



## A Proteome-Level Investigation Into *Plasmodiophora brassicae* Resistance in *Brassica napus* Canola

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Clubroot of Brassicaceae, an economically important soil borne disease, is caused by Plasmodiophora brassicae Woronin, an obligate, biotrophic protist. This disease poses a serious threat to canola and related crops in Canada and around the globe causing significant losses. The pathogen is continuously evolving and new pathotypes are emerging, which necessitates the development of novel resistant canola cultivars to manage the disease. Proteins play a crucial role in many biological functions and the identification of differentially abundant proteins (DAP) using proteomics is a suitable approach to understand plant-pathogen interactions to assist in the development of gene specific markers for developing clubroot resistant (CR) cultivars. In this study, P. brassicae pathotype 3 (P3H) was used to challenge CR and clubroot susceptible (CS) canola lines. Root samples were collected at three distinct stages of pathogenesis, 7-, 14-, and 21-days post inoculation (DPI), protein samples were isolated, digested with trypsin and subjected to liquid chromatography with tandem mass spectrometry (LC-MS/MS) analysis. A total of 937 proteins demonstrated a significant (q-value < 0.05) change in abundance in at least in one of the time points when compared between control and inoculated CR-parent, CR-progeny, CS-parent, CS-progeny and 784 proteins were significantly (q < 0.05) changed in abundance in at least in one of the time points when compared between the inoculated- CR and CS root proteomes of parent and progeny across the three time points tested. Functional annotation of differentially abundant proteins (DAPs) revealed several proteins related to calcium dependent signaling pathways. In addition, proteins related to reactive oxygen species (ROS) biochemistry, dehydrins, lignin, thaumatin, and phytohormones were identified. Among the DAPs, 73 putative proteins orthologous to CR proteins and quantitative trait loci (QTL) associated with eight CR loci in different chromosomes including chromosomes A3 and A8 were identified. Proteins including BnaA02T0335400WE, BnaA03T0374600WE, BnaA03T0262200WE, and BnaA03T0464700WE are orthologous to identified CR loci with possible roles in mediating clubroot responses. In conclusion, these results have contributed to an improved understanding of the mechanisms involved in mediating response to P. brassicae in canola at the protein level.

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## INTRODUCTION

Canola (Brassica napus L.) is a widely cultivated, cruciferous oilseed crop. Globally, 68.87 million metric tons of canola was produced in 2020, making it an important agricultural commodity (Shahbandeh, 2021). Since its development  $\sim$ 50 years ago, Canada has led the world in canola production (Canola Council of Canada, 2021 - Industry overview), generating \$29.9 billion annually in economic activities (Canola Council of Canada, 2021 - Industry Overview). For over a century, Brassica crop yield has been negatively affected by P. brassicae Woronin, which causes clubroot disease (Cook and Schwartz, 1930; Buczacki, 1983). This disease has impacted > 56 countries in all of Asia, Europe, Americas, some parts of Oceania and Africa, leading to yield losses of ~15% globally (Dixon, 2009; Czubatka-Bieńkowska et al., 2020). Development of CR canola cultivars has been successful in combating the disease but the evolution of new pathotypes of P. brassicae that can break down the resistance continues to be a major problem (Strelkov et al., 2018).

*Plasmodiophora brassicae*, the soil-borne, obligate, biotrophic protist pathogen produces resting spores, which can survive in soil for many years and, when the conditions are favorable, serve as the inoculum to infect susceptible plants resulting in the death of ~50% of the infected plants (Gibbs, 1931; Wallenhammar, 1996; Howard et al., 2010; Hwang et al., 2012). Clubroot development in Brassica spp. is favored by warm temperatures (20-25°C) and wet soils with relatively lower soil pH (<6.5; Dixon, 2009; Gossen et al., 2013). Disease progression in roots is characterized by a primary plasmodia stage marked by the infection of root hairs, followed by secondary plasmodia phase indicated by cortical cell infection, which escalates into increased and irregular vascular cambium activity, reduction in xylem tissue, increase in parenchyma cells, and eventually the deformity of roots (Kageyama and Asano, 2009; Hwang et al., 2012; Malinowski et al., 2012). Pathogen progression in root tissues trigger phytohormone (auxin and cytokinin) imbalance leading to hypertrophy and hyperplasia, which results in the formation of clubs in the roots (Siemens et al., 2006; Ludwig-Müller et al., 2009, 2017; Jahn et al., 2013). Once the pathogen advances to the secondary plasmodia phase, significant damage on the root tissues occur and infection sites become physiological sinks, leading to the disruption of water and mineral conduction, resulting in stunting, wilting and eventual death of plants (Keen and Williams, 1969; Walerowski et al., 2018).

In different parts of the world, clubroot management programs involve the application of soil amendments (e.g., lime) or chemical fungicides, crop rotation, and field equipment sanitization; however, many of these conventional methods are either too expensive or ineffective in large farm settings (Donald and Porter, 2009). As alluded to earlier, recently, growers have depended heavily on CR cultivars; however, the emergence of novel *P. brassicae* pathotypes and their rapid spread have posed serious threats to the management of clubroot disease in canola (Strelkov et al., 2018; Hollman et al., 2021). Numerous studies have been conducted to

understand the biology and races of *P. brassicae* (Williams, 1966; Kageyama and Asano, 2009), and to identify quantitative trait loci (QTL) related to the disease resistance in *Brassica* species (Nagaoka et al., 2010; Hasan et al., 2021). To date approximately 24 QTL and over 400 putative clubroot resistance genes have been reported and mapped to the A and C genomes of different *Brassica* spp. (Nagaoka et al., 2010; Dakouri et al., 2021; Hasan et al., 2021). Among these, multiple QTL were mapped to each of chromosomes A3 and A8 in *B. rapa*. Although many of the putative clubroot resistance related genes are yet to be confirmed, their putative physical location has been reported by different researchers (Yu et al., 2017; Hasan et al., 2021).

Advances in next generation sequencing (NGS) and other "omics"-based approaches have contributed to the identification of many putative genes involved in mediating responses in Brassica spp. to the clubroot pathogen. For instance, RNA sequencing studies in B. oleracea, B. rapa, B. napus, Arabidopsis thaliana have implicated a role for hormonal signaling in response to the pathogen and profiled several putative clubroot resistance related gene transcripts and lncRNAs (Chen et al., 2016; Zhang et al., 2016; Zhao et al., 2017; Summanwar et al., 2019, 2020, 2021; Galindo-González et al., 2020). Similarly, there are a few reports of metabolomicsbased approaches in Brassica spp. that have documented metabolome-level changes in response to the pathogen including a potential role for gluconasturtiin in conferring resistance against the pathogen (Pedras et al., 2008; Wagner et al., 2012, 2019). Proteome-level changes in canola in response to the clubroot pathogen have also been described, which included characterization of root proteomes in response to infection in susceptible canola (Cao et al., 2008). Ji et al. (2018) reported DAPs during early stages of infection and suggested a role for phytohormone regulation during the host-pathogen interaction. Song et al. (2016) characterized the proteome-level changes in B. rapa genotypes with and without clubroot resistance gene (Rcr1) and reported 527 DAPs, which could be potentially involved in mediating clubroot resistance. Su et al. (2018) and Lan et al. (2019) have also identified DAPs during the secondary stage of infection in CR and CS genotypes of B. rapa. Their studies suggested that the proteins involved in brassinosteroids (BR) and cytokinin (CK) pathways were potentially involved in the regulation of clubroot development. Due to the important role played by proteins in mediating various cellular processes, including hostpathogen interactions, the identification and characterization of DAPs may be a viable approach to identify the proteins potentially involved in mediating resistance to the pathogen (Quirino et al., 2010; Song et al., 2016; Ji et al., 2018; Lan et al., 2019).

In the current study, a comprehensive proteomics analysis at three different stages of infection in CR and CS canola representing CR/CS parent and CR/CS doubled-haploid (DH) progeny lines was conducted and DAPs potentially involved in mediating clubroot resistance were identified. Our results have significance in not only further understanding the molecular mechanisms of clubroot resistance but also in the potential development of protein-specific markers to assist in the selection of CR lines.

## MATERIALS AND METHODS

## **Plant Material**

In this study, a set of doubled haploid (DH) lines and their parents were used; detailed information on these lines are available in Hasan and Rahman (2016). Briefly, the DH lines were derived from  $F_1$  plants of Rutabaga Brookfield-9005 P1 x A07-29NI and A07-29NI x Rutabaga Brookfield-9005 P1 crosses. The parent Rutabaga Brookefield-9005 is a CR inbred line developed from rutabaga (*Brassica napus* var. *napobrassica*) cv. Brookefield-9005 through self-pollination (Hasan and Rahman, 2016) and showed complete resistance to Canadian *P. brassicae* pathotypes 2, 3, 5, 6, and 8. The CS parent used in this study was A07-29NI, which is an open pollinated spring *B. napus* canola cultivar grown in Canada as UA AlfaGold (Rahman, 2016).

## Preparation of *Plasmodiophora brassicae* Inoculum, Infection of Plants and Disease Assessment

A single spore isolate of *P. brassicae*, SACAN-ss1 belonging to pathotype 3 (P3H) was used to challenge the seedlings. Resting spores of the pathogen were isolated essentially as described by Williams (1966). Briefly, infected root tissues with visible clubroot galls (rated disease score of 3/3) were collected, washed, and stored at  $-80^{\circ}$ Celsius until use. Frozen gall tissues (38 g) were thawed and homogenized in sterile deionized water (400 ml) using a mechanical blender. Homogenized tissue was filtered through eightfold layers of cheesecloth (American Fiber & Finishing Inc., Albemarle, NC, United States) and the density of the resting spores in the filtrate was determined using a hemocytometer (VWR, Mississauga, ON, Canada). Final resting spore density was adjusted to  $1 \times 10^7$  spores/ml with sterilized deionized water (600 ml).

The resistance and susceptibility of all 11 CR and 9 CS DH lines used in this study was confirmed prior to experimentation. All experimental plants were grown in a greenhouse maintaining 400 µmol/m<sup>2</sup> of light intensity at 22/15°C (light/dark) with a 16/8 h photoperiod in cells filled with Sunshine Professional Growing Mix (Sunshine Horticulture, Bellevue, NE, United States). When seedlings were at 2-3 true leaf stage they were directly injected, close to the root, with the spore suspension (1 ml of 1  $\times$   $10^7$ spores/ml). Control plants were treated with sterile water and maintained in separate trays under identical growth conditions. The soil was saturated with water for the first week following inoculation and subsequently watering was performed as required to keep the soil moist. Mixed fertilizer, Nitrogen-Phosphorous-Potassium (20-20-20) was applied once in a week.

Disease screening was performed at 45 DPI as follows: roots were washed thoroughly with tap water, and symptoms were rated visually on a 0-3 scale as described by Kuginuki et al.

(1999), where 0 = no visible symptom of clubroot galls, 1 = a small gall mainly on lateral roots, 2 = small galls on the main and lateral roots, and 3 = severe clubbing or the entire root system was deformed. The disease index (DI) was calculated as follows: DI =  $[(n_1 + 2n_2 + ...3n_n)]/(N_Tx3)] \times 100$ , where  $n_1$ ,  $n_2$  ...  $n_n$  indicate the individuals from a line with the level of disease severity (0–3) and  $N_T$  indicates the total number of individuals in the line (Li et al., 2016).

## Confirmation of Infection Using Microscopy

To assess the changes in the root tissues, histology and scanning electron microscopy (SEM) of the infected and uninfected tissues at 7–, 14–, and 21– DPI were performed. For histology, samples were prepared as described by Summanwar et al. (2019). Stained tissue sections were dried at 37°C and visualized with a Leica DMRXA light microscope (Meyer Instruments, Houston, TX, United States). Images were acquired with a QI Click digital camera and processed using Q Capture Pro 7 software (Q Imaging, Surrey, BC, Canada).

For SEM, all steps until tissue sectioning were followed according to Summanwar et al. (2019). Tissue sections were collected on SEM pin stubs, air dried, and dewaxed in toluene twice for 5 min and then washed with 100% ethanol twice for 2 min. Samples were further air dried and subjected to sputter coating with palladium (Pd) for 120 seconds using Hummer 6.2 Sputter Coater (Anatech Ltd, Battle Creek, MI, United States). Once coated, tissues were subjected to high resolution electron imaging using ZEISS EVO 10 Scanning Electron Microscope (SEM) (Carl Zeiss Canada Ltd, Toronto, ON, Canada) (White Plains, NY, United States).

