Insights from the proteome profile of *Phytophthora capsici* in response to the novel fungicide SYP-14288

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**ABSTRACT**

*Phytophthora capsica* is a destructive oomycete plant pathogen that causes huge losses to crop production worldwide. However, the novel fungicide SYP-14288 has shown excellent activity against various stages of the oomycete life cycle as well against fungal plant pathogens. The current study utilized isobaric tags for relative and absolute quantitation technology to generate proteome profiles of *P. capsici* in the presence or absence of SYP-14288 in order to gain a greater understanding of the SYP-14288 mode of action. A total of 1,443 individual proteins were identified during the investigation, of which 599 were considered to have significantly altered expression. Further investigation using Cluster of Orthologous Groups of proteins analysis and Kyoto Encyclopedia of Genes and Genomes pathway analysis indicated most of the proteins with altered expression were associated with carbohydrate metabolism, energy metabolism and their downstream biological functions, especially with regard to oxidoreductase activity and subsequent adenosine triphosphate (ATP) production associated pathways. Quantitative expression analysis using qRT-PCR validated the proteomic data. These results seem to indicate that SYP-14288 treatment caused a shift in energy metabolism that resulted in the activation of compensatory mechanisms affecting carbohydrate and lipid metabolism. The study also found evidence that the up-regulation of transmembrane transporters and proteins associated with stress response might also be coopted to compensate for the disrupted proton gradient and other downstream effects. Taken together these results provide strong evidence that SYP-14288 has a similar mode of action to the oxidative phosphorylation uncoupler Fluazinam but further investigation, including molecular studies, is required to completely characterize the SYP-14288 mode of action in *P. capsici*. However, the proteomic data collected in the current study does provide important insight into the overall effect of SYP-14288 in *P. capsici*, which could be useful for the registration and application of this novel fungicide.

**INTRODUCTION**

The oomycete *Phytophthora capsici* Leonian is a destructive pathogen, which causes crown, root and fruit rot, in a wide range of solanaceous and cucurbit crops leading to...
significant economic losses throughout the world every year (Lamour et al., 2012). Indeed, its economic impact is so great that it is currently listed as one of the top 10 most important oomycete pathogens in plant pathology (Kamoun et al., 2015). Although genetic modification of susceptible species and preventative management practices have improved the control of diseases caused by Phytophthora capsici, the application of fungicides is still a critical component of integrated control programs (Ristaino & Johnston, 1999).

The novel fungicide SYP-14288, which was developed by Shenyang Research Institute of Chemical Industry of China, belongs to the diarylamine family, and shares a similar chemical structure with Fluazinam (Fig. S1), which is reported to be an uncoupler of mitochondrial oxidative phosphorylation (Guo et al., 1991). Like other diarylamine fungicides, SYP-14288 exhibits broad-spectrum fungicidal activity that provides excellent control of many plant diseases including Colletotrichum orbiculare and cucumber downy mildew Pseudoperonospora cubensis, as well as outstanding control of the pepper blight caused by Phytophthora capsici. Although SYP-14288 works as an uncoupler to control various plant pathogens, its exact target site(s) remains unclear (Wang et al., 2018). Given the great potential of SYP-14288 within the fungicide market, it is important to characterize its mode of action in greater detail.

The completion of the Phytophthora capsici genome sequencing project has provided the opportunity for new analytical techniques to be applied to the study of fungicide mode of action in this important pathogen including proteomics, which has been widely applied for the toxicological evaluation of novel chemicals, and especially target site discovery in recent years (Bruneau et al., 2003; Hughes et al., 2006; Bantscheff et al., 2007). For example, two-dimensional gel electrophoresis has been successfully used to investigate the response of Phytophthora cactorum to zoxamide (Mei et al., 2014), while MALDI-TOF-MS/MS has been used to explore the mode of action of the novel fungicide JS399-19 in Fusarium graminearum (Hou et al., 2013). Similarly, the isobaric tags for relative and absolute quantitation (iTRAQ) method has been used to study the effect of pyrimorph in Phytophthora capsici (Pang et al., 2015). Proteomic methods provide important information that is difficult to obtain using traditional biochemical studies and have therefore become popular tools to study the mode of action of novel compounds.

