The Powdery Mildew Effector CSEP0027 Interacts With Barley Catalase to Regulate Host Immunity

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Powdery mildew is one of the most important fungal pathogen diseases. The genome of barley mildew fungus, Blumeria graminis f. sp. hordei (Bgh), encodes a large number of candidate secreted effector proteins (CSEPs). So far, the function and mechanism of most CSEPs remain largely unknown. Here, we identify a Bgh effector CSEP0027, a member of family 41, triggering cell death in Nicotiana benthamiana. CSEP0027 contains a functional signal peptide (SP), verified by yeast secretion assay. We show that CSEP0027 promotes Bgh virulence in barley infection using transient gene expression and host-induced gene silencing (HIGS). Barley catalase HvCAT1 is identified as a CSEP0027 interactor by yeast two-hybrid (Y2H) screening, and the interaction is verified in yeast, in vitro and in vivo. The coexpression of CSEP0027 and HvCAT1 in barley cells results in altered localization of HvCAT1 from the peroxisome to the nucleus. Barley stripe mosaic virus (BSMV)-silencing and transiently-induced gene silencing (TIGS) assays reveal that HvCAT1 is required for barley immunity against Bgh. We propose that CSEP0027 interacts with barley HvCAT1 to regulate the host immunity and likely reactive oxygen species (ROS) homeostasis to promote fungal virulence during barley infection.

Keywords: powdery mildew, Blumeria graminis, effector, CSEP, virulence, barley catalase

INTRODUCTION

Powdery mildews are widespread fungal diseases that affect more than 10,000 plant species, such as important cereal crops, economic, and ornamental plants (Glawe, 2008; Dean et al., 2012; Takamatsu, 2013). As obligate biotrophic pathogens, powdery mildew fungi totally depend on the living plant cells for survival and reproduction. Mildew conidiospores attach to the epidermal tissue of the host, germinate and produce fungal infection structures, such as the appressorium and penetration peg to penetrate the plant cell wall, subsequently, the haustoria are developed within the lumen of the host cells but separated from the host cell cytoplasm by extrahaustorial membrane (EHM) and extrahaustoral matrix (EHMX) (Panstruga, 2003; Both et al., 2005). The haustorium is believed to be a site for nutrient uptake and signaling exchange (Panstruga and Dodds, 2009;
Stergiopoulos and de Wit, 2009), and effector proteins are believed to deliver into the plant cells through haustorium to promote fungal virulence.

*Blumeria graminis*, the powdery mildew fungus causing disease on the cereal crop species and grasses (Poaceae), has been classified into at least eight formae speciales (f.sp.), each adapted to a host genus (Troch et al., 2014). *B. graminis* f.sp. *hordei* (Bgh) and *B. graminis* f.sp. *triticii* (Bgt) colonize barley and wheat, respectively. The *Bgh* and *Bgt* genomes code for ∼700 and 800 candidate secreted effector proteins (CSEPs), respectively (Godfrey et al., 2010; Spanu et al., 2010; Pedersen et al., 2012; Wicker et al., 2013; Frantzeskakis et al., 2018; Müller et al., 2019). Many *Bgh* CSEPs are overlapped with the so called *Blumeria* effector candidate (BEC) proteins identified from the proteomic analyses (Bindschedler et al., 2009, 2016; Godfrey et al., 2009). A majority of these *Bgh* CSEPs have a predicted amino-terminal signal peptide (SP) and a putative Y/F/WxC motif (Bindschedler et al., 2009; Godfrey et al., 2010; Spanu et al., 2010; Pedersen et al., 2012). A large proportion of *Bgh* CSEPs (c. 25%) are structurally predicted similar to RNase and/or RNA-binding activity, and these CSEPs are termed as RNase Like Proteins expressed in Haustoria (RALPHs) (Pedersen et al., 2012; Spanu, 2017). Interestingly, most of the so far identified *Bgh* AVR2 effectors, each recognized by a cognate barley MLA receptor, are also RALPHs with fungal RNase folds but lacking the residues required for RNase activity (Lu et al., 2016; Saur et al., 2019; Bauer et al., 2021). So far, several *Bgh* CSEPs/BECs have been functionally characterized with respect to fungal virulence through transient gene expression and host-induced gene silencing (HIGS) approaches (Bindschedler et al., 2009; Godfrey et al., 2009; Nowara et al., 2010; Spanu et al., 2010; Pedersen et al., 2012; Pliego et al., 2013; Ahmed et al., 2015, 2016; Menardo et al., 2017; Frantzeskakis et al., 2018; Pennington et al., 2019; Li et al., 2021). The host targets have been identified for some CSEPs that are involved in plant immunity and stress responses (Zhang et al., 2012; Schmidt et al., 2014; Ahmed et al., 2015; Pennington et al., 2016, 2019; Saur et al., 2019). Recently, few *Bgh* CSEPs have been showed or proposed to play a role in regulating the host cell death (Pennington et al., 2019; Li et al., 2021). A CSEP0064/BEC1054, one of the *Bgh* RALPHs, binds to RNA and may act as a pseudoenzyme to inhibit the action of the host ribosome-inactivating proteins (RIPs) that would otherwise induce cell death (Pennington et al., 2019). The CSEP0139 and CSEP0182 are capable of suppressing programmed cell death (PCD) induced by various cell death inducers in plant cells (Li et al., 2021). Despite these intensive studies, the function and mode of action of many CSEPs remain largely unclear.