## Sample Collection for Protein Extraction

Tissue samples (72 in total) comprised of four genotypes [CRparent (10 individuals pooled per biological replicate), CS-parent (10 individuals pooled per biological replicate), CR-progeny-bulk (20 individuals pooled per biological replicate representing 10 lines), and CS-progeny-bulk (16 individuals pooled per biological replicate representing 8 lines)]. Root samples were collected at three different times points, 7–, 14–, and 21-DPI as per our previous studies (Summanwar et al., 2019). Three independent experiments comprised of three biological replicates in each experiment and two treatments (inoculated and control) made up the 72 samples (4 genotypes  $\times$  3 independent experiments  $\times$  3 biological replicates  $\times$  2 treatments = 72).

# Extraction of Protein for Proteome Analysis

Frozen root tissue samples described above were homogenized in liquid nitrogen using a mortar and pestle. Homogenized tissue (200 mg) was used for protein extraction using a 1:2 (w/v) ratio of 50 mM HEPES-KOH pH 8.0. 50 mM NaCl, and 4% SDS. Samples were solubilized by vortexing and incubated at 95°C with agitation at 600 rpm for 15 min. After cooling to room temperature, samples were centrifuged at 20,000 g for 5 min and the supernatants were transferred to fresh Eppendorf microcentrifuge tubes (2.0 ml). Protein estimation was then performed using a bicinchonic acid (BCA) assay (23225; ThermoScientific, Mississauga, ON, Canada) and samples were normalized to 100  $\mu$ g each in a total volume of 250  $\mu$ L. Samples were then reduced with 10 mM dithiothreitol (DTT) at 95°C for 5 min, cooled to room temperature and alkylated with 30 mM iodoacetamide for 30 min in the dark and quenched with 10 mM DTT.

Proteins were digested using a magnetic bead based protocol using a 1:1 mix of hydrophilic and hydrophobic Sera-Mag SpeedBead Carboxylate-modified magnetic particles (45152105050250 and 65152105050250; GE Life Sciences, Boston, MA, United States) (Leutert et al., 2019). Tryptic hydrolysis of protein samples was performed overnight at 37°C using sequencing-grade trypsin (V5113; Promega, Madison, WI, United States). Digested peptides were quantified using a Nanodrop and acidified by adding formic acid (FA) to a final concentration of 5%. The peptide samples were then dried by vacuum centrifugation and dissolved in 3% acetonitrile (ACN) containing 0.1% trifluoroaceticacid (TFA) and desalted using ZipTip C18 pipette tips (ZTC18S960; Millipore, Oakville, ON, Canada). Desalting was performed by pre-equilibrating each tip sequentially with 100% ACN, 60% ACN/0.1% TFA, and 3% ACN/0.1% TFA. Samples were adsorbed by pipetting ten times followed by three washes with 3% ACN/0.1% TFA and eluted with 10 µL of 60% ACN/0.1% TFA five times. Desalted peptides were dried and dissolved in 3.0% ACN/0.1% FA prior to liquid chromatography with tandem mass spectrometry (LC-MS/MS) analysis.

## Nanoflow Liquid Chromatography With tandem Mass Spectrometry Analysis

Peptides were analyzed using a Fusion Lumos Tribrid Orbitrap mass-spectrometer (ThermoFisher Scientific) using a BoxCarDIA data acquisition scheme as described previously (Mehta et al., 2022). Briefly, the dissolved peptides were injected using an Easy-nLC 120 system (LC140; ThermoFisher Scientific) and separated using a 50 cm Easy-Spray PepMap C18 column (ES803A; ThermoFisher Scientific). The column was equilibrated with 100% solvent A (0.1% FA) and the samples were eluted using a segmented solvent B gradient (80% ACN/0.1% FA) from 4 to 41% B (0–65 min). MS<sup>1</sup> acquisition was performed using two multiplexed tSIM scans of 10 BoxCar windows each spanning a range of 350-1400 m/z. Detection was performed at 120,000 resolution and normalized AGC targets of 100% per isolation window. MS<sup>2</sup> acquisition was performed using twenty-eight 38.5 m/z windows with an overlap of 1 m/z and a minimum m/z of 200. Resolution was set to 30,000 using a dynamic maximum injection time and the minimum desired points across each peak was set to 6.

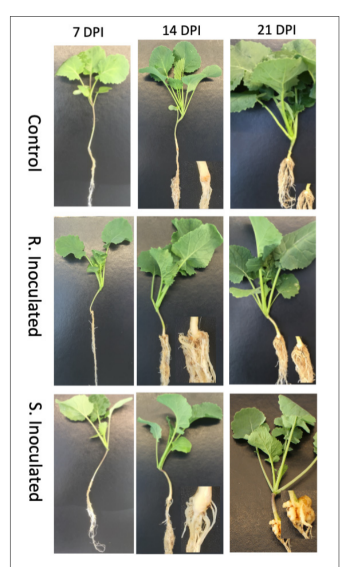
# Data Processing and Bioinformatics Analysis

BoxCarDIA raw files were processed using Spectronaut 14 (Biognosys AG, Wagistrasse, Schlieren Switzerland) under a directDIA analysis mode and default search parameters without

*N*-acetyl variable modification as described in Mehta et al. (2022). Spectra were searched using a published *B. napus* Westar proteome (Song et al., 2020). Trypsin specificity was set to two missed cleavages and a protein and PSM false discovery rate of 1% each. Data filtering criteria was set to *q*-value  $\leq$  0.01 and global normalization with MS2-level quantification was performed. Comparative analysis between samples was performed using Spectronaut with a significance threshold of FDR-corrected *p*-value (Dunn, 1961) to identify differentially abundant proteins.

# Functional Classification and Annotation of Differentially Abundant Proteins

Proteome analysis was carried out following standard method similar to published works (Kanehisa et al., 2016;



**FIGURE 1** Clubroot gall development following inoculation with *P. brassicae* pathotype 3. Control at 7-, 14-, and 21-DPI, Clubroot resistant (CR) inoculated line at 7-, 14-, and 21-DPI Clubroot susceptible (CS) inoculated 7-, 14-, and 21-DPI.

Mistry et al., 2021) predicted proteins were annotated using the gene descriptions from KEGG, Pfam, and Uniprot databases to gain additional insights into their biological roles. TAIR hit IDs corresponding to proteins with significant differentially expressed values (*q*-value < 0.05, after Dunn, 1961) were taken for gene ontology (GO) enrichment analysis using AgriGo (Du et al., 2010; Tian et al., 2017). *Arabidopsis* gene model (TAIR9) was used in the background and the following parameters were applied to run the analysis – statistical test method (Hypergeometric), multi-test adjustment statistical method [Yekutieli (FDR under dependency)], significance level (0.05), minimum number of mapping entries (5), and gene ontology type (complete GO).

## RESULTS

## Disease Progression Following Plasmodiophora brassicae Challenge

We compared the changes occurring in the roots of CR and CS lines following the pathogen infection at three different time points using light and scanning and electron microscopy (SEM). The results obtained at each time points are described below.

#### 7 Days Post Inoculation

At this stage of infection, the morphology of plant roots and shoots were similar in both control and inoculated groups, in both CR and CS lines with roots being free of galls (**Figure 1**). Microscopic examination of root tissues indicated that *P. brassicae* was present in the early developmental stage in the vegetative plasmodia phase, colonized around the root hairs, penetrated the cell wall barrier, and established in the root system (**Figures 2**, **3**). Infected cells in both CR and CS lines appeared to be similar in histology sections and appeared as a mass of pink matrices at the infected sites (**Figure 2**) and under SEM, cells appeared as a tight mass of pathogen cells, which could be distinguished from the uninfected cells (**Figure 3**).

#### 14 Days Post Inoculation

At 14 DPI, we observed that the root tissues and the cortical region of roots was also penetrated by the pathogen forming mature secondary plasmodia, and larger areas of the root tissue were found to be infected (**Figures 2**, **3**). Histology sections showed that the disease progression was apparent in the CS lines, with a condensed mass of cells arranged at the center (**Figure 2**). However, in the CR lines, the pink matrix had condensed and loosened giving a granular pink dot like appearance indicating a slower progression of the pathogen (**Figure 2**). Similarly, under SEM, loosened granular materials were observed in the infected cells in the CR and CS line (**Figure 3**).

#### 21 Days Post Inoculation

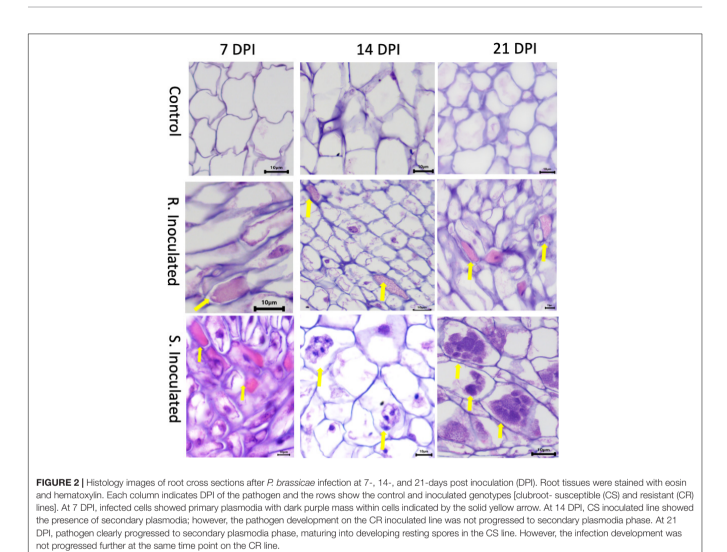
At 21 DPI, not only the clubroot galls were clearly visible on the CS roots (**Figure 1**), but also the plasmodia, sporangia, and resting spores of the pathogen could also be clearly visualized in under light microscopy as well as SEM, indicating disease progression and infection of the root cortical region (**Figures 2**, **3**). The color of pathogen cells changed to dark purple with multiple clusters of maturing secondary plasmodium (**Figure 2**) and SEM micrographs indicated maturing secondary plasmodium in a cluster of pathogen cells within the infected host tissues (**Figure 3**). The whole plant roots appeared swollen with clubs and deformed the root architecture (**Figure 1**). However, in the CR roots, disease progression was slow and appearance of infected cells was similar at 14 and 21 DPI, showing pink matrix in the cells under histology sections and loosely packed granular pathogen in the infected cells under SEM.

#### 45 Days Post Inoculation

At 45 DPI, 98% of the CR lines did not develop any galls and there were clear visual differences between the roots of CR and CS plants (**Supplementary Figure 1**). All inoculated CS and CR lines had a disease severity index (DSI) rating > 98% and <5%, respectively (**Supplementary Table 1**). We selected 7-, 14-, and 21-DPI for our proteome studies since these stages appear to be important for disease progression as evidenced by our microscopic studies and correspond with our previous transcriptome-level studies (Summanwar et al., 2019).

# Identification of Differentially Abundant Proteins

Protein profiles from control and inoculated CR and CS root samples, from both progeny and parent lines were analyzed. We detected an average of 3281 proteins across all the genotypes, treatments, and timepoints, with a low of 2296 proteins in the 14 DPI control CR-parent lines and a high of 3689 proteins at 21 DPI control CS-progeny lines (Table 1). Among these proteins, 937 were significantly (q < 0.05) differentially abundant in at least in one of the time points when compared between control and inoculated root proteomes of CR-parent, CR-progeny, CS-parent, CS-progeny across the three time points (Supplementary Table 2). Similarly, 784 proteins were significantly (q < 0.05) differentially abundant in at least in one of the time points when compared between the inoculated-CR and CS root proteomes of parent and progeny across the three time points (Supplementary Table 3). Finally, while there were 684 differentially abundant proteins were common to both CR and CS control samples, 99 proteins demonstrated significant (q < 0.05) differential abundance only in the inoculated CR and CS samples, including both progeny and parent lines (Supplementary Table 4). In the pool of unique proteins, there were 36 proteins that showed similar pattern of accumulation in parent and progeny, of them 20 proteins were increased in accumulation and 16 were decreased as the infection progressed to 21 DPI (Supplementary Table 4). In this pool, nine putative proteins were potentially involved in plant and pathogen interaction, proteins with increased accumulation included: orthologs such as fascilin-like (BnaA02T0010200WE), annexin (BnaA06T0293900WE), glutathione S-transferase (Bna A03T0164200WE and BnaA03T0240300WE); and proteins with decreased accumulation pattern included: heat shock protein (Bnascaffold286T0010300WE and BnaA01T01 12500WE), patellin-4-like (BnaC05T0253300WE), ankyrin (BnaA01T0020600WE), and thioredoxin (BnaA03T0276100WE) (Supplementary Table 4).