The current study utilized a quantitative proteomics approach based on iTRAQ technology to investigate the changes in protein expression that occur in Phytophthora capsici exposed to SYP-14288 in order to gain a global perspective of the biochemical pathways affected by SYP-14288, as well as to identify specific target sites that could be validated by qRT-PCR. It is hoped that the results of the study will provide an expanded understanding of the SYP-14288 mode of action as well as critical data that can be utilized for the design of novel fungicides against a broad range of plant diseases.

**MATERIALS AND METHODS**

**Sensitivity of Phytophthora capsici to SYP-14288**

Mycelial growth inhibition method as previously described in Cai et al. (2016) was used to determine the sensitivity of Phytophthora capsici in response to treatment with SYP-14288. Six Phytophthora capsici isolates collected from peppers of different breeding fields in
China were evaluated. All the isolates were cultured in darkness on potato dextrose agar plates (200 g potato, 20 g Glucose, 15 g agar, distilled water to one L) containing SYP-14288 (kindly provided by Shenyang chemical research institute) at the following concentrations: 0, 0.3, 0.6, 1, 5, 10, 50 µg/mL, respectively. Dimethyl sulfoxide was used as a solvent control. The plates were then incubated at 25 °C for 3 days before the diameter of each colony was measured at two perpendicular angles. Each treatment consisted of four replicate plates and the entire experiment conducted three times. The half maximal effective concentration (EC$_{50}$) was then calculated using a regression equation (Supplementary Information) derived by correlating the log of the SYP-14288 concentration and the probit of percentage inhibition based on the average radial growth of the *Phytophthora capsici* isolates compared to the control treatment.

**Protein extraction from *Phytophthora capsici***

According to the previously described method in Pang *et al.* (2015), three g of mycelial samples harvested from the wild type isolate HD3 were incubated in 80 mL PDB medium amended with one µg/mL SYP-14288 (the EC$_{50}$ determined above). Equivalent control samples were prepared in the absence of SYP-14288. The liquid cultures were then placed in dark-incubation at 25 °C for 24 h in a shaker (220 rpm) before the mycelia were harvested by centrifugation. The resulting mycelia were washed and dried under vacuum before being stored at −80 °C until required. A total of 10 separate cultures were used for each treatment and the entire experiment conducted once.

**Preparation of protein hydrolysate**

The frozen mycelia from each treatment were ground under liquid nitrogen in a ball mill (MM400; Retsch, Haan, Germany) using two mL tubes and five mm sterile steal balls, before two volumes of cold (4 °C) lysis buffer (8 M urea, 30 mM HEPES, one mM PMSF, two mM EDTA, and 10 mM dithiothreitol (DTT)) were added to each tube. The samples were then subjected to low temperature sonication (Sonics Vibra-Cell VCX-500 Ultrasonic Processor; Artisan, Champaign, IL, USA) (pulse on 2 s, pulse off 3 s, power 180 W) for 5 min to enhance cell lysis, before centrifugation at 20,000×g for 30 min. The supernatant containing the protein fraction was then transferred to new tubes, while the pellet containing undisrupted cells was discarded. DTT was added to each sample to a final concentration of 10 mM before incubation at 56 °C. After 1 h iodoacetamide was added to a final concentration of 55 mM before a second period of dark-incubation (1 h). The protein was then precipitated by the addition of four volumes of cold acetone buffer and incubation at −20 °C for 3 h before being collected by centrifugation at 20,000×g for 20 min at 4 °C. The resulting protein samples were then washed in cold acetone to remove any residual DTT before being dissolved in 500 μL resuspension buffer (50% w/v tetraethylammonium bromide, 0.1% SDS). The concentration of the protein samples was measured using the Bradford dye-binding assay (Bradford, 1976), and an equal amount of protein from each of the 10 replicates pooled together to produces 100 µg samples for each treatment. The resulting samples were then hydrolyzed by one µg/µL trypsin at 37 °C overnight and dried in vacuum before being analyzed.
iTRAQ labeling
The peptides from the two treatments (SYP-14288 and negative control) were labeled using the iTRAQ Reagent-8 Plex Multiplex kit (Applied Biosystems, Waltham, MA, USA) according to the protocol of the manufacturer. Different iTRAQ tags were used for the two treatments. The labeled peptides were then dried under a speed-vac before being assessed by chromatography.