Reactive oxygen species (ROS), produced from aerobic metabolism in plants, have been appreciated as major signaling molecules in plant development and in response to the biotic and abiotic stresses (Apel and Hirt, 2004; Nanda et al., 2010; Waszczak et al., 2018). In plant–pathogen interactions, ROS can directly kill pathogens or can serve as signaling molecules to regulate the plant defense responses (Mittler et al., 2011; Mittler, 2017). Hydrogen peroxide (H$_2$O$_2$) and superoxide anion (O$_2^-$) are the two major ROS molecules accumulating in the plants in response to the pathogen infections. The plants rely on an intricate network to control the levels of ROS at different subcellular compartments (Hückelhoven and Kogel, 2003; Nanda et al., 2010; Petrov and Van Breusegem, 2012). Catalases are part of “the ROS network,” playing a central role in maintaining the cellular H$_2$O$_2$ balance and in signaling crosstalk (Du et al., 2008; Chaouch et al., 2010; Nanda et al., 2010; Sharma and Ahmad, 2014; Li et al., 2015; Zhang et al., 2015; Murota et al., 2017; Yuan et al., 2017; Chen and Jarosz, 2020; Chen et al., 2020).

In barley/wheat response to *B. graminis* infection, ROS are involved in immune responses at early and late stages of the pathogen infections (Hückelhoven and Kogel, 2003). In barley under attack by *Bgh* or *Bgt* spores, H$_2$O$_2$ is detected to locally accumulate in papillae (cell wall appositions) or in the whole cell, which is generally associated with host cell inaccessibility (Thordal-Christensen et al., 1997; Hückelhoven et al., 1999, 2001, 2003). The ROS are also detected in *Bgt*-attacked wheat epidermal cells and are involved in both pattern-triggered immunity (PTI) and effector-triggered immunity (ETI; Altpeter et al., 2005; Schweizer, 2008; Chang et al., 2019). On the other hand, superoxide radical anion (O$_2^-$) is believed to act in restricting cell death. In barley epidermal cells under attack by *Bgh* spores, O$_2^-$ accumulation is strictly associated with a successful penetration and O$_2^-$ also accumulates in the living cells neighboring the HR cells (Hückelhoven and Kogel, 1998; Hückelhoven et al., 2000). These studies suggest that ROS play a complex role in the plant–biotrophic fungal interactions, not only in early cell wall-associated defense and in late defense signaling but also in the cell-death suppression.

In this study, we screen ∼100 *Bgh* CSEPs through agroinfiltration in *Nicotiana benthamiana* and identify CSEP0027 triggering cell death. We show that CSEP0027 promotes fungal virulence in barley infection. We further identify CSEP0027 interactors by yeast two-hybrid (Y2H) screening and barley HvCAT1 is shown to interact with CSEP0027 in yeast, in vitro and in vivo. Coexpression of CSEP0027 and HvCAT1 in barley cells induces the nuclear accumulation of HvCAT1 that is normally localized to the peroxisome. The functional analyses indicate that HvCAT1 is involved in barley immunity against *Bgh*. We propose CSEP0027 target barley HvCAT1 to regulate host immunity and promote fungal virulence in barley infection.