Temporal Changes in the Root Proteome of Clubroot Resistant and Clubroot Susceptible Progeny Lines in Response to the Pathogen

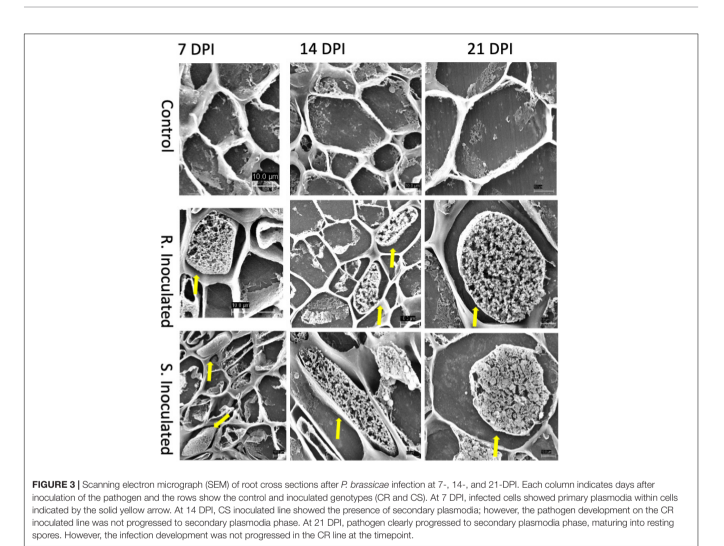
We profiled the proteome level differences that were induced in the CR- and CS-progeny lines in response to the pathogen at the three time points. In this study, comparisons were made between the uninoculated controls and pathogen-challenged CR- and CSplants at these time points.

### 7 Days Post Inoculation

At 7 DPI, there were 321 and 205 DAPs that increased in abundance in CR- and CS- lines, respectively (**Figure 4A**); whereas 122 and 244 were decreased in abundance in the CR- and CS- lines, respectively in response to the pathogen (**Figure 4A**). Among these, six proteins increased, while seven decreased in abundance in both lines (**Figure 4A**).

Among the proteins that exhibited similar trend of change in abundance in both CR- and CS-progeny lines, hereinafter referred to as 'shared proteins' (**Supplementary Table 2-4A**). Plant and pathogen interaction related proteins such as major intrinsic protein (aquaporin TIP1-2) (BnaA02T0335400WE) and germin-like protein (BnaA01T0301400WE) from the cupin family were observed to decrease in abundance in both CR and CS lines. In contrast, cysteine rich receptor like protein kinase (BnaA06T0336500WE) involved in salt and antifungal stress responses was observed to increase in abundance in both the CR- and CS-progeny lines following pathogen challenge (**Supplementary Table 2-4A**).

In addition to the "shared proteins" above, we also identified the proteins that exhibited an increase in abundance in the CR- and decreased in the CS-progeny lines and viceversa, hereinafter referred to as "contrasting proteins." There were 52 DAPs that exhibited these contrasting trends in abundance in the progeny lines (Supplementary Table 5-4A). These DAPs that increased in the CR- and decreased in CS-progeny lines included stress related proteins, for instance, ferredoxin (BnaC03T0212500WE), peroxidase (BnaC09T0553700WE), chitin elicitor-binding protein (BnaC03T0429600WE), and remorin (BnaC08T0350700WE) (Supplementary Table 5-4A). Other proteins such as thaumatin



(BnaC08T0208400WE and BnaA08T0229700WE) and peroxidase C3 (BnaC04T0547800WE) were observed to be decreased in the CR- and increased in the CS-progeny lines in response to the pathogen infection (**Supplementary Table 5-4A**).

#### 14 Days Post Inoculation

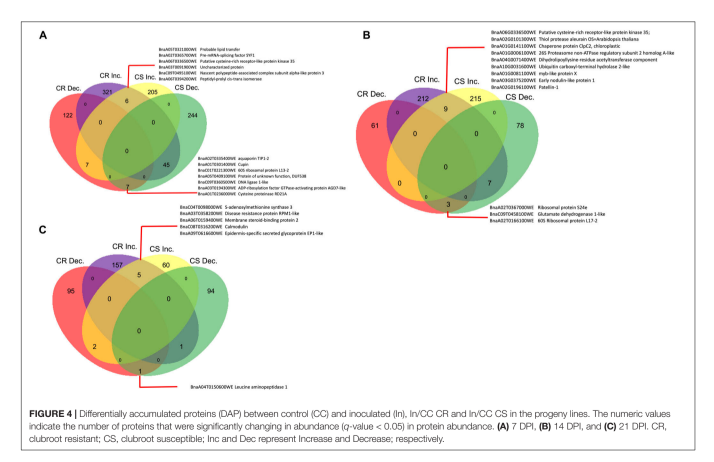
When a similar comparison was made at 14 DPI, there were 212 and 215 DAPs, that increased in abundance in CR- and CS-progeny lines, respectively, whereas 61 and 78 decreased in the CR- and CS-progeny lines, respectively, following pathogen challenge (Figure 4B). Nine proteins increased and three proteins decreased in abundance in both CR- and CS-progeny lines (Figure 4B). Seven proteins demonstrated contrasting trends in the CR- and CS- lines including one protein (BnaA05T0429000WE), orthologous to coiled coil domain which increased in abundance in the CR-progeny lines and decreased in the CS-progeny line (Supplementary Table 5-4B).

In the pool of "shared proteins" between CR- and CS-progeny lines, putative protein, BnaA06G0336500WE (orthologous to cysteine rich receptor like kinase) increased in abundance by  $>3.93 \log_2$  fold in the CR-progeny line in response to the pathogen (**Supplementary Table 2-4B**). Similarly, two proteins, BnaA10G0031600WE (orthologous to ubiquitin) and BnaA01G0006100WE (orthologous to 26S proteasome) increased significantly in both the CR- and CS-progeny lines (**Supplementary Table 2-4B**). Other plant stress related proteins orthologous to patellin (BnaA02G0196100WE), chaperone (BnaA01G0141100WE), cathepsin (BnaA02G0101300WE) were also observed to be significantly increased in both CR- and CS-progeny lines in which the relative accumulation of the first two were higher in CR-progeny lines whereas the latter (BnaA02G0101300WE) was higher in CS-progeny lines (**Supplementary Table 2-4B**).

#### 21 Days Post Inoculation

At 21 DPI, there were 157 and 60 DAPs that increased in abundance in CR- and CS-progeny lines, respectively, whereas 95 and 94 DAPs decreased in abundance in CR- and CS-progeny lines, respectively, following pathogen challenge (**Figure 4C**). Five proteins were found to increase in both CR- and CS-progeny lines (**Supplementary Table 2-4C**), while three proteins exhibited contrasting trends in abundance in the CR- and CS-progeny lines including one of the putative proteins (BnaC06T0464800WE) orthologous to metacaspase-4 which was increased in the CR-

Sample description	Condition	AVG precursors	AVG peptides	AVG protein groups	AVG proteins
Progeny 14 DPI control resistant	PG14CR	8539.00	6351.67	1904.33	3431.67
Progeny 14 DPI control susceptinle	PG14CS	8526.67	6427.67	1902.67	3496.33
Progeny 14 DPI inoculated resistant	PG14IR	7812.33	5768.67	1778.67	3088.67
Progeny 14 DPI inoculated susceptinle	PG14IS	9049.33	6794.33	2010.67	3657.67
Progeny 21 DPI control resistant	PG21CR	7829.67	5871.33	1797.33	3216.67
Progeny 21 DPI control susceptinle	PG21CS	9106.67	6774.00	2024.33	3689.33
Progeny 21 DPI inoculated resistant	PG21IR	7593.33	5670.33	1769.67	3077.67
Progeny 21 DPI inoculated susceptinle	PG21IS	9249.33	6881.33	2011.33	3703.33
Progeny 7 DPI control resistant	PG7CR	9775.00	7162.67	2061.67	3779.00
Progeny 7 DPI control susceptinle	PG7CS	8518.00	6295.00	1908.67	3408.33
Progeny 7 DPI inoculated resistant	PG7IR	8608.67	6462.00	1936.67	3539.33
Progeny 7 DPI inoculated susceptinle	PG7IS	8618.67	6360.67	1881.33	3343.00
Parent 14 DPI control resistant	PR14CR	5336.00	4044.33	1320.67	2296.33
Parent 14 DPI control susceptinle	PR14CS	7720.33	5841.33	1779.00	3110.00
Parent 14 DPI inoculated resistant	PR14IR	7889.67	5828.00	1762.67	3118.33
Parent 14 DPI inoculated susceptinle	PR14IS	7001.67	5223.00	1621.67	2797.00
Parent 21 DPI control resistant	PR21CR	9818.33	7351.67	2121.00	3922.00
Parent 21 DPI control susceptinle	PR21CS	6334.00	4900.00	1533.33	2735.67
Parent 21 DPI inoculated resistant	PR21IR	6497.33	4930.00	1543.67	2788.33
Parent 21 DPI inoculated susceptinle	PR21IS	8413.33	6325.00	1910.00	3474.67
Parent 7 DPI control resistant	PR7CR	6483.67	4927.00	1603.67	2753.33
Parent 7 DPI control susceptinle	PR7CS	7494.33	5672.00	1718.00	3151.67
Parent 7 DPI inoculated resistant	PR7IR	8560.00	6394.67	1923.33	3532.67
Parent 7 DPI inoculated susceptinle	PR7IS	9224.33	6784.67	2001.00	3652.33



and decreased in the CS-progeny line (**Supplementary Table 5-4C**). In the pool of shared proteins, proteins orthologous to calmodulin (BnaC08T0316200WE), disease resistance protein (RPM1-like) (BnaA03T0358200WE), *s*-adenosylmethionine synthase (BnaC04T0098000WE), and epidermis specific secreted glycoprotein (BnaA09T0616600WE) were increased significantly in both CR- and CS-progeny lines (**Supplementary Table 2**).

## Temporal Changes in the Root Proteome of Clubroot Resistant and Clubroot Susceptible Parent Lines in Response to the Pathogen

In addition to the aforementioned comparison of the pathogeninduced, temporal proteome changes at the three different time points CR- and CS-progeny lines, we also performed a similar study in the CR- and CS-parent lines. Similar to the progeny lines, in this case, comparisons were made between the uninoculated controls and pathogen-challenged CR and CS plants at these time points.

#### 7 Days Post Inoculation

At 7 DPI, we observed 87 and 88 proteins that increased in abundance in CR and CS-parent lines, respectively, and 362 and 188 proteins were observed to decrease in CR- and CS-parent lines, respectively, in response to the pathogen (Figure 5A). In addition, we observed that four proteins increased in both CR- and CS-parent lines and 25 proteins decreased in both parent lines (Figure 5A). Among these, proteins orthologous to dehydrin (BnaC06T0291300WE, BnaA07T02 39200WE, and BnaC06T0439800WE) were increased in abundance in the root proteome of the CS-parent line, while cysteine proteinase inhibitor (BnaA02T0044300WE), remorin (BnaC04T0592600WE), auxilin (BnaC03T0627000WE), heat shock 70 kDa (BnaC06T0068300WE and BnaC03T0112700WE), calmodulin (BnaC08T0316200WE), and ferritin (BnaA07T01 87000WE) were decreased in abundance in both the CR- and CS-parent lines in response to the pathogen (Supplementary Table 2-5A).

Among the contrasting proteins, there were 27 DAPs that exhibited opposing trends in abundance following pathogen inoculation at 7 DPI (**Supplementary Table 5-5A**). They included stress related proteins, for instance, cysteine proteinase RD19a (Bnascaffold1716T0016400WE), calnexin (BnaA06T0265000WE), allene oxide cyclase 4 (BnaA08T0253000WE), peroxidases (BnaC01T0310600WE and BnaA05T0420500WE) (**Figure 5A** and **Supplementary Table 5-5A**). The latter two peroxidases and RD19a were increased in the CS lines and decreased in the CR lines; whereas the calnexin and AOC were decreased in the CS line and increased in the CR line.

#### 14 Days Post Inoculation

When a similar comparison was made at 14 DPI, there were 55 and 35 DAPs increased in abundance in CR- and CSparent lines, respectively, whereas 34 and 155 DAPs decreased in abundance in the CR- and CS- parent lines, respectively in response to the pathogen (**Figure 5B**). One of the putative proteins (BnaC06T0291300WE) orthologous to dehydrin ERD14 demonstrated a contrasting pattern, increasing in the CR and decreasing in the CS-parent line (**Figure 5B** and **Supplementary Table 5-5B**). No shared proteins were detected in this time point.