Segregation of peptides by chromatography
The peptide mixtures were separated according to the reported protocol in Pang et al. (2015). The samples were first dissolved in 100 µL buffer A (10 mM KH₂PO₄, pH 3.0; acetonitrile ACN/H₂O 25/75, v/v) before flowing through a 100 × 4.6 mm fractionation column (Luna®; Phenomenex, Inc., Torrance, CA, USA). The proteins were then eluted using a gradient of two buffers: 100% buffer A for 10 min, 0–30% buffer B (10 mM KH₂PO₄, pH 3.0; 2 M KCl, ACN/H₂O 25/75 v/v) for 15 min, 30–100% buffer B for 15 min and 100% buffer B for 10 min at a flow rate of one mL/min for 1 h. The UV absorbance of the eluents was detected at 214 nm. A total of 16 fractions were acquired and dried before being desalted using a C18 column (Strata-X; Phenomenex, Inc., Torrance, CA, USA).

Analysis of peptides by LC–MS/MS
Peptide fractions were analyzed according to the previously described proteomics study in Pang et al. (2015). In brief, the LC–MS/MS system consisting of the Thermo Fisher Scientific Easy-nLC nanoflow chromatograph and the Thermo Fisher Scientific Q Exactive™ Mass Spectrometer was used. The samples were fractionated in a 100 × 75 mm C18 column (PepMap™; Thermo Fisher Scientific, Waltham, MA, USA) at a flow rate of 400 nL/min. The MS spectra in the range of 350–2,000 m/z were acquired in the positive ion reflection mode using a data-dependent Top 10 method which dynamically selected the most abundant precursor ions from the survey scan (300–1,650 Th) for higher energy collisional dissociation fragmentation (Michalski et al., 2011).

Database searches for peptide and protein identification
The raw MS/MS data were converted into the .MGF file format using the Proteome Discoverer 1.3 software (Thermo Fisher Scientific, Waltham, MA, USA), before the exported files were used to search the NCBI Phytophthora Database employing the automatic decoy database search function of Mascot 2.3 (Matrix Science, London, UK). Several parameters in Mascot were optimized for peptide searching, including iTRAQ 8-plex for quantification, tolerance of one missed cleavage of trypsin, carbamidomethylation for cysteine as a fixed modification, and oxidation for methionine as a variable modification. The precursor mass tolerance was set to 15 ppm and the mass tolerance to 20 mmu. A concatenated target-decoy database-search strategy was used to check the rate of false positives, which were found to be less than 1% in all cases. High-confidence proteins which contained at least two unique peptides with a protein score greater than 1.3 were selected for further quantification.
Quantitative data analysis of iTRAQ labeled peptides

The iTRAQ quantification was determined based on the reporter ion intensities detected from Proteome Discoverer version 1.3 (Thermo Fisher Scientific, Waltham, MA, USA). The iTRAQ channels were normalized by summing all the reporter ion intensities in each iTRAQ channel and equalizing each channel contribution by dividing individual reporter ion intensities by the corresponding channel-specific correction factor. The quantitative data was analyzed using statistical procedures previously described in Ross et al. (2004), where the log2-transformation for the iTRAQ ratios was first performed to yield a normal distribution. Proteins were considered to be differentially expressed if they displayed at least 1.5-fold change in abundance (±) in response to SYP-14288 treatment, and therefore were selected as proteins of interest in the subsequent Kyoto Encyclopedia of Genes and Genomes pathway (KEGG) analysis. The KEGG analysis was performed according to the formula described previously (Liu et al., 2012), and the pathways with a P-value <0.05 were considered significantly “enriched” to be correlated with the altered expression of proteins in response to SYP-14288. A Cluster of Orthologous Groups of proteins (COG) analysis was further performed according to the previously described method in Ye et al. (2006).