**RESULTS**

CSEP0027 Specifically Induces Cell Death in *N. benthamiana*

The *Bgh* genome encodes several hundreds of potential effectors, and ∼491 effector-like proteins were initially identified to be CSEPs (Spanu et al., 2010; Pedersen et al., 2012). We selected a hundred of these *CSEP* genes for further characterization based on their expression levels and abundance in haustoria (Godfrey et al., 2010; Pedersen et al., 2012). The cDNA sequences of 101 CSEPs from 34 families were amplified with specific primers using RNA samples derived from barley leaf materials infected with the compatible isolate *Bgh*A6 (*Supplementary Table 1*). All CSEP cDNA sequences excluding the predicted
signal peptide (ASP) were subcloned into vector pGR107 for Agrobacterium tumefaciens-mediated transient expression in N. benthamiana (Wang et al., 2011). We identified several CSEPs suppressing cell death in plants (Li et al., 2021), but much fewer CSEPs inducing cell death. As shown in Figure 1A, CSEP0027 is one of the CSEPs inducing clear water-soaked-like cell death phenotype in N. benthamiana, as compared to GFP alone, which serves as a negative control. The AVRr13 effector and its cognate receptor MLA13 were also coexpressed and triggered cell death in N. benthamiana (Lu et al., 2016), which severely as a positive and technique control here (Supplementary Figure 1). Trypan blue staining confirmed the localized cell death and immunoblotting verified the expression of the HA-tagged fusion proteins (Figure 1A), and DAB (3,3′-diaminobenzidine) staining also revealed H$_2$O$_2$ accumulation in the infiltrated area (Supplementary Figure 2).

The CSEP0027, CSEP0028, and CSEP0340 are the three members from the same Bgh CSEP family 41 (Pedersen et al., 2012), in addition, BgtE-10117 and BgtE-20000 are the two potential Bgt homologs being identified as highly related sequences to CSEP0027 (Supplementary Figure 3A) (Praz et al., 2017). All these five CSEPs harbor a predicted SP, a Y/FxX motif, and a conserved C-terminal cysteine, with some conserved residues in the middle (Figure 1C and Supplementary Figure 3A). We tested if any of the other four CSEPs trigger cell death, unexpectedly none of them induced cell death in N. benthamiana (Figure 1B and Supplementary Figure 3B). CSEP0027, thus represents a unique Bgh effector protein to induce cell death in N. benthamiana.

**CSEP0027 Is a Secreted Protein**

To validate the secretory function of the CSEP0027 signal peptide, we used a yeast secretion assay based on invertase secretion and yeast growth on sucrose or raffinose media (Lee et al., 2006; Oh et al., 2009). The predicted SPs were fused in frame to the mature sequence of invertase in the vector pSUC2 and expressed in the invertase mutant yeast strain YTK2 that otherwise cannot grow on YPRAA medium (Gu et al., 2011). CSEP0027-SP derived construct enabled transformed yeast cells that otherwise cannot grow on YPRAA medium (Gu et al., 2011). To further analyze the expression pattern of CSEP0027, we conducted a time course experiment (Figure 3C). The transcript level of CSEP0027 remained low from 0 to 12 hpi and was markedly induced at 24 and 48 hpi in both the haustorial containing samples (H) and epiphytic structures (E), with highly enriched transcripts in H sample but not in E sample at 48 hpi (Figure 3C). This expression pattern supports CSEP0027 functioning during barley infection and likely at the post-penetration stages.

**CSEP0027 Interacts With Barley Catalase HvCAT1**

To identify host targets of CSEP0027, we performed a Y2H screening of a cDNA prey library derived from Bgh infected barley leaves. Using a bait of CSEP0027 without the SP, we identified two independent clones harboring the fragments of a barley catalase gene, HvCAT1. The targeted Y2H analysis showed that CSEP0027 interacted with full-length HvCAT1 but not with HvCAT2 (Figure 4A), another reported barley catalase that shares more than 70% amino acid identity with HvCAT1 (Supplementary Figure 4; Skadsen et al., 1995). Further interaction analysis indicated that HvCAT1 interacts with CSEP0027 likely through the N-terminal catalase domain but not the C-terminal domain (Supplementary Figure 5).