#### 21 Days Post Inoculation

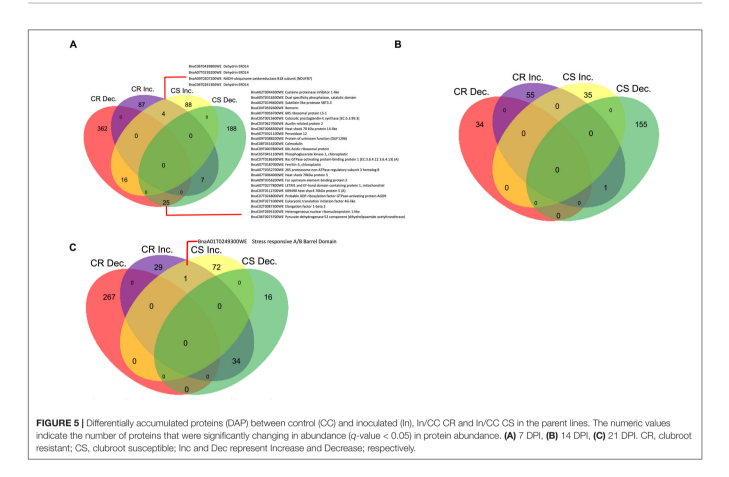
At 21 DPI, when the root proteomes of CR- and CS-parent lines were compared, we observed 29 and 72 DAPs that were increased in abundance in CR- and CS-parent lines, respectively, whereas 267 and 16 DAPs decreased in abundance in CR- and CS-parent lines, respectively, following pathogen challenge (Figure 5C). One protein (BnaA01T0249300WE), orthologous to stress responsive A/B barrel domain, was observed to increase in both CR- and CS-parent lines (Supplementary Table 2-5C), whereas 34 proteins exhibited contrasting trends in abundance in the CR- and CS-parent lines including ten putative proteins that are potentially involved in mediating plant and pathogen responses (Figure 5C and Supplementary Table 5-5C). These contrasting proteins included fasciclin-like arabinogalactan (BnaA01T0140800WE, BnaA10T0023700WE, and BnaA09T0184900WE), KDELtailed cysteine endopeptidase (BnaA06T0163700WE), thioredoxin-like domain (BnaA09T0477700WE), peroxidase (BnaA09T0646900WE), and patellin-4 (BnaC07T0112600WE) (Supplementary Table 5-5C). All the orthologs were increased in the CR-parent and decreased in the CS-parent lines following pathogen challenge (Figure 5C and Supplementary Table 5-5C).

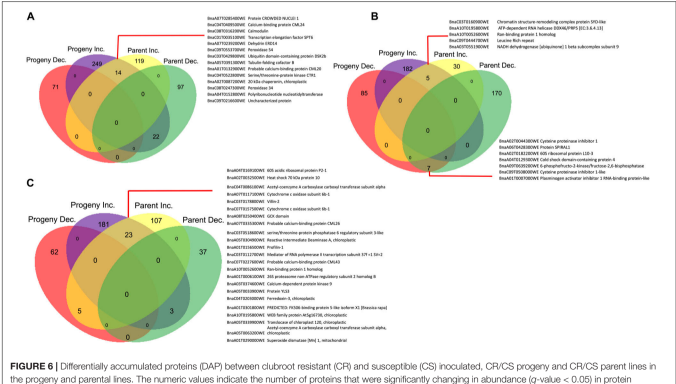
## Pathogen-Induced Temporal Changes in Root Proteome—A Comparison of Clubroot Resistant/Clubroot Susceptible Progeny and Clubroot Resistant/Clubroot Susceptible Parent Lines

In this case, we compared the pathogen-induced proteome level changes in the CR-parent and CR-progeny lines against their respective CS-counterparts. In other words, we compared the pathogen-induced temporal changes in the proteome of the CR-progeny against similar changes taking place in the CS-progeny lines and the CR-parent against the CS-parent lines at the three different time points. Our results at each time point are described below.

#### 7 Days Post Inoculation

At 7 DPI, 249 and 119 DAPs increased in abundance in the CS-progeny and CS-parent lines, respectively following pathogen challenge (**Figure 6A**). While 71 and 97 DAPs decreased in abundance in the CS-progeny and CS-parent lines, respectively (**Figure 6A**). There were 14 shared proteins increased in both CR-progeny and in the CR-parent, which included stress responsive metabolic compounds such as, calmodulin (BnaC08T0316200WE), dehydrin ERD14 (BnaA07T0239 200WE), peroxidase (BnaC09T0553700WE, BnaC08T0247 300WE), and calcium binding protein (BnaC04T0409500WE) (**Figure 6A** and **Supplementary Table 3-6A**).





abundance. (A) 7 DPI, (B) 14 DPI, (C) 21 DPI. Inc and Dec represent Increase and Decrease; respectively.

#### 14 Days Post Inoculation

When a similar comparison was made at 14 DPI, 182 and 30 proteins increased in abundance in the CS-progeny and CS-parent lines, respectively, following pathogen challenge (Figure 6B). While 85 and 170 proteins decreased in abundance in the CS-progeny and CS-parent lines, respectively (Figure 6B). There were five shared proteins that increased in abundance in both the CS-progeny and the CS-parent, including a protein orthologous to leucine rich repeat domain (BnaC09T0444700WE), RNA binding protein 1 (BnaA10T0052600WE), and chromatin structureremodeling complex protein SYD-like (BnaC03T0160900WE) (Supplementary Table 3-6B). One of the putative proteins (BnaA03T0551900WE) orthologous to NADH dehydrogenase [ubiquinone] 1 beta subcomplex subunit 9 increased > 9.0 log<sub>2</sub> fold in both CR-progeny and CR-parent lines (Supplementary Table 3-6B). Among the shared proteins, there were seven proteins increased in abundance in both CS-progeny and CS-parent group. These proteins included the orthologs of cysteine proteinase inhibitor (BnaA02T00443 00WE and BnaC09T0508000WE), SPIRAL1-like protein (BnaA06T0428300WE), Cold shock domain-containing protein (BnaA04T0129300WE) and 60S ribosomal protein (BnaA02T0182200WE) (Supplementary Table 3-6B).

#### 21 Days Post Inoculation

At 21 DPI, there were 181 and 107 DAPs that were increased in the CR-progeny and CR-parent lines, respectively; while 62 and 37 were decreased in the CR-progeny and CR-parent lines, respectively (**Figure 6C**). Twenty-three proteins were commonly increased in CR-progeny and CR-parent lines, nine of them were orthologous to stress response proteins including heat shock 70 (BnaA02T0032500WE), villin (BnaC03T0178800WE), calcium binding protein (CML43) (BnaC07T0227600WE), calcium dependent kinases (BnaA03T0374600WE), ferredoxin (BnaC04T0203000WE), and superoxide dismutase (**Figure 6C**). One of the proteins orthologous to superoxide dismutase [Mn] 1 (BnaA01T0290000WE) was increased > 2 folds in both CR-progeny and CS-parent lines (**Supplementary Table 3-6C**).

### Gene Ontology Enrichment Analysis of Differentially Abundant Proteins in the Root Proteomes of Progeny Lines

The DAPs that were identified in the various comparisons described above, that were temporally regulated as a result of pathogen challenge in the CS-progeny, CR-progeny, CS-parent, and CR-parent were further analyzed for their potential biological functions. The results obtained with respect to the enriched categories in the GO biological processes and molecular functions are provided in **Supplementary Tables 6A–D**. Our key findings are described below.

#### **Clubroot Resistant-Progeny**

By 21 DPI, 58 unique GO terms associated with various biological processes were significantly (FDR < 0.05) enriched in the CR-progeny lines (**Supplementary Table 6A**). The

vast majority of these biological processes included: cellular process (GO:0009987), response to stimulus (GO:0050896), protein metabolic process (GO:0019538), and response to stress (GO:0006950) in the CR-progeny lines (**Supplementary Table 6A**). In addition, there were several categories that are associated with plant-microbe interactions including: response to - biotic stimulus (GO:0009607), – bacterium (GO:0009617), – other organism (GO:0051707), -oxidative stress (GO:0006979), and – ROS (GO:0000302) (**Supplementary Table 6A**).

In terms of molecular functions, there were 24 enriched categories in the CR-progeny lines (**Supplementary Table 6A**). Categories that were highly enriched included: ion binding (GO:0043167), cation binding (GO:0043169), and calcium ion binding (GO:0005509) (**Supplementary Table 6A**). Molecular function categories related to ROS such as oxidoreductase (GO:0016491), antioxidant activity (GO:0016209), acting on peroxide as acceptor (GO:0016684), and peroxidase activity (GO:0004601) were greatly enriched indicating their potential involvement in signal transduction processes in response to the pathogen in the CR-progeny lines.

#### **Clubroot Susceptible-Progeny**

By 21 DPI, 59 unique GO terms associated with biological processes were significantly (FDR < 0.05) enriched in the CS-progeny lines (**Supplementary Table 6B**). The highest-level categories included response to -stimulus (GO:0050896), -stress (GO:0006950), and -abiotic stimulus (GO:0009628) (**Supplementary Table 6B**). Similar to the results described above for the CR-progeny lines, there were several categories directly associated with plant-microbe interactions such as response to -other organism (GO:0051707), defense response (GO:0006952), and – biotic stimulus (GO:0009607) (**Supplementary Table 6B**).

In terms of molecular functions, there were 18 enriched categories in the CS-progeny lines (Supplementary Table 6B). Categories that were enriched included: ion (GO:0043167), cation binding binding (GO:0043169), metal ion binding (GO:0046872), oxidoreductase activity (GO:0016491), and structural molecule activity (GO:0005198) (Supplementary Table 6B). Molecular function categories related to ROS, for instance, antioxidant activity (GO:0016209), oxidoreductase (GO:0016491), acting on peroxide as acceptor (GO:0016684), and intramolecular oxidoreductase activity (GO:0016860) were highly enriched indicating their potential role in plant and pathogen interaction in the CS-progeny lines as well (Supplementary Table 6B). In addition, GO categories related to metal binding such as calcium ion-(GO:0005509), copper ion- (GO:0005507), heme- (GO:0020037), and iron ion binding (GO:0005506) were highly enriched (Supplementary Table 6B).

## Shared and Contrasting Gene Ontology Categories in the Clubroot Resistant and Clubroot Susceptible Progeny Lines

In terms of biological process, 39 categories in the CR- and CSprogeny lines were similar and majority of them are potentially associated with plant-pathogen interactions, including GO categories in response to – stress (GO:0006950), – biotic stimulus (GO:0009607), – bacterium (GO:0009617), – other organism (GO:0051707), – ROS (GO:0000302), and defense response (GO:0006952).

Nineteen GO categories were unique to the root proteome of the CR-progeny lines including cellular process (GO:0009987), which was one of the highest enriched categories in the CR progeny (**Supplementary Tables 6A,B**). Other categories such as protein metabolic process (GO:0019538), cellular protein metabolic process (GO:0044267), translation (GO:0006412) were also among the unique categories in the CR-progeny lines (**Supplementary Tables 6A,B**). Eleven GO categories were unique to CS-progeny including defense response (GO:0006952), cellular component biogenesis (GO:0044085), and cellular component assembly (GO:0022607) (**Supplementary Tables 6A,B**).

In terms of molecular functions, 17 GO categories were identical in the CS and CR progeny including ion binding (GO:0043167), metal ion binding (GO:0046872), and structural molecule activity (GO:0005198) (**Supplementary Tables 6A,B**). The GO categories such as oxidoreductase activity (GO:0016491) and acting on sulfur group of donors (GO:0016667) were the only GO terms unique to the CS-progeny line (**Supplementary Table 6B**). Similarly, seven unique GO categories were found in CR-progeny line (**Supplementary Table 6A**) including binding (GO:0005488), endopeptidase activity (GO:0004175), translation factor activity, and nucleic acid binding (GO:0008135) (**Supplementary Table 6A**).

## Gene Ontology Enrichment Analysis of Differentially Abundant Proteins in the Root Proteomes of Parent Lines Clubroot Resistant-Parent

By 21 DPI, 55 unique GO terms associated with biological processes were significantly (FDR < 0.05) enriched in the root proteome of the CR-parent line (**Supplementary Table 6C**). The highest-level categories included cellular process (GO:0009987), response to -stimulus (GO:0050896), -stress (GO:0006950), cellular protein metabolic process (GO:0019538), and response to chemical stimulus (GO:0042221) in the parent lines (**Supplementary Table 6C**). There were several categories associated with plant-microbe interactions including response to - biotic stimulus (GO:009607), -bacterium (GO:0009617), -other organism (GO:0051707), and response to ROS (GO:0000302) that were observed. Similarly, enriched categories associated with response to -metal ion (GO:0010038), and -cadmium ion (GO:0046686) were also observed to be highly enriched (**Supplementary Table 6C**).