Total RNA extraction and qRT-PCR validation of differentially expressed proteins

The qRT-PCR analysis of selected Phytophthora capsici genes belonging to different KEGG pathways were performed using Total RNA extracted from mycelial samples prepared according to the protocol described above regarding protein samples. The RNA was extracted using a TRIzol method, and first strand cDNA synthesized using the TransScript™ Two-Step RT-PCR SuperMix kit (Transgen, Beijing, China). The qRT-PCR analysis was performed using SYBR Green II fluorescent dye and the ABI PRISM 7500 Real-Time PCR system (Applied BioSystems, Foster City, CA, USA). The 20 µL reaction mixtures contained 10 µL of SYBR Premix DimerEraser (2×), 0.4 µL of ROX Reference Dye Dye II (50×conc.), two µL of template DNA, 0.6 µL of each of the forward and reverse primers (10 µM), and 6.4 µL of ddH2O, and processed according to the following program: an initial melting at 95 °C for 30 s; followed by 40 cycles of 95 °C for 5 s, 57.3 °C for 30 s, and 72 °C for 34 s, with a final extension at 95 °C for 30 s. The melting curves were obtained using a standard protocol, and used to confirm that the target products did not contain primer dimers after the amplification had been completed. Two housekeeping genes, WS21 and UBC, were selected as the internal controls, and their geometric mean Ct values used to determine the uniformity of the cDNA concentrations (Yan & Liou, 2006). Relative expression values were then calculated according to the 2−ΔΔCT method (Livak & Schmittgen, 2001). Three biological replicates were performed for each of the two treatments using fresh cultures for the RNA extraction, while three technical replicates were performed for each biological replicate during the qRT-PCR itself. Statistical analysis for the qPCR data (P = 0.05) was carried out as described previously (Yuan et al., 2006). The gene-specific primers used during the qRT-PCR analysis are listed in Table S1.
RESULTS

Inhibitory effect of SYP-14288 on the mycelial growth of Phytophthora capsici

Treatment with SYP-14288 was found to inhibit the mycelial growth of all the test isolates in a dose-dependent manner (Fig. 1). The EC$_{50}$ of SYP-14288 varied, but was found to be approximately one µg/mL in isolate HD3, the isolate subsequently selected for the proteomic analysis. Previous studies have shown that a reduction of 50–70% during the exponential growth phase is ideal for proteomic analysis of fungicide effects (Bandow et al., 2003; Mei et al., 2014), while prolonged exposure can induce non-specific effects such as cell lysis, which can complicate the interpretation of proteomic data. Consequently the proteomic analysis was performed on samples collected from isolate HD3 after 24 h exposure to one µg/mL SYP-14288 (Mei et al., 2014; Pang et al., 2015).

Overview of the quantitative proteomics analysis

A total of 1,443 proteins were identified during the iTRAQ–LC–MS/MS analysis of the two proteome libraries (Table S2). The majority of the proteins had a molecular mass in excess of 10 kDa, although the mass of individual proteins varied greatly, ranging from 5.5 to 726.6 kDa (Fig. S2).

Changes in functional pathways in response to SYP-14288

The KEGG analysis indicated that as many as 186 pathways were affected by SYP-14288 treatment, including six primary pathways and 39 secondary pathways. However, only 18 specific pathways belonging to 10 different secondary pathways and three primary pathways were found to be significant ($P < 0.05$). These pathways included selenoamino acid metabolism, fatty acid metabolism, and the citrate cycle, and mostly belonged to the metabolism category, with only one pathway being allocated to the cellular processes category, and one to the organismal systems category (Table 1). The KEGG analysis indicated that the 10 secondary pathways affected by SYP-14288 were associated with the differential expression of 405 proteins, 126 of which were up-regulated and 279 of which
were down-regulated. However, the majority of the proteins affected belonged to just five secondary pathways including carbohydrate metabolism, energy metabolism, metabolism of other amino acids, amino acid metabolism, transport and catabolism, which were associated with 162, 86, 42, 30 and 30 proteins, respectively (Fig. 2).

### Changes in the expression of individual proteins in response to SYP-14288

A total of 599 individual proteins were found to have altered expression in response to SYP-14288, when a 1.5-fold change in abundance was used as a threshold for differential expression. Of these, 291 were found to be up-regulated, while 308 were down-regulated. Furthermore, 40 unique proteins were identified that were only expressed in Phytophthora capsici isolate HD3 in the presence of SYP-14288.

The subsequent COG analysis revealed that the proteins exhibiting differential expression were primarily associated with eight functional groups reflecting different molecular functions (Fig. 3). The oxidoreductase activity category contained the highest number of proteins, 78 in total, which was almost double in comparison to the other categories including protein binding, nucleic acid binding, ATP binding, metal ion binding, cofactor binding, transferase activity (transferring phosphorus-containing groups) and pyrophosphatase activity, which contained 50, 48, 43, 42, 41, 40 and 35 proteins, respectively. The COG analysis also indicated that most of the proteins affected