The interaction between CSEP0027 and HvCAT1 was further verified by in vitro and in vivo assays (Figures 4B–D). For glutathione S-transferase (GST) pull-down assay, GST-CSEP0027 fusion or GST alone derived from E. coli was incubated with HvCAT1-HA containing crude lysate of N. benthamiana. An immunoblotting analysis indicated that GST-CSEP0027 pulled down HvCAT1-HA whereas GST did not (Figure 4B). In luciferase complementation imaging (LCI) assays, CSEP0027-nLuc interacted with cLuc-HvCAT1, thus generated luminescence signal, the reciprocal pair HvCAT1-nLuc and cLuc-CSEP0027 also generated strong luminescence signal in N. benthamiana (Figures 2, 4), while two pairs of negative control did not produce any detectable signal (Figures 1, 3, 4C). In addition, the HvCAT2-nLuc and cLuc-CSEP0027 did not generate detectable signal (Figures 4C, 5). In co-immunoprecipitation (co-IP) analysis, the HvCAT1-Flag fusion did immuno-precipitate with CSEP0027-HA in N. benthamiana, whereas GFP-Flag did not (Figure 4D).

Together, these results indicate that CSEP0027 specifically interacts with barley HvCAT1.
FIGURE 1 | CSEP0027 triggers cell death in Nicotiana benthamiana. (A) Expression of CSEP0027 triggers cell death in N. benthamiana. Agrobacterium tumefaciens was used to transiently express CSEP0027 or GFP in N. benthamiana leaves. The picture was taken at 5 dpi, and cell death was visualized by trypan blue staining. The numbers in each circled area indicate numbers of cell death in total number of leaf areas infiltrated with the construct. Total protein extract was obtained from N. benthamiana leaves at 60 hpi and protein expressions were confirmed by immunoblotting using anti-HA antibody. Ponceau staining was used to show equal loading. (B) CSEP0028 and CSEP0340 do not trigger cell death in N. benthamiana. The experimental procedures are the same as in (A). The stars in the Western blots in panels (A,B) indicate non-specific signals. (C) Sequence alignment of CSEP0027, CSEP0028, and CSEP0340, performed using the DNAMAN software. The signal peptides are highlighted in red box, Y/FxC motif in blue box, and C-terminal conserved cysteine in green box.

FIGURE 2 | CSEP0027 is a secreted protein. Yeast invertase secretion assay was used to confirm the function of the predicted SP of CSEP0027. A construct expressing a fusion of SP sequence of CSEP0027 and a yeast invertase was transformed into the yeast strain YTK12 and tested in the assay, with the N-terminal sequence of Magnaporthe oryzae Mg87 protein and SP sequence of Phytophthora sojae PsAvr1b used as negative and positive controls, respectively. CMD-W plates were used to select yeast strain YTK12 carrying the pSUC2 vector. YPRAA media were used to indicate invertase secretion. An enzymatic activity test based on the reduction of 2,3,5-triphenyltetrazolium chloride (TTC) to red-colored formazan was also used to confirm invertase secretion.
CSEP0027 Induces the Nuclear Localization of HvCAT1