In terms of molecular functions, there were 22 enriched categories in the root proteome of the CR-parent line in response to the pathogen (**Supplementary Table 6C**). These highly enriched categories include binding (GO:0005488), ion binding (GO:0043167), cation binding (GO:0043169), and metal ion binding (GO:0046872) (**Supplementary Table 6B**). Molecular function categories related to ROS such as antioxidant

activity (GO:0016209), peroxidase activity (GO:0004601), and intramolecular oxidoreductase activity (GO:0016860) were highly enriched. Similarly, GO categories associated with calcium ion binding (GO:0005509) and heme binding (GO:0020037) were among some of the highest-level enriched categories in the molecular function (**Supplementary Table 6C**).

#### **Clubroot Susceptible-Parent**

By 21 DPI, 36 unique GO terms associated with biological processes were significantly (FDR < 0.05) enriched in the CS-parent line (**Supplementary Table 6D**). GO enriched categories in the root proteome of the CS-parent line was similar to that of the CR-parent line. The highest-level categories included response to -stimulus (GO:0050896), -stress (GO:0006950), and protein metabolic process (GO:0019538) (**Supplementary Table 6D**). There were several categories directly associated with plant-microbe interactions such as response to – other organism (GO:0051707), – defense response (GO:0006952), -biotic stimulus (GO:0009607), and -bacterium (GO:0009617) (**Supplementary Table 6D**).

In terms of molecular function, there were 22 enriched categories in the CS-parent lines. Categories including binding (GO:0005488), cation binding (GO:0043169), ion binding (GO:0043167), and metal ion binding (GO:0046872) were highly enriched in the CS-parent line in response to the pathogen challenge (**Supplementary Table 6D**). Molecular function categories related to ROS such as antioxidant activity (GO:0016209), peroxidase activity (GO:0004601), and oxidoreductase (GO:0016491) were highly enriched (**Supplementary Table 6D**). Likewise, metal ion binding related categories, such as calcium ion binding (GO:0005509) and heme binding (GO:0020037) were also highly enriched (**Supplementary Table 6D**).

## Shared and Contrasting Gene Ontology Categories in the Clubroot Resistant and Clubroot Susceptible Parent Lines

In terms of biological process, 34 categories in the root proteome of the CR-parent and the CS-parent line were identical and majority of them were potentially associated with plant and pathogen interaction, including GO categories in response to stimulus (GO:0050896), -stress (GO:0006950), cellular protein metabolic process (GO:0019538), and response to chemical stimulus (GO:0042221) (**Supplementary Tables 6C,D**). There were also several categories associated with plant-microbe interactions such as response to – biotic stimulus (GO:0009607), -bacterium (GO:0009617), and -oxidative stress (GO:0006979) that were observed (**Supplementary Tables 6C,D**).

Three GO categories were unique to the root proteome of the CS-parent line including protein metabolic process (GO:0019538), protein import (GO:0017038), and protein localization in organelle (GO:0033365) (**Supplementary Tables 6C,D**). Similarly, 22 GO categories were unique to CR-parent including cellular process (GO:0009987), post-embryonic development (GO:0009791), and response to metal ion (GO:0010038) (**Supplementary Tables 6C,D**). In terms of molecular function, 22 GO categories were identical in the root proteome of the CS- and CR-parent line including binding (GO:0005488), ion binding (GO:0043167), cation binding (GO:0043169), and metal ion binding (GO:0046872) (**Supplementary Tables 6C,D**). The GO terms, oxidoreductase activity (GO:0016491) and isomerase activity (GO:0016853) were unique in the root proteome of the CR-parent line (**Supplementary Table 6C**). Similarly, two unique GO categories, *s*-acyltransferase activity were found in the root proteome of the CS-parent line (**Supplementary Table 6D**).

In the following sections, we describe the major groups of proteins that were found to be differentially abundant, and which are potentially involved in mediating the resistance phenotype in the CR lines used in this study.

## **Plant Hormone Related Proteins**

Due to the important roles of phytohormones in the progression of clubroot disease in B. napus, we analyzed our proteome data for DAPs associated with phytohormone biosynthesis and/or signaling. When the control and inoculated lines were compared, i.e., inoculated vs. control-CR and inoculated vs. control -CS lines, in the case of both progeny and parents, there were six DAPs that were related to salicylic acid (SA) regulation. Four of them (BnaA03T0555800WE, BnaC07T0044900WE, Bnascaffold286T0028200WE, and BnaA01T0310200WE) were orthologous to Arabidopsis regulatory protein, non-expressor of pathogenesis related 1 (NPR1) and NPR4, whereas two of them (BnaA09T0109100WE and BnaA10T0130500WE) were orthologous to systematic acquired resistance (SAR) deficient 1 protein (Supplementary Table 7A). Three of the proteins [(Bnascaffold286T0028200WE and BnaC07T004 4900WE, orthologous to regulatory protein NPR1), and (BnaA02T0079400WE, orthologous to aminotransferase ALD1)] showed consistent pattern in both parent and progeny lines. Two of the NPR1 orthologs (Bnascaffold286T0028200WE BnaC07T0044900WE) showed contrasting pattern between CR and CS lines, thereby the proteins increase in CR and decrease in CS in both the parent and progeny lines, while the protein BnaA02T0079400WE consistently increased in accumulation in both CR and CS root proteomes across both parent and progeny lines (Supplementary Table 7A).

There were 12 DAPs that were orthologous to proteins related to ethylene (ET) and jasmonic acid (JA) biosynthetic pathways (Supplementary Table 7A). Seven DAPs were orthologous to serine/threonine-protein kinase CTR1, which functions as a negative regulator of the ET biosynthetic pathway (Kieber et al., 1993). The protein, (BnaA06T0336500WE) orthologous to serine/threonine-protein kinase CTR1, increased in abundance in both CR and CS and the pattern was consistent in both the parent and progeny lines (Supplementary Table 7A). Similarly, there were three proteins (BnaA09T0013600WE, BnaA03T0361400WE, and BnaC04T0098000WE) orthologous to s-adenosylmethionine (SAM) synthase that exhibited a similar pattern of protein accumulation, i.e., an increase in both CR and CS, in both the parent and progeny lines (Supplementary Table 7A). As the pathogen infection progressed to 21 DPI, the protein, BnaA03T0361400WE, increased in the CR lines and decreased in abundance in the CS lines in both the parent and progeny (**Supplementary Table 7A**). Similarly, the proteins, BnaA09T0013600WE and BnaC04T0098000WE, were increased in abundance in the CR lines in both the parent and progeny and the increasing pattern was similar in CS-progeny, however, the pattern in CS parent at 21 DPI was decreasing (**Supplementary Table 7A**).

Two plant hormones, auxin and cytokinins (CK), are involved in clubroot gall formation. We identified three auxin related proteins (BnaA10T0208100WE, BnaA05T040 9700WE, and BnaC03T0302500WE) and three CK related proteins (BnaC07T0399800WE, BnaA06T0322500WE, and BnaA01T0059700WE) (Supplementary Table 7A). As the infection progressed to 21 DPI, one of the putative proteins (BnaA05T0409700WE) orthologous to auxin-induced in root cultures protein 12 was increased in abundance in both CR and CS root proteomes and the pattern was similar in both the parent and progeny lines (Supplementary Table 7A). Similarly, one of the orthologs of histidine kinase (BnaA01T0059700WE) also increased at 21 DPI, showing shared pattern between CR-progeny, CR-parent, and CS-progeny; however, in the CS-parent, the protein decreased in abundance at 21 DPI (Supplementary Table 7A).

In an addition to the comparisons mentioned above, root proteomes were compared between inoculated CR-parent vs. inoculated CS-parent and inoculated CR-progeny vs. inoculated vs. CS-progeny. There were eight proteins representing five important phytohormones, including auxin, CK, ET, JA, and SA (Supplementary Table 7B). Two SA related proteins (BnaC07T0044900WE and Bnascaffold286T0028200WE) were detected in both CR and CS root proteomes, and the latter showed a shared pattern with an increase in both the CR-parent and CR-progeny as the infection progressed to 21 DPI (Supplementary Table 7B). Other proteins involved in ET mediated (BnaA09T0013600WE) and JA mediated (BnaA08T0253000WE) defense responses were increased in the CR-progeny at all three times points. Four proteins related to phytohormones, auxin (BnaA05T0409700WE, BnaC03T03 02500WE) and CK (BnaC07T0399800WE, BnaA01T0059 700WE) were differentially abundant in both parent and progeny lines (Supplementary Table 7B). While both the CK related proteins were increased in CR-progeny, one CK related protein (BnaA01T0059700WE), that is orthologous to a histidine kinase, was consistently increased in both the CR- parent and progeny lines at 21 DPI (Supplementary Table 7B). Similarly, the auxin related protein (BnaA05G0409700WE), orthologous to auxin-induced in root cultures protein 12, also exhibited a shared pattern of abundance and increased in both CR-parent and CR-progeny lines (Supplementary Table 7B).

## Proteins Involved in Calcium Signaling

When the root proteomes were compared between the inoculated vs. control CR and inoculated vs. control CS lines, 32 differentially accumulated proteins orthologous to calcium signaling proteins were detected (**Supplementary Table 8A**). Among these proteins, 21 contained helix-loop-helix

structural motifs and were categorized as EF-hand domain containing proteins, nine were labeled as calreticulin family, and the remaining two were related to calmodulin binding protein (**Supplementary Table 8A**). As the infection progressed to 21 DPI, three putative calcium signaling proteins (BnaA07T0335300WE, BnaA03T0464700WE, and BnaA03T0374600WE) were consistently increased in the root proteome of the CR-progeny and CR-parent lines (**Supplementary Table 8A**). In contrast, all three proteins consistently decreased in both the CS-parent and CS-progeny lines at 21 DPI (**Supplementary Table 8A**).

In an addition to the comparison mentioned above, DAPs were assessed between inoculated CR-parent vs. inoculated CS-parent and inoculated CR-progeny vs. inoculated CS-progeny and 22 orthologs of calcium signaling proteins were identified (**Supplementary Table 8B**). Three of these putative proteins (BnaC08T0316200WE, BnaC07T0227600WE, and BnaA07T0335300WE), were orthologous to calcium binding proteins, and were consistently increased at all three time points in both the CR-parent and CR-progeny lines (**Supplementary Table 8B**).

## **Lignin Related Proteins**

The comparative analysis of the root proteomes between the inoculated vs. control CR and inoculated vs. control CS lines revealed that there were 24 DAPs that were orthologous to peroxidases, caffeic acid 3-O-methyltransferase, lignin forming anionic peroxidase, and s-adenosylmethionine (SAM) synthase 3 (METK3) (Figure 7 and Supplementary Table 9A). The enzyme METK3 is involved in lignin biosynthesis (Shen et al., 2002) and there were four protein orthologs for METK3 (BnaC03T0264500WE, BnaA09T0013600WE, BnaA03T0361400WE, and BnaC04T0098000WE), among which, BnaC03T0264500WE, was increased and the pattern was consistent at 21 DPI in CS-progeny and the CSparent (Supplementary Table 9A). Similarly, there were 13 proteins orthologous to peroxidases, proteins involved in pathogen response as well as during the biosynthesis and degradation of lignin. Among these proteins, three putative proteins (BnaA10T0260700WE, BnaC01T0221000WE, and BnaC08T0247300WE), were increased >  $1.0 - \log_2$  fold in the CR-progeny lines and decreased in the CS-progeny lines when the pathogen infection progressed to 21 DPI (Figure 7 and Supplementary Table 9A). The protein BnaC08T0247300WE (orthologous to peroxidase) was increased in the CS- and CR-parent line, thus exhibiting a trend that was similar to the CR-progeny lines (Supplementary Table 9A).

Root proteomes when compared between inoculated CRparent vs. inoculated CS-parent and inoculated CR-progeny vs. inoculated CS-progeny, there were 18 DAPs that are related to lignin (**Supplementary Table 9B**). While majority of the proteins showed contrasting pattern between the CR- parent and progeny lines, three of the proteins (BnaA03T0561300WE, BnaC02T0522200WE, and BnaC08T0247300WE) that are orthologous to peroxidases have shared pattern between parent and progeny, with an increase in the CR root proteomes of both the parents and the progeny lines (**Supplementary Table 9B**).

