). The CRG/UPF Proteomics Unit is a member of the ProteoRed PRB3 consortium which is supported by grant PT17/0019 of the PE $I+D+i 2013-2016$ from the Instituto de Salud Carlos III (ISCIII) and ERDF. We acknowledge support from the Spanish Ministry of Economy and Competitiveness, "Centro de Excelencia Severo Ochoa 2013–2017", SEV-2012-0208, and "Secretaria d'Universitats i Recerca del Departament d'Economia i Coneixement de la Generalitat de Catalunya" (2017SGR595).

Author Contributions

F.J.F.-A. conceived and directed the experiments; E.L. conducted the experiment(s); E.S. and C.C. performed the MS/MS analyses. A.E.-N., C.F., R.C. and F.A. developed the bioinformatics approach for the analysis of the results; A.E.-N., F.J.F.-A., B.B.-U. and J.M.C.F. wrote the paper. All authors reviewed the manuscript.

Additional Information

Supplementary information accompanies this paper at [https://doi.org/10.1038/s41598-019-46270-0.](https://doi.org/10.1038/s41598-019-46270-0)

Competing Interests: The authors declare no competing interests.

Publisher's note: Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional afliations.

Co O Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit [http://creativecommons.org/licenses/by/4.0/.](http://creativecommons.org/licenses/by/4.0/)

 $© The Author(s) 2019$