Since CSEP0027 interacts with HvCAT1, we examined the subcellular localization of CSEP0027 and catalases in barley cells. The plasmids expressing CSEP0027^{ΔSP}·CFP (Cyan Fluorescent Protein), YFP (Yellow Fluorescent Protein)-HvCAT1, and YFP-HvCAT2 fusions were constructed and delivered into barley cells by particle bombardment. Confocal imaging indicated that CSEP0027^{ΔSP}·CFP was localized in both cytosol and nucleus, similar to YFP alone (Figure 5, the top panels), while YFP-HvCAT1 was localized in many small dots in the cytoplasm, totally different from that of CFP alone (Figure 5, 2nd panels). Since many plant catalases are localized to peroxisomes, we tested the localization of YFP-HvCAT1 in peroxisomes by coexpression of YFP-HvCAT1 with a peroxisomal marker, PST1-RFP (Red Fluorescent Protein). As expected, YFP-HvCAT1 was almost fully co-localized with PST1-RFP in many cytoplasmic foci in the same cells (Figure 5, 3rd panels). Interestingly, YFP-HvCAT2 was also co-localized with PST1-RFP in most of the cytoplasmic dots (Figure 5, 4th panels). Next, we tested the localization of CSEP0027^{ΔSP}·CFP and YFP-HvCAT1 in barley cells by coexpression analysis. Remarkably, confocal imaging indicated that YFP-HvCAT1 was detected not only in the peroxisomal dots but also in the nucleus, and CSEP0027^{ΔSP}·CFP appeared to co-localize with YFP-HvCAT1 in some of the cytoplasmic dots but fully overlapped with YFP-HvCAT1 in the nucleus (Figure 5, 5th panels). Interestingly, when YFP-HvCAT2 was coexpressed with CSEP0027^{ΔSP}·CFP in barley cells, YFP-HvCAT2 remained to localize in the peroxisomal dots and some dots appeared to overlap with CSEP0027^{ΔSP}·CFP in the cytoplasm (Figure 5, the bottom panels). These localization analyses suggest that HvCAT1 and CSEP0027 have overlapped subcellular localization in the cytosol and CSEP0027 specifically induces the nuclear localization of HvCAT1.

HvCAT1 Is Involved in Barley Immunity

The plant catalases play an important role in biotic stress responses by regulating ROS signaling and homeostasis (Du et al., 2008; Chaouch et al., 2010; Sharma and Ahmad, 2014). To evaluate the function of HvCAT1 in barley immunity, we knocked down the HvCAT1 expression through barley stripe mosaic virus vector (BSMV)-mediated virus-induced gene silencing (VGIS) approach followed by the inoculation of a compatible Bgh isolate. An antisense fragment of HvCAT1 used efficiently silenced HvCAT1 but not HvCAT2 (Figure 6A and Supplementary Figure 3). Scoring of Bgh microcolony formation rate (i.e., microcolony index, MI%) in barley leaf cells at 60–72 hpi indicated that the relative MI% increased by ~30% in the HvCAT1-silenced leaves as compared to the EV control (Figure 6B). Staining of the Bgh infected barley leaves showed more microcolonies and better hyphae growth on the leaf surface of HvCAT1-silenced barley, as compared with the EV control (Figure 6C). Furthermore, transiently-induced gene silencing (TIGS) technique was used to silence HvCAT1 in barley leaf epidermal cells (Himmelbach et al., 2007; Bai et al., 2012). The RNAi-HvCAT1 construct was delivered into the barley cells by particle bombardment followed by Bgh spores inoculation. Relative haustorium formation rate (i.e., relative haustorium index, HI%) scored at 48 hpi almost significantly increased by ~50% as compared with the EV control (Figure 6D). By contrast, TIGS-silencing of the barley Mlo, a gene required for full susceptibility to Bgh (Kusch and Panstruga, 2017), drastically reduced Bgh HI% in barley cells by ~80% (Figure 6D). Together, these data indicate that HvCAT1 is involved in barley immunity against Bgh.
FIGURE 4 | CSEP0027 specifically interacts with barley catalase HvCAT1. (A) Yeast two-hybrid (Y2H) assay shows CSEP0027-HvCAT1 interaction. Yeast was transformed with indicated bait and prey constructs. Serial dilutions from cell suspension of yeast expressing bait and prey constructs are shown. Growth on SD-Trp-Leu plates indicates yeast clones carrying the bait and prey constructs. The interactions were detected as yeast growth on SD-Trp-Leu-Ade-His plates. (B) Glutathione S-transferase (GST) pull-down assay confirms CSEP0027-HvCAT1 interaction. HvCAT1-HA was extracted from N. benthamiana leaves at 2 dpi, while GST-CSEP0027 and GST alone were purified from E. coli. GST pull-down fractions were detected by immunoblotting using anti-HA antibody and by Ponceau staining. (C) LCI assay confirms CSEP0027-HvCAT1 interaction. The N- or C-terminal fragment of LUC (nLuc or cLuc) was fused with indicated proteins. Indicated fusion pairs were coexpressed in N. benthamiana by agroinfiltration. The luminescent signal was collected at 48 hpi with a charge-coupled device (CCD) imaging apparatus. (D) Co-immunoprecipitation (Co-IP) analysis validates CSEP0027 and HvCAT1 interaction. HvCAT1-Flag or GFP-Flag was transiently coexpressed with CSEP0027-HA in N. benthamiana. The crude proteins were extracted at 48 hpi and subjected to Co-IP analysis.