# Reactive Oxygen Species Related Proteins

A comparison of the root proteomes between inoculated vs. control CR and inoculated vs. control CS lines indicated that there were 20 putative, differentially abundant, ROS related proteins (Supplementary Table 10A). Five exhibited contrasting trends in abundance between the CR- and CSprogeny lines. Three of these proteins (BnaA09T0646900WE, BnaC04T0547800WE, and BnaC09T0113900WE), orthologous to l-ascorbate peroxidase 5, were decreased in abundance in the CR-progeny and increased in abundance in the CS-progeny lines (Figure 7 and Table S10). The remaining two proteins, BnaA05T0420500WE (orthologous to peroxiredoxin-2F) and BnaA10T0260700WE (orthologous to l-ascorbate peroxidase 5) increased in abundance in the CR-progeny and decreased in the CS-progeny (Figure 7 and Supplementary Table 10). In the case of these two proteins, the pattern of accumulation observed in the CR-progeny lines was also consistent in the CR-parent line with an increase in accumulation as the infection progressed to 21 DPI (Supplementary Table 10A).

Three other ROS-related proteins, BnaA03T0167800WE (orthologous to glutathione peroxidase 2), BnaA06T0044600WE peroxidase (orthologous L-ascorbate to 1), and BnaA09T0631100WE (orthologous to superoxide dismutase) exhibited a similar pattern of protein accumulation in CR-, CS-progeny, and CR-parent lines where the proteins increased in abundance (Supplementary Table 10A). However, two of these proteins (BnaA03T0167800WE and BnaA06T0044600WE) in the CS-parent decreased in abundance as the infection progressed to 21 DPI (Supplementary Table 10A). Nevertheless, among these three proteins, the protein BnaA09T0631100WE (orthologous to superoxide dismutase) has shared pattern of protein abundance between parent and progeny in both the CS and CR lines (Supplementary Table 10A).

When the inoculated CR-parent vs. CS-parent and inoculated CR-progeny vs. inoculated CS-progeny were compared, 31 DAPs were identified. Among them, six proteins [BnaC02T0522200WE-, BnaA03T0561300WE-, BnaC08T0247300WE-, orthologous to peroxidase; and BnaC06T0450900WE, orthologous to thioredoxin-like domain; BnaA01T0290000WE, orthologous to superoxide dismutase; and BnaC07T0044900WE, orthologous to ankyrin repeat domain) exhibited shared patterns between both the parent and progeny lines, with all of them increased in accumulation in the CR lines of both the parents and progeny as the disease progressed to 21 DPI (Supplementary Table 10B). The protein BnaA01T0290000WE increased > 2.0-log<sub>2</sub> fold change in both CR- parent and CR-progeny lines (Supplementary Table 10B).

## Dehydrins

Between the inoculated vs. control CR and inoculated vs. control CS lines, there were nine proteins orthologous to dehydrin, three of them (BnaC02T0286500WE, BnaC06T0291300WE, and BnaA07T0239200WE) were increased in abundance in the CR-progeny and decreased in the CS-progeny lines (**Supplementary Table 11A**). All three proteins increased in abundance at

7 DPI in the CS-progeny lines; however, as the infection progressed, they were consistently decreased at 14 and 21 DPI, demonstrating an opposite pattern of protein accumulation relative to the CR-progeny lines (**Supplementary Table 11A**). In the case of BnaA07T0239200WE, we observed that it was increased in abundance in both the CR-progeny as well as

the CR-parent lines as the infection progressed to 21 DPI (Supplementary Table 11A).

In a separate comparison of root proteomes between inoculated CR-parent vs. inoculated CS-parent and inoculated CR-progeny vs. inoculated CS-progeny, four orthologs of dehydrin (BnaC06T0291300WE, BnaC06T0439800WE, BnaA

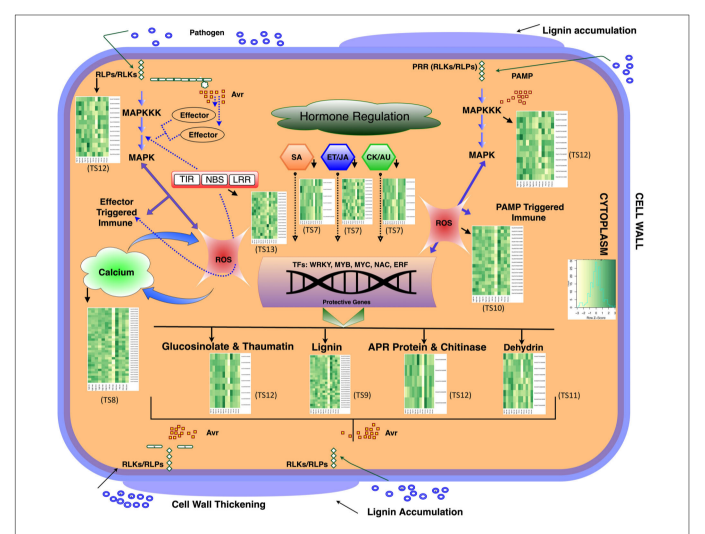


FIGURE 7 | Schematic diagram of plant-pathogen interaction upon P. brassicae inoculation in B. napus. Heatmaps were developed using log<sub>2</sub> fold change values at 7-, 14-, and 21-DPI and were shown besides each major metabolic processes in the plant-pathogen interaction pathway indicated by a dark solid arrow. Dark Green indicates increase in abundance and light color indicates decrease in abundance. Predicted proteins that were included in the heatmaps were significantly (q < 0.05) differentially abundant at least in one time point across all genotypes investigated. During P. brassicae infection, pathogen releases signal metabolites such as PAMPs and effectors and these elicitors were recognized by host cells. Plant receptors such as receptor like -kinases and -proteins (RLKs/RLPs) and resistance (R) proteins interact with the pathogen signal compounds, thereby activates the MAP kinases (signal transduction). Transcription factors are regulated and successively plant defense related metabolic pathways are activated, proteins related to phytohormone biosynthesis pathway, especially SA, JA, and ET biosynthetic pathway and other secondary metabolism were activated. Pathogen defense related secondary metabolites such as glucosinolates, small molecules thaumatin, polyssacharides (lignin), glycosyl hydrolases (chitinases), and a spectrum of antimicrobial and defense related proteins including dehydrin proteins were differentially accumulated in response to the pathogen. Growth regulators, auxin (AU) and cytokinin (CK) were manipulated by the pathogen, the hormones are regulated abnormally, thereby leading to the hypertrophy in the root tissues. There is an interplay between ROS and calcium signaling compounds during plant and pathogen (P. brassicae) interaction. Calcium acts as a second messenger and with the involvement of Ca<sup>2+</sup> -pumps, -buffers, and -exchangers, they orchestrates and regulates cellular processes and participates in an interplay with other signaling transduction processes such as ROS. Detail annotation of protein ids shown in the heatmaps are provided in the supplementary tables: TS7 – TS13 (Supplementary Tables 7–13). Avr, avirulence factor; RLP, receptor like proteins; RLK, receptor like kinase; SA, salicylic acid; JA, jasmonic acid; ET, ethylene; CY, cytokinin; AU, auxin; PRR, pathogen recognition receptor; PAMP, pathogen associated molecular pattern; MAPK, mitogen-activated protein kinase; TIR, toll/interleukin-1 receptor; NBS, nucleotide binding site; LRR, leucine rich repeat; APR, adult plant resistance. PgR-7d, CR progeny 7 DPI; PgR-14d, CR progeny 14 DPI; PgR-21d, CR progeny 21 DPI; PgS, CS progeny; PrR, CR parent; PrS, CS parent.

07T0354800WE, and BnaA07T0239200WE) were identified (**Supplementary Table 11B**). These proteins decreased in the CR root proteomes of both parent and progeny lines as the infection progressed to 21 DPI (**Supplementary Table 11B**).

## **Heat Shock Proteins**

A comparison of the root proteomes between inoculated vs. control CR and inoculated vs. control CS lines revealed that there were 16 differentially accumulated proteins that were related to heat shock proteins (**Supplementary Table 12A**). Three of the proteins (BnaC06T0068300WE, BnaC04T0303200WE, and BnaC06T0195100WE) orthologous to HSP70 and HSP90, were increased as the infection progressed to 21 DPI in CR lines and decreased at 21 DPI in the CS lines in the case of both the parent and progeny lines except for BnaC06T0195100WE, which increased at 21 DPI in CS-parent lines (**Supplementary Table 12A**).

Furthermore, root proteomes when compared between inoculated CR-parent vs. inoculated CS-parent and inoculated CR-progeny vs. inoculated CS-progeny, we identified 14 heat shock proteins orthologous to HSP70 and HSP90 kDa (**Supplementary Table 12B**). Four of the proteins (BnaA02T0032500WE, BnaC06T0068300WE, BnaC04T0303200WE, and Bnascaffold286T0010300WE) were consistently increased in the CR-parent and CR-progeny as the infection progressed to 21 DPI Table S12-B).

## **Resistance Proteins**

When root proteomes were compared between inoculated vs. control CR and inoculated vs. control CS lines, 24 differentially accumulated R proteins were identified. Seven of these proteins (BnaA01T0054600WE, BnaA03T0182000WE, BnaA03T0374600WE, BnaA06T0336500WE, BnaA10T0238 900WE, BnaC04T0522800WE, and BnaC02T0029600WE) were orthologous to the disease resistance X-TIR-NB-LRR-X family and three (BnaC08T0367400WE, BnaC09T0444700WE, and BnaA06T0288700WE) were orthologous to proteins containing leucine rich repeat (LRR) domains and the remaining were orthologous to stress response and calcium dependent protein kinases (Figure 7 and Supplementary Table 13A). Three of these proteins (BnaA01T0054600WE, BnaA03T0374600WE, and BnaA06T0336500WE) orthologous to WRKY19 increased in abundance in the CR-progeny as well as in the CR-parent lines; whereas, the latter two (BnaA01T0054600WE and BnaA06T0336500WE) were consistently increased at all time points in both the CR and CS genotypes (progeny and parent), while one of the proteins (BnaA03T0374600WE) decreased in the CSparent and CS-progeny as the infection progressed to 21 DPI (Supplementary Table 13A).

When the inoculated CR-parent vs. inoculated CS-parent and inoculated CR-progeny vs. inoculated CS-progeny were compared, 15 DAPs that were related to disease resistance were identified (**Supplementary Table 13B**). Three proteins orthologous to WRKY19 showed shared pattern between parent and progeny lines; while two of them (BnaA03T0374600WE and BnaC01T0013800WE) were increased in the CR-parent and CRprogeny lines, and BnaA03T0397600WE was decreased in both CR parent and CR progeny as the infection progressed to 21 DPI (**Supplementary Table 13B**).

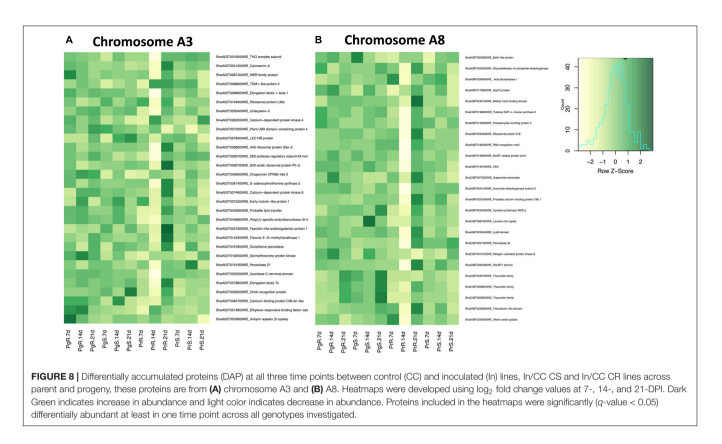
## Differentially Abundant Proteins Encoded by Genes Located on Chromosome A3 and A8

Resistance to clubroot has been frequently mapped to QTLs located on chromosomes A3 and A8 (Hasan et al., 2021). When we compared the root proteomes between inoculated vs. control CR and inoculated vs. control CS in both parent and progeny, we detected 30 and 25 DAPs that were located on chromosome A3 and A8, respectively (**Supplementary Tables 14A,C**).

Among the DAPs, located on chromosome A3, we identified 15 that were orthologous to a range of plant-pathogen interaction related proteins, including fasciclin-like arabinogalactan (BnaA03T0031800WE), peroxidase (BnaA03T0191900WE), calumenin (BnaA03T0041500WE), chitin recognition protein (BnaA03T0306200WE), and ankyrin (BnaA03T0555800WE) (Figure 8A and Supplementary Table 14A). Five of these proteins increased in abundance by 21 DPI in the CR-progeny lines, with two of the five DAPs, BnaA03T0142300WE (orthologous to flavone 3'-O-methyltransferase 1) and BnaA03T0374600WE (orthologous to calcium-dependent protein kinase 9) increased in CR and decreased in CS lines in both the progeny as well as parent lines (Supplementary Table 14A). Similarly, three proteins, BnaA03T0167800WE (orthologous to glutathione peroxidase), BnaA03T0306200WE (chitin recognition protein), and BnaA03T0262200WE (calcium-dependent protein kinase 4) were observed to be increased in both the CR-progeny as well as the CR-parent lines (Supplementary Table 14A).