| Pathway                                           | P-value | Pathway ID | Functional category                  | Secondary category          | Primary category          |
|---------------------------------------------------|---------|------------|--------------------------------------|-----------------------------|---------------------------|
| Selenoamino acid metabolism                       | 0.003   | ko00450    | Metabolism of other amino acids      | Metabolism                  |                           |
| Fatty acid metabolism                             | 0.005   | ko00071    | Lipid metabolism                     | Metabolism                  |                           |
| Citrate cycle (TCA cycle)                         | 0.006   | ko00020    | Carbohydrate metabolism              | Metabolism                  |                           |
| NOD-like receptor signaling pathway               | 0.010   | ko04621    | Immune system                        | Organismal systems          |                           |
| Pyruvate metabolism                               | 0.013   | ko00620    | Carbohydrate metabolism              | Metabolism                  |                           |
| Carbon fixation in prokaryotes                    | 0.013   | ko00720    | Energy metabolism                    | Metabolism                  |                           |
| Beta-alanine metabolism                           | 0.013   | ko00410    | Metabolism of other amino acids      | Metabolism                  |                           |
| Propanoate metabolism                             | 0.016   | ko00640    | Carbohydrate metabolism              | Metabolism                  |                           |
| Cysteine and methionine metabolism                | 0.019   | ko00270    | Amino acid metabolism                | Metabolism                  |                           |
| Histidine metabolism                              | 0.020   | ko00340    | Amino acid metabolism                | Metabolism                  |                           |
| Lysosome                                          | 0.021   | ko04142    | Transport and catabolism             | Cellular processes          |                           |
| Methane metabolism                                | 0.021   | ko00680    | Energy metabolism                    | Metabolism                  |                           |
| Taurine and hypotaurine metabolism                | 0.028   | ko00430    | Metabolism of other amino acids      | Metabolism                  |                           |
| Limonene and pinene degradation                   | 0.028   | ko00903    | Metabolism of terpenoids and polyketides | Metabolism                  |                           |
| Sulfur metabolism                                 | 0.031   | ko00920    | Sulfur metabolism                    | Metabolism                  |                           |
| Carbon fixation in photosynthetic organisms       | 0.035   | ko00710    | Energy metabolism                    | Metabolism                  |                           |
| Glycolysis/Gluconeogenesis                        | 0.042   | ko00010    | Carbohydrate metabolism              | Metabolism                  |                           |
| Chloroalkane and chloroalkene degradation         | 0.047   | ko00625    | Xenobiotics biodegradation and metabolism | Metabolism                  |                           |
by SYP-14288 exposure belonged to five primary functional categories including ATP binding, pyrophosphatase activity, oxidoreductase activity, transferase activity (transferring phosphorus-containing groups) and metal ion binding which mediated the changes occurring in a multitude of downstream pathways, thus, these should be the original pathways of the effect of SYP-14288 application. However, it should be noted that individual proteins can be assigned to more than one category during COG analysis (Fig. 4).
Phytophthora capsici proteins associated with the SYP-14288 mode of action and stress response

Given that SYP-14288 has a similar chemical structure to the oxidative phosphorylation uncoupler Fluazinam, and likely shares the same mode of action, the results of the metabolic pathway enrichment analysis, which assigns metabolic function of differentially expressed proteins by matching them to previously annotated proteins in the KEGG pathway database, was used to investigate proton producing and transporting proteins and their interactions.
those associated with generalized stress responses (Table 2). The analysis revealed that many of the proteins associated with oxidative phosphorylation had altered expression in response to SYP-14288 were associated with proton production as well as proton-motive force formation and ultimately impacted the production of ATP. Other proteins associated with proton translocation and supplemental energy production, including ATP-binding Cassette (ABC) transporters and fatty acid metabolism were also affected. Similarly the heat shock proteins associated with energy release to protect organisms under stress were
also found to be affected. These results provide strong evidence that SYP-14288 affects the energy metabolism of *Phytophthora capsici* and has a mode of action similar to Fluazinam.

**Gene expression analyses for differentially expressed proteins**

A total of 20 proteins belonging to different metabolic pathways associated with the effect of SYP-14288 were selected to validate the results of the proteomic analysis using qRT-PCR. The results confirmed that 16 of the 20 genes had altered trends of expression consistent with the proteomic data, which implied the proteomic data was reliable (Fig. 5). According to the statistical analysis, eight of them differed significantly \( (P < 0.05) \). Besides, four genes including ATPase, ATP phosphoribosyltransferase, heat shock protein 90 and a putative acyl-CoA dehydrogenase had different altered trends with proteomic data, but the difference were not huge (with protein/gene levels at 1.957/0.941, 0.475/2.602, 1.576/0.297 and 1.341/0.621, respectively).