DISCUSSION
The genomes of many filamentous plant pathogens interacting biotrophically with plants encode hundreds of predicted effectors, and yet loss of function of some individual effectors can have measurable effect on fungal virulence. B. graminis fungi also encode several hundreds of CSEPs, and it is expected that many of the CSEPs contribute to the obligate biotrophy life style of the B. graminis fungi, for example, co-survival with the host cells or tissues. It is thus of particular interests to understand the functions and mechanisms of CSEPs in regulating host immune responses and cell-death related processes. Here, we identify Bgh CSEP0027 that triggers cell death when heterologously expressed in N. benthamiana. Importantly, CSEP0027 promotes fungal virulence in barley and interacts with HvCAT1 that is involved in host immunity, most likely, in the maintenance of ROS homeostasis in host cells. In this study, the primary aim in ectopically expressing the Bgh CSEPs in N. benthamiana is to identify those who may have cell-death related functions, either suppressing or inducing cell death, hoping to better understand the biotrophic lifestyle of the Bgh fungus. Indeed, we have predominately identified CSEPs suppressing cell death in N. benthamiana (Li et al., 2021), but unexpectedly, CSEP0027 triggering cell death as shown in the present study. We speculate that this cell death activity of CSEP0027 and related pathway may not be fully conserved in dicots and monocots. For example, the co-receptors BAK1 and SOBIR1 are important immune signaling components required for PTI and cell death in dicots (Liu et al., 2016; van der Burgh et al., 2019), while whether the co-receptors are also required for CSEP0027-induced cell death in N. benthamiana is not yet resolved here, and importantly, whether the signaling pathway for CSEP0027 induced cell death is shared between N. benthamiana and barley awaits for further investigation. Nevertheless, this study findings suggest that B. graminis fungi may utilize the CSEPs to target host catalase, a likely component of host ROS networks, presumably to manipulate the ROS homeostasis and signaling for the benefit of the pathogens.

CSEP0027 Functioning in Fungal Virulence
The well-established HIGS technique has been used for identifying Bgh CSEPs with functions in promoting fungal virulence (Nowara et al., 2010). So far, a few dozens of Bgh CSEPs have been shown to contribute to Bgh pathogenicity (Nowara et al., 2010; Zhang et al., 2012, 2019; Pliego et al., 2013; Aguilar et al., 2015; Ahmed et al., 2015, 2016; Pennington et al., 2019;
Li et al., 2021). In the present study, HIGS of CSEP0027 led to the reduction of HI% by ∼37% in the infected barley cells. Together with the transient overexpression results, our data support the role of CSEP0027 in promoting fungal virulence during barley infection. Our data also suggest that CSEP0027 is most likely a cytoplasmic effector and HvCAT1 is one of its virulence targets. By affecting the subcellular localization of HvCAT1, CSEP0027 may facilitate Bgh infection of host barley.

Bgh CSEP genes are usually induced and/or differentially expressed during the infection of barley. Some CSEP genes are induced at early stages of barley infection, for example, from ∼6 to 12 hpi, whereas others are induced at later stages from 24 to 48 hpi (Godfrey et al., 2009; Zhang et al., 2012; Hackenberg et al., 2013; Schmidt et al., 2014; Aguilar et al., 2015; Ahmed et al., 2015, 2016). CSEP0027 is induced from 24 to 48 hpi and is enriched in fungal haustoria (Figure 3B). We thus believe CSEP0027 functions at later stages of infection, most likely during and after haustorium formation.