In separate root proteome comparisons between inoculated CR-parent vs. inoculated CS-parent and inoculated CR-progeny vs. inoculated CS-progeny, there were 26 DAPs that are located on chromosome A3. Four proteins among these, BnaA03T0003400WE (orthologous to mitochondrial ATPase inhibitor), BnaA03T0041500WE (calumenin-A), BnaA03T0402800WE (lipid transfer), and BnaA03T0135900WE (C2 domain) showed consistent pattern of accumulation in parent and progeny lines (**Supplementary Table 14B**). Three of these proteins (BnaA03T003400WE, BnaA03T0402800WE, and BnaA03T0041500WE) were increased in the CR-parent and CR-progeny line, and one of the proteins (BnaA03T0135900WE) was decreased in the CR-lines in both parents and the progeny (**Supplementary Table 14B**).

proteomes When root compared between were inoculated vs. control CR and inoculated vs. control CS in both parent and progeny, 16 putative DAPs were related to plant and pathogen interactions and were located on chromosome A8 (Supplementary Table 14C). Some of the major plant-microbe interaction related included proteins peroxidase (BnaC08T0247300WE), thaumatin (BnaC08T0208400WE, BnaC08T0208200WE, and BnaA08T0229700WE), and villin (BnaA08T0139700WE)



(Figure 8B and Supplementary Table 14C). As the infection progressed to 21 DPI, three proteins (BnaA08T0229700WE, BnaC08T0208200WE, and BnaC08T0208400WE) orthologous to thaumatin family proteins were increased by > 2.0-log<sub>2</sub> fold in the CR progeny lines (Supplementary Table 14C). The pattern of accumulation for the latter two were consistent in the CR parent line as well (Supplementary Table 14C). On the contrary, protein (BnaC08T0247300WE) orthologous to peroxidase 34 exhibited a contrasting pattern, wherein it increased in the CR-progeny and decreased in the CS-progeny lines (Supplementary Table 14C).

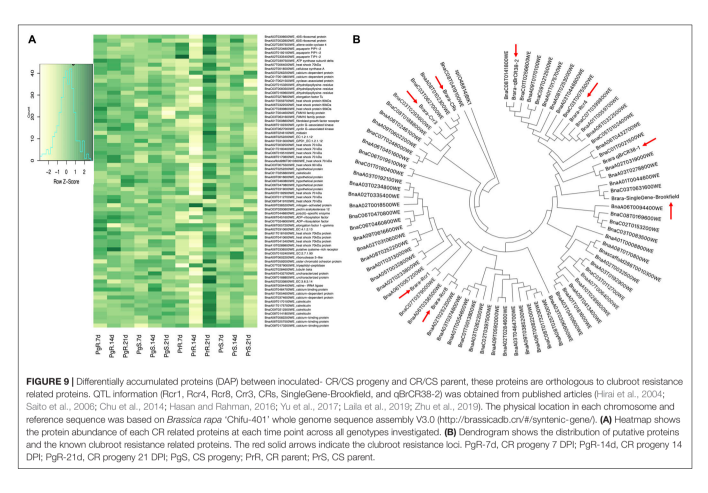
Furthermore, when the root proteomes of the inoculated CR-parent vs. inoculated CS-parent and inoculated CR-progeny vs. inoculated CS-progeny were analyzed, 22 DAPs were located on chromosome A8. Four proteins among these, BnaA08T0019900WE (orthologous to eukaryotic translation initiation factor 2alpha kinase 1), BnaA08T0170800WE (heat shock 70 kDa protein 9), and BnaA08T0250400WE (mitochondrial intermembrane space import and assembly protein 40-like), and BnaA08T0284200WE (ran-binding protein 1 homolog a-like) showed shared pattern of protein accumulation where each protein increased in abundance in the CR-lines of both parents and the progeny (**Supplementary Table 14D**).

## Proteins Associated With Clubroot Resistance Related Genes and Quantitative Trait Loci

Our current study has identified 74 DAPs (Figure 9A) that are orthologous to the eight CR related loci (Rcr8, Crr3, Rcr4,

Rcr1, CRs, SingleGene-Brookfield, and qBrCR38-1, qBrCR38-1) (Figure 9B) that were reported in the earlier studies (Hasan et al., 2021). The physical location of the DAPs in each chromosome was based on Brassica rapa 'Chifu-401' whole genome sequence assembly V3.01. There were 57 unique proteins potentially involved in plant and pathogen interaction, orthologs of calcium binding-, heat shock-, and aquaporin- related proteins were frequently observed in the dataset (Supplementary Table 15). There were two orthologs of calcium binding proteins on chromosome A3, BnaA03T0464700WE (chromosome location, A3 24574673 25715499 Rcr1) and BnaA03T0374600WE (chromosome location, A3 23823130 26806601 Rcr4) that were increased in both the CR-progeny as well as CR-parent lines and decreased in both the CS-progeny and CS-parent lines (Supplementary Table 15). Similarly, among the heat shock proteins (HSP), Hsp70 and Hsp90 were predominant and BnaA03T0410600WE (chromosome A8 20254038 21748038 gBrCR38-2) orthologous location to Hsp70, consistently increased in both CR-parent and CRprogeny lines. Other putative proteins, BnaA09T0616600WE (chromosome location A7\_20107291\_20209491\_qBrCR38-1) orthologous to D-mannose binding lectin, was increased in CR-progeny and CR-parent lines. Notably, BnaA06T0336500WE (chromosome location A3\_23823130\_26806601\_Rcr4), orthologous to cysteine-rich receptor-like protein kinase), exhibited increasing pattern of protein abundance in the CR and CS lines in both the progeny and parent upon pathogen

<sup>&</sup>lt;sup>1</sup>http://brassicadb.cn/#/syntenic-gene/



infection (**Figure 9A**). Finally, three putative orthologs of aquaporin proteins [BnaA03T0234800WE (chromosome location A3\_23823130\_26806601\_Rcr4), BnaA03T0192100WE (chromosome location A8\_12116024\_13046214\_SingleGene-Brookfield), and BnaA02T0335400WE (chromosome location A2\_22502031\_26348249\_Rcr8)] were decreased in both CR- and CS-progeny lines upon pathogen infection. One of the proteins (BnaA02T0335400WE) was consistently decreased at all time points in both the CS and CR lines in the case of both the parent and progeny lines (**Figure 9A**).

## DISCUSSION

Our current study has provided novel insights into the possible mechanisms underlying resistance to the clubroot pathogen. Although there are limited reports of proteomics and transcriptomics based investigation into the clubroot pathogen and *B. napus* interaction (Cao et al., 2008; Summanwar et al., 2019, 2020, 2021; Galindo-González et al., 2020; Zhou et al., 2020), our results align with earlier reports on *B. rapa* especially as it relates to calcium signaling protein kinases and calcium binding proteins in CR and CS root proteomes (Song et al., 2016). Similarly, the accumulation patterns of some of the vital defense-related glycosyl hydrolases including chitinases and thaumatin proteins, and phytohormone

(e.g., SA)-related proteins were consistently increased in the CR root proteome which is in agreement with previous transcriptome studies (Summanwar et al., 2019; Galindo-González et al., 2020). Moreover, in this study we have identified several other calcium signaling proteins, dehydrins, and proteins that were orthologous to CR loci in different chromosomes particularly, chromosome A3 and A8. The DAPs that may be related to clubroot resistance are discussed in detail below.

# Calcium Signaling in Resistance to *Plasmodiophora brassicae*

When plants are challenged by pathogens, activation of multilayered intra- and extracellular signaling pathways takes place and defense responses are triggered (Tuteja and Mahajan, 2007). Calcium (Ca<sup>2+</sup>) ion concentration alters rapidly and dynamically in the cytosol in response to environmental or pathogen stimuli and plays a vital role as a second messenger, and communicates with other signal transduction pathways to elicit cellular responses (Zhang et al., 2014). Our current study identified 33 DAPs that are orthologous to various intracellular Ca<sup>2+</sup> sensors, such as calreticulin, annexin, calmodulin (CaM), calmodulin-like (CMLs), and Ca<sup>2+</sup> dependent protein kinases (**Figure** 7 and **Supplementary Table 8A**). We observed that the orthologs of sensor relay (CMLs) proteins (e.g., BnaA03T0464700WE, BnaC04T0461900WE,

BnaC07T0227600WE, and BnaC08T0316200WE) consistently increased in the CR-progeny lines in response to the pathogen. This is similar to CMLs gene expression in Arabidopsis in response to Pseudomonas syringae, where the ortholog was rapidly induced under pathogen infection and its overexpression resulted in the hypersensitive response (Chiasson et al., 2005). Similarly, calcium dependent protein kinase (CPK; BnaA03T0374600WE) was increased in CRparent and progeny lines and decreased in CS- parent and progeny indicating that the increase in accumulation in the CR lines could potentially be involved in the resistance phenotype. Similar observations have been made, albeit at the transcript level, where the CPK transcript was rapidly induced in response to fungal elicitors and its overexpression resulted in resistance to the pathogen (Coca and San Segundo, 2010).

In support of a potential role for calcium signaling proteins in mediating responses to *P. brassicae*, GO functional categories of response to - biotic (GO:0009607), and calcium ion binding (GO:0005509) were highly enriched. GO terms such as calcium ion binding (GO:0005509) is involved in the regulation of  $Ca^{2+}$  concentration in cytosol and  $Ca^{2+}$  sensor activity (Priest et al., 2014). Eight DAPs were associated with the term (GO:0005509) exhibiting a shared pattern between the CR-parent and progeny lines, with an increase in protein abundance as the infection progressed to 21 DPI (**Supplementary Table 8B**). Thus, the accumulation of calcium signaling-related proteins was strongly affected by *P. brassicae*, suggesting their involvement in mediating defense response against the pathogen.

# Reactive Oxygen Species in Response to *Plasmodiophora brassicae*

While excessive production of ROS in a cell can be detrimental, small and sub-toxic levels of ROS generation is essential for signaling purposes in order to maintain cellular processes, including plant-pathogen responses (Huang et al., 2019). Putative proteins related to diverse ROS-scavenging systems, such as peroxidases, ascorbate peroxidase, superoxide dismutase, glutathione, and thioredoxin have been reported to be differentially accumulated, potentially to maintain ROS homeostasis in cells and to limit the ROS-dependent damage (Mittler et al., 2004; Liu J. et al., 2021). Furthermore, increase in accumulation of some of the proteins, such as BnaA05T0420500WE (orthologous to peroxiredoxin-2F) and BnaA10T0260700WE (orthologous to l-ascorbate peroxidase 5) in the CR-progeny and a corresponding decrease in the CS-progeny, related to ROS could also suggest the involvement of ROS-dependent signaling pathways in clubroot resistance. For instance, GO enriched categories such as response to oxidative stress (GO:0006979), oxidoreductase (GO:0016491), antioxidant activity (GO:0016209), and acting on peroxide as acceptor (GO:0016684) were some of the highly enriched categories observed in both the genotypes. Furthermore, it has also been reported that ROS generation is linked to the formation of defensive barriers against biotrophic fungus in barley and inhibiting the pathogen colonization (Hückelhoven and Kogel,

2003). While ROS production in the cell was associated with the defense responses, some necrotrophic pathogens such as Botrytis cinerea promote ROS accumulation and triggers cell death aiding in the proliferation of the pathogen (Govrin and Levine, 2000). Earlier studies have demonstrated that overexpression of ROS related gene, l-ascorbate peroxidase 6, in Nicotiana benthamiana improved resistance against Pseudomonas solanacearum and Fusarium solani var. coeruleum in sugarcane (Liu et al., 2018). Similarly, the overexpression of ascorbate peroxidase lines in rice exhibited increased tolerance to Xanthomonas orvzae (Jiang et al., 2016). In both the cases, the genes were associated with ROS scavenging. However, to our knowledge, there are no reports of similar studies in the Brassica-P. brassicae pathosystem. Collectively, our results imply that the differential accumulation of ROS-related proteins may be the result of a complex network of signals against the pathogen attack and may be viable targets for genetic manipulation in order to afford durable resistance to P. brassicae in canola.