**DISCUSSION**

Proteomic analysis is a powerful tool to determine the mode of action of fungicides by identifying proteins and metabolic pathways that have altered patterns of expression in response to exogenous compounds (*Vido et al., 2001; Teixeira et al., 2005; Bantscheff et al., 2007; Santos, Simoes & Sa-Correia, 2009*). The current study employed an iTRAQ LC–MS/MS
based approach to produce quantitative proteome profiles to characterize the response of *Phytophthora capsici* to the novel fungicide SYP-14288. A total of 1,443 proteins were identified in the control and EC$_{50}$ treatment libraries. The subsequent gene ontology analysis categorized these proteins into 186 biological processes, 18 of which had significantly altered expression in the fungicide treatment, including metabolic pathways such as carbohydrate and amino acid metabolism. Although only 18 pathways were affected, a total of 599 individual proteins were found to have significantly altered expression (change fold $\geq 1.5$), the majority of which were associated with carbohydrate and energy metabolism. The subsequent COG analysis categorized the proteins into eight groups including oxidoreductase activity, protein binding and nucleic acid binding. Several of the proteins identified by the COG and KEGG pathway analysis were selected for validation using qRT-PCR, which primarily focused on proteins associated with oxidative phosphorylation, ATP consumption, proton transportation and responses to stress that would all be affected if SYP-14288 had a similar mode of action as the fungicide Fluazinam. For the most part, the results of the qRT-PCR were consistent with those of the iTRAQ analysis, with most of the genes having similar patterns of expression, which indicated that the proteomic analysis was convincing and reliable. However, there were a few disparities between the proteome and RNA data, which most likely resulted from a time lag between transcription and the accumulation of the resulting protein.

Many of the proteins (86 in total) that had altered expression in response to SYP-14288 were associated with energy metabolism, with many being significantly up-regulated. For example, the ATP synthase subunit beta, which is a component of the ATP synthase complex that catalyzes ATP synthesis by utilizing the electrochemical gradient of protons across the inner membrane of the mitochondrion produced during oxidative phosphorylation (Wang & Oster, 1998; Elston, Wang & Oster, 1998; Tyler et al., 2006; Leyva, Blanchet & Amzel, 2009). The vacuolar proton translocating ATPase A subunit and an uncharacterized ATPase (GI: 262108152) were also found to be highly up-regulated. It is well established that ATPase enzymes can function as pumps (including proton pumps) and as co-transporters, catalyzing ATP hydrolysis during the process to provide metabolic energy (Martin & Senior, 1980; Njus, Knoth & Zallakian, 1981). Previous research has shown that the up-regulation of these proteins and the supplemental energy produced by their activity can play an important role in drug resistance and during severe oxidative stress (Milgrom et al., 2007). The up-regulation of these two proteins appears to corroborate the hypothesis that SYP-14288 acts as an uncoupler of oxidative phosphorylation or as an ATP biosynthesis inhibitor similar to Fluazinam. The current study also found that several proteins associated with the oxidative phosphorylation pathway itself were significantly down-regulated, including cytochrome c oxidase subunit 3 (also known as complex IV of respiratory electron transport chain) and NADH dehydrogenase flavoprotein 2, which are both involved in proton transfer and the establishment of the transmembrane proton gradient in mitochondria that provides the electrochemical potential for the ATP synthase (Hochstein & Dalton, 1973; Tsukihara et al., 1995; Balsa et al., 2012). The down-regulation of these proteins suggests that SYP-14288 may cause a high proton gradient between the inner membrane of the
mitochondrion that can inhibit the formation of proton pumps on the outer surface of the inner membrane via a negative feedback mechanism. In addition, it was interesting to note that the NADH dehydrogenase subunit 7 was significantly up-regulated (change fold = 1.505) in response to SYP-14288. This protein, which is one of the main sources of superoxide and is responsible for the translocation four protons across the mitochondrial membrane, has previously been reported to be the primary target of the uncoupler Fluazinam (Lee et al., 2012), which can be taken as further evidence that the two fungicides have shared a mode of action.