## Phytohormones and Immunity Responses to *Plasmodiophora brassicae*

Phytohormones play a vital role in responding to environmental stresses. In our study, proteins related to several phytohormones were detected including auxin, CKs, ET, JA, and SA. There are many reports of changes to endogenous phytohormone levels in plant roots during the development of clubroot infection (Ludwig-Müller et al., 2009). Primarily, SA and JA/ET are regulated synchronously to respond to the invading pathogen (Verma et al., 2016). SA plays a fundamental role in activating both systemic and local resistance in response to biotrophic pathogens (White, 1979; Vlot et al., 2009) while JA/ET is vital for defense against necrotrophic invaders (Glazebrook, 2005). Additionally, auxin and CKs, which are primarily involved in the regulation of cell division and cell elongation, are manipulated by P. brassicae leading to the hypertrophy of infected root cells (Robin et al., 2020). Our current study identified three auxin related proteins (BnaA10T0208100WE, BnaA05T0409700WE, and BnaC03T0302500WE) in which the putative protein orthologous to auxin efflux carrier component 5 (BnaA10T0208100WE) increased in the CR progeny and decreased in the CS progeny root proteomes as the infection progressed to 21 DPI (Figure 7). There are reports in the literature that the genes related to auxin efflux carrier was upregulated in the later stages of clubroot infection at 14-, 28-, and 35-DPI (Robin et al., 2020). In our study, at 21 DPI, one putative protein, BnaA06T0322500WE, orthologous to histidine kinase (Nishimura et al., 2004), was increased in the CR and decreased in the CS root proteomes of both parent and progeny lines (Supplementary Table 7B). Although it is unclear why these proteins are decreasing in the CS line as compared to CR line, it is a clear indication that they are important in mediating responses to clubroot pathogen. Additional studies are underway in our laboratory to further characterize the roles of these proteins in clubroot response/resistance.

NPR1 is one of the key positive regulators of SA dependent pathway and it has been demonstrated that it is important for the systemic acquired resistance in *Arabidopsis* (Cao et al., 1997). Overexpression of NPR1 in *B. juncea* conferred resistance against *Alternaria brassicae* and *Erysiphe cruciferarum* pathogen (Ali et al., 2017). Several studies in *Arabidopsis* and rice, have reported that accumulation of NPR1 conferred disease resistance against various pathogens (Fu and Dong, 2013; Kumar et al., 2013; Molla et al., 2016). Therefore, NPR1 related proteins exhibiting a contrasting pattern in abundance, especially those with an increase in CR root proteome and a decrease in CS, such as Bnascaffold286T0028200WE and BnaC07T0044900WE, may be potential candidates for further investigation into their role(s) in mediating resistance to clubroot.

Along with SA, ET, and JA also play crucial roles in the activation of plant defense pathways (Glazebrook, 2005). We have identified three proteins (BnaA03T0361400WE, BnaA09T0013600WE, and BnaC04T0098000WE) that are orthologous to *s*-adenosylmethionine (SAM) synthase, which increased in CR root proteomes of both parent and progeny lines in response to the pathogen. SAM is a precursor for the synthesis of ET, and the overexpression of SAM synthase related gene resulted in enhanced tolerance of Arabidopsis to oxidative stress (Ma et al., 2017). Moreover, earlier studies have demonstrated that, transcripts related to ET biosynthesis were induced in response to biotic stress (Vilanova et al., 2017; Yang et al., 2017). These proteins, especially the ones that increased in CR root proteomes and decreased in the CS may be appropriate candidates for further investigation.

# Dehydration Induced Proteins in Response to *Plasmodiophora brassicae*

Dehydrins are stress proteins and accumulate during abiotic stress and protect cells from dehydration (Kiyosue et al., 1994; Kovacs et al., 2008; Liu et al., 2017). Although their mechanisms of action are poorly understood, the accumulation of dehydrins in plants have been reported during salinity, drought, and low temperature stresses (Kovacs et al., 2008; Hanin et al., 2011). In this study, several dehydrin proteins that were observed to be decreased (BnaC02T0286500WE, and BnaA07T0239200WE) in the CS lines and some proteins (BnaA07T0354800WE, BnaC06T0291300WE, and BnaC06T0439800WE) were increased in the CS root proteome in response to the pathogen (Figure 7 and Supplementary Table 11A). Earlier studies have suggested that the expression of dehydrins may confer tolerance to desiccation stress in plants by acting as chaperones, thereby stabilizing cellular membranes by sequestering electrolytes, ions, buffering water molecules, and lowering electrolyte leakage (Rorat, 2006; Tunnacliffe and Wise, 2007; Graether and Boddington, 2014).

Microscopic observation of the disease progression in this study, especially as the pathogen progresses to secondary plasmodia phase, confirmed severe tissue damage presumably due to uncontrolled cell division and accumulation of callus around the vascular cambium. The proliferation of vascular cambium results in a concomitant inhibition of xylem development (Malinowski et al., 2012). Due to suppression of xylogenesis, plants experience dehydration due to poor water and mineral conduction (Malinowski et al., 2012) as evidenced by the wilting and stunting of plants at later stages of disease. Infected plants may trigger the expression of dehydrin genes leading to increase in dehydrin abundance in order to cope with the dehydration stress. Similarly, three proteins (BnaA08T0045300WE, BnaA10T0060800WE, and BnaC02T0364000WE) orthologous to other drought related genes, including cysteine proteinase RD21a (responsive to desiccation 21) (Yamaguchi-Shinozaki et al., 1992) were identified and all three proteins were increased in CR and CS lines; of note, the protein accumulation was relatively higher in the CS lines than in the CR lines (Supplementary Table 11A) supporting the hypothesis that clubroot infection results in a water-deficit stress in canola. To our knowledge, this is the first report of dehydrins being regulated in response to the clubroot pathogen. As alluded to above, our current data suggests that clubroot infection results in water deficit stress in canola plants mimicking drought stress as the xylogenesis is strongly repressed and uptake of water and mineral is obstructed in the CS lines. Additionally, it is reasonable to speculate that the increase in abundance of specific dehydrins (BnaC02T0286500WE and BnaA07T0239200WE) in the CR lines may be a contributing factor for the observed resistance in these lines and these proteins may be appropriate candidates for further investigation.

## Lignin Related Proteins in Plant Defense

Lignin, a phenolic polymer, forms a physical barrier between plant cell and pathogens and it is non-degradable by most microorganisms (Moura et al., 2010). Often a reinforcement of cell walls with lignin occurs when host encounters microbial pathogens (Moura et al., 2010; Vanholme et al., 2010). In this study, we identified 24 putative proteins that were orthologous to different types of peroxidases and proteins involved in lignin biosynthesis (Figure 7). For instance, the enzyme METK3 is involved in lignin biosynthesis (Shen et al., 2002) and there were four putative orthologs (BnaC03T0264500WE, BnaA09T0013600WE, BnaA 03T0361400WE, and BnaC04T0098000WE), of which, BnaA03T0361400WE (located in Chromosome A3) was increased consistently in the CR and decreased in CS lines especially when the infection progressed to 21 DPI. Similarly, one of the proteins orthologous to caffeic acid 3-O-methyltransferase (BnaA03T0142300WE, located at chromosome A3 genome) demonstrated shared pattern in protein accumulation in CR and CS lines in the case of both parent and progeny (Supplementary Table 9A). Similar to our results Zhang et al. (2007) reported an increase in peroxidase activity in Chinese cabbage in response to Erwinia carotovora. Increase in lignification has also been observed in Pinus nigra in response to Sphaeropsis sapinea infection (Bonello and Blodgett, 2003). There are also multiple reports in the literature describing an increase in the expression of lignin biosynthetic genes and lignin levels upon pathogen infection (Bhuiyan et al., 2009; Miedes et al., 2014; Summanwar et al., 2021). Over expression of lignin biosynthesis related genes such as BnaC03T0264500WE could potentially deposit lignin that could form a non-degradable barrier, which could restrict pathogen penetration of plant cells, thereby reducing disease infection. Others have reported that the overexpression of cinnamoyl-coA reductase 2, a rate limiting enzyme in lignin synthesis pathway, in *B. napus* resulted in increased lignin accumulation and a reduction in *Sclerotinia scletotiorum* disease symptoms (Liu D. et al., 2021). A similar approach could be viable for the development of CR canola.

## Clubroot Resistant Related Loci in Chromosome A3 and A8

Clubroot resistance-related genes have been reported on both A and C genomes of Brassica spp. A total of 24 CR loci have been reported, where 11 and five loci were reported from the chromosome A3 and A8, respectively; and the remaining were associated with other chromosomes (Hasan et al., 2021). In this study, we have identified 73 putative proteins from eight possible CR loci in Brassica that were reported earlier (Hirai et al., 2004; Saito et al., 2006; Chu et al., 2014; Hasan and Rahman, 2016; Yu et al., 2017; Laila et al., 2019; Zhu et al., 2019). Six proteins (BnaA02T0335400WE related to Rcr8; BnaA06T0336500WE-, BnaC06T0195100WE-, BnaA03T0374600WE-, BnaA03T0262200WErelated to Rcr4, and BnaA03T0464700WE- related to Rcr1) increased in abundance and are potentially involved in plant and microbe interactions (Supplementary Table 8). The latter (BnaA03T0374600WE, BnaA03T0262200WE, three and BnaA03T0464700WE) are involved in calcium signaling, and the first three BnaA02T0335400WE, BnaA06T0336500WE, and BnaC06T0195100WE are related to aquaporin TIP1, cysteine rich receptor, and HSP70 protein, respectively (Supplementary Table 15). Putative proteins linked to Rcr4 are located at the chromosomal location between 23823130 26806601 bp in chromosome A3 of Brara\_Chiifu\_V3.0 genome, and the proteins linked to Rcr1 and Rcr8 are located at the chromosomal position 24574673\_25715499 bp in A3 and 22502031\_26348249 bp in A2, respectively. The locus labeled as Rcr4 on A03 was demonstrated to confer resistance against P. brassicae pathotypes 2, 3, 5, 6, and 8 (Yu et al., 2017). Likewise, the locus designated as Rcr8 on A2, and Rcr1 on A3 conferred resistance against pathotype 5X and 3, respectively (Chu et al., 2014; Yu et al., 2017). Many of the reported loci have been documented as major loci, QTL or genes, the putative proteins from our dataset were based on the sequence similarity with the specific loci and they will be suitable candidates for further investigation on their biological roles especially in the context of CR. They may also be suitable candidates for the design of gene-specific molecular markers to assist with the selection of CR lines.

## CONCLUSION

Our results have contributed to a better understanding of the protein level changes in *B. napus* in response to clubroot causing obligate biotrophic protist pathogen, *P. brassicae* and

has identified major proteins and metabolic pathways potentially involved in clubroot resistance. Our results indicate that calcium signaling pathway is involved in intra- and extracellular flow of information during the pathogen detection and response against it and indicates that the accumulation of calcium ions enhances tolerance potential of host against biotic stress. Additionally, with the detection of important ROS related proteins, our results suggest that there could be a crosstalk between calcium and ROS in mediating a response to the clubroot pathogen. A range of metabolic pathways and host defense related proteins involving distinctive biological processes and molecular functions were either increased in the CR line indicating their potential role in the resistance mechanism or decreased in the CS line signifying the susceptible trait in the disease development. The former is supported by the higher abundance of lignin related proteins, ROS related peroxidases along with glycosyl hydrolases, PRRs, and phytohormone related proteins. This study has identified 74 differentially accumulated proteins that were linked to 7 possible clubroot resistance loci in four chromosomes (A2, A3, A7, and A8) and several of them were directly related to plant and pathogen interaction including calcium signaling and ROS related proteins. Many of the genes encoding the aforementioned proteins could be viable candidates for enhancing clubroot resistance in canola either through genetic manipulation or through classical breeding.

## DATA AVAILABILITY STATEMENT

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found below: Proteomics raw data, Xcalibur parameters, and Spectronaut search settings have been deposited to the ProteomeXchange Consortium via the PRIDE partner repository with a dataset identifier PXD031070.

## **AUTHOR CONTRIBUTIONS**

NK and HR designed the project with input from RGU on the proteomics component. NK supervised the project, interpreted the results, and edited multiple versions of the manuscript. DA designed, with the input of HR and NK, and conducted all the experiments, analyzed the results, and wrote the manuscript. DM and RGU performed the LC-MS/MS analysis and wrote the corresponding materials and methods. HR provided all plant material used in this study and input into draft version of this manuscript. All authors contributed to the article and approved the submitted version.

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## SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fpls.2022. 860393/full#supplementary-material

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