Many of the proteins with altered expression were found to be involved in carbohydrate metabolism. The most down-regulated protein in this category was pyruvate phosphate dikinase, which is responsible for phosphoenolpyruvate biosynthesis, a process that requires ATP (Evans & Wood, 1968; Pocalyko et al., 1990; Mei et al., 2014), the down regulation of which could have been caused be a reduction in ATP as a result of SYP-14288 treatment. In contrast, several glycolytic enzymes, including glycoside hydrolases that assists in the hydrolysis of glycosidic bonds in complex sugars (Sinnott, 1990; Davidson et al., 2006), were up-regulated. For example, glyceraldehyde-3-phosphate dehydrogenase, which catalyzes the sixth step in glycolysis (Tarze et al., 2007), was highly up-regulated (change fold = 2.275). The increased expression of enzymes associated with sugar metabolism enzymes could be another response to increase energy generation to compensate for the disrupted proton gradient caused by the putative uncoupling effect of SYP-14288. The activation of glucose consumption is not only a means to produce essential energy but has also been associated with mechanisms that respond to metabolic stress (Owen et al., 2008). The results of the current study also indicated that many downstream proteins associated with the processing of acetyl-CoA in citric acid cycle, pyruvic acid metabolism and the pentose phosphate pathway were down-regulated. It is possible that these changes occurred in response to reduced acetyl-CoA levels, which could imply that SYP-14288 has an effect on enzymes that catalyze the conversion of pyruvic acid to acetyl-CoA increasing the production of ATP via the citric acid cycle, another mechanism that could compensate for the reduced levels of ATP caused by uncoupler activity.

Aside from changes that directly affected energy metabolism, the current study also found that SYP-14288 changed the expression of proteins involved in amino acid and lipid metabolism including aminotransferase, which catalyzes the interconversion of certain amino acids and α-keto acid (Grenville-Briggs et al., 2005); argininosuccinate lyase (change fold = 0.521), which is involved in arginine biosynthesis (Knudsen, Burton & Færgeman, 2003); translation elongation factor 1 alpha, which is involved in polypeptide elongation (Van’t Klooster et al., 2000); and the hypothetical protein PHYSODRAFT-485399; which all had reduced levels of expression. The negative impact of SYP-14288 on such a wide range of proteins associated with amino acid metabolism indicates that it has a significant effect on protein synthesis in Phytophthora capsici. Similar results have also been observed in the response of Phytophthora capsici to Pyrimorph (Pang et al., 2015). In contrast, almost all of the proteins involved in lipid metabolism were up-regulated including Acyl-CoA dehydrogenase, which catalyzes the initial step in each cycle of fatty acid β-oxidation resulting in the introduction of a trans double-bond between C2 and C3.
of the acyl-CoA thioester substrate (Thorpe & Kim, 1995); and a member of the enoyl-CoA hydratase/isomerase family (change fold = 2.388), which is involved in fatty acid beta-oxidation (Müller-Newen, Janssen & Stoffel, 1995). The up-regulation of proteins associated with lipid metabolism, which can produce large amounts of energy, indicates that SYP-14288 cause a dramatic redistribution of metabolic processes associated with energy production. Indeed, it is interesting to note that this result demonstrates an increase in the production of acetyl-CoA associated with up-regulation of lipid metabolism, while the other findings of the current study (detailed above) suggest levels of acetyl-CoA associated with carbohydrate metabolism were reduced, which again indicates a compensatory mechanism in response to the putative uncoupling activity of SYP-14288.

Some of the proteins with altered expression were allocated to functional categories completely unrelated to energy metabolism. For example, several proteins belonging to the ABC Superfamily were found to be significantly up-regulated. Previous studies have shown that ABC transporters, which transfer various substrates across membranes, have been implicated in the detoxification of fungicides by excretion and the response to environmental stress (Van Den Brule et al., 2002; Jones & George, 2004). Given the strong evidence presented above that SYP-14288 act as an uncoupler of oxidative phosphorylation, it is possible that the up-regulation of ABC proteins could be an attempt to excrete SYP-14288 in order to maintain a proton gradient in the mitochondria of Phytophthora capsici and finally offset the toxicity of the fungicide. It was also found that elicitin, which is associated with the induction of hypersensitive cell death and other biochemical changes associated with the defense response of plants (Kamoun et al., 1993), was significantly up-regulated. This implies that signal transduction is an essential factor in the response of Phytophthora capsici to SYP-14288. The current study also found that an inorganic phosphate transporter was highly up-regulated. Given that phosphate enters the mitochondria via a phosphate transporter in exchange for OH\(^-\) or by co-transport with H\(^+\) (Karandashov et al., 2004), it is possible that the increased levels of this transporter could be another compensatory mechanism to address the disrupted proton gradient and reduced energy production resulting from SYP-14288 treatment. Several heat shock proteins, which are known to be involved in generalized stress responses (De Maio, 1999), were also found to be up-regulated. In this case it is possible the altered expression was induced in order to respond to the thermolytic requirements of Phytophthora capsici if its energy dissipative pathways were inhibited by SYP-14288. Although none of the proteins discussed in this section are directly linked to energy metabolism, they are consistent with indirect consequences resulting from the disruption of oxidative phosphorylation, and therefore provide further evidence that SYP-14288 has a similar mode of action as the uncoupler Fluazinam.

Some functional groups of protein were affected dramatically under SYP-14288 treatment. The pyrophosphatases, which could hydrolyse ATP and produce phosphate (Clair et al., 1997), may cause the activation of extra phosphate transfer. Acceleration of respiratory rate caused by activation of oxidoreductases could be a complementary mechanism of ATP shortage. And some ABC proteins and metal ion binding proteins could acted as transmembrane transporters to transfer protons crossing the inner membrane of mitochondria thus compensate for the disrupted proton gradient. All these
functional groups may be the primary step of SYP-14288 function and then caused the downstream intricate biological reaction.

In summary, the results of the current study found extensive evidence that SYP-14288 has a wide ranging effect on the metabolism of *Phytophthora capsici* involving a multitude of different biochemical pathways. However, the study also indicated that the primary effect of SYP-14288 was a disruption of the proton gradient within the mitochondria by perturbing proton transporter or altering membrane permeability (*Guo et al., 1991*), and that most of the other changes were downstream responses to compensate for the reduced ATP of both the ATP synthetase and ATPase being significantly up-regulated, as well as other proteins associated with energy metabolism (especially those associated with respiration) in response to the altered proton motive force. This shift in energy metabolism resulted in the activation of compensatory mechanisms affecting carbohydrate and lipid metabolism as well as a reduction in amino acid biosynthesis. The study also found evidence that the up-regulation of transmembrane transporters and proteins associated with stress response might also cooperate to compensate for the disrupted proton gradient and other downstream effects. However, despite characterizing the metabolic pathways affected by SYP-14288 and possible target sites, the proteomic analysis failed to identify a specific target protein. Further investigation, including affinity chromatography assay, are therefore required to completely characterize the SYP-14288 mode of action in *Phytophthora capsici*. However, the proteomic data collected in the current study does provide important insight into the overall effect of SYP-14288 in *Phytophthora capsici*, which could be useful for the registration and application of this novel fungicide.

**CONCLUSION**

The novel fungicide SYP-14288 has shown excellent activity against various plant pathogens including *Phytophthora capsici*, which is the causal agent of blight on hundreds of plants. The current study utilized iTRAQ to gain a greater understanding of the mode of action of SYP-14288. A total of 599 out of 1,443 proteins were considered to have significantly altered expression; most of these proteins were associated with carbohydrate metabolism, energy metabolism and their downstream biological functions, especially oxidoreductase activity and subsequent ATP production associated pathways. All results of the current study seem to indicate that SYP-14288 causes a shift in energy metabolism that results in the activation of compensatory mechanisms affecting carbohydrate and lipid metabolisms. All the data strongly implies that SYP-14288 works as an oxidative phosphorylation uncoupler. However, further investigation is still needed to completely characterize the mode of action of SYP-14288.

**ADDITIONAL INFORMATION AND DECLARATIONS**

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The authors declare that they have no competing interests.

Author Contributions
- Meng Cai conceived and designed the experiments, performed the experiments, analyzed the data, prepared figures and/or tables, authored or reviewed drafts of the paper.
- Zhiwen Wang conceived and designed the experiments, performed the experiments, analyzed the data, prepared figures and/or tables, authored or reviewed drafts of the paper.
- Xiaoxia Ni performed the experiments.
- Yanhua Hou performed the experiments.
- Qin Peng analyzed the data, contributed reagents/materials/analysis tools.
- Xiang Gao contributed reagents/materials/analysis tools, he revised this paper draft.
- Xili Liu conceived and designed the experiments, authored or reviewed drafts of the paper, approved the final draft.

Data Availability
The following information was supplied regarding data availability:
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