Regulation of ROS Signaling and Homeostasis

Reactive oxygen species, as major regulatory and signaling molecules, can be generated in different subcellular compartments of plant cells and are regulated by an array of antioxidant systems (Waszczak et al., 2018). During plant–fungus interaction, one of the early events in plant response to fungal penetration is an oxidative burst in the apoplastic space, generated mainly by the phagocyte respiratory burst oxidase homologous nicotinamide adenine dinucleotide phosphate (NADPH) oxidases, cell wall peroxidases, and oxalate oxidases (Hückelhoven, 2007; Lehmann et al., 2015). In barley/wheat and B. graminis interactions, H$_2$O$_2$ and some other ROS molecules are generated in plant cells during the early stages of fungal penetration, participating in the cell wall lignification and apposition (Zhang et al., 1995; Thordal-Christensen et al., 1997; Hückelhoven et al., 1999; Hückelhoven et al., 2001, 2003; Hückelhoven, 2007; Schweizer, 2008; Li et al., 2015). Interestingly, Bgh fungus also secretes an extracellular catalase that may function in H$_2$O$_2$ scavenging in the apoplastic space of host cells (Zhang et al., 2004). The catalases have been known as a class of ROS scavenging enzymes catalyzing the conversion of H$_2$O$_2$ into H$_2$O and O$_2$, thereby regulating the homeostasis of the intracellular ROS level (Mhamdi et al., 2010; Sharma and Ahmad, 2014). ROS homeostasis is maintained in a very complex manner, involving different ROS-scavenging enzymes, such as catalases, ascorbate peroxidases, glutathione, superoxide dismutases (Mittler et al., 2004; Torres et al., 2006). The peroxisomal ROS levels are closely regulated by CAT activity, and in Arabidopsis, the primary peroxisomal H$_2$O$_2$ scavenger is CAT2 (Mhamdi et al., 2012). Here, we show that barley HvCAT1 and HvCAT2 are also peroxisomal catalases. It is unclear whether and when these two HvCAT1 and HvCAT2 are involved in the H$_2$O$_2$ decomposition and signaling in peroxisomes during barley interaction with Bgh fungus. Since CSEP0027 expression is induced and most likely functions post haustorium formation, we speculate that HvCAT1 may play a role in regulating ROS homeostasis at later stages, e.g., during and post haustorium formation. Our preliminary data suggest that CSEP0027 triggered-cell death involves H$_2$O$_2$ accumulation in N. benthamiana, however, it is not clear if the expression CSEP0027 induced disturbance of ROS homeostasis thus cell death, or vice versa. On the other hand, it is also not yet clear if CSEP0027 has activity in cell death during barley interaction.
with \textit{Bgh} fungus. Undoubtedly, more data are needed for fully understanding the role of CSEP0027 in interacting with barley catalases and in regulating ROS homeostasis during barley interaction with \textit{Bgh} fungus, particularly, the cell death signaling pathway that might be a primary target of the biotrophic fungal pathogen.

The current data are in line with the notion that ROS, in particular cellular \(H_2O_2\), may play an important role in the barley interactions with the \textit{B. graminis} fungi. It is not unexpected that peroxisomal ROS signaling/homeostasis and ROS signaling cross-talk among the organelles are integral and important parts of barley defense responses to the biotrophic \textit{Bgh} fungal pathogen.

The Regulation of Plant Catalases
Apart from being regulated at transcriptional level, plant catalases are also regulated at post-translational level (Mhamdi et al., 2010). A variety of plant proteins have been reported to affect the activity and stability of plant catalases (Yang and Poovaiah, 2002; Fukamatsu et al., 2003; Verslues et al., 2007; Li et al., 2013, 2015; Zou et al., 2015; Kneeshaw et al., 2017). In addition, some pathogen secreted proteins are also identified to interact with the plant catalases and affect their activity, stability, and subcellular localization (Inaba et al., 2011; Mathioudakis et al., 2013; Zhang et al., 2015; Murota et al., 2017; Sun et al., 2017). In line with these examples, the current study data provide new evidence that biotrophic fungal pathogen also secretes an effector to target and affect host catalase subcellular localization in plants.

The plant catalases are mostly peroxisomal proteins and imported into the peroxisome matrix via the peroxisomal targeting signal 1 (PTS1) pathway, i.e., relying on the C-terminal tripeptide PTS1 signal to interact with a peroxisomal receptor and translocate into the peroxisome (Gatto et al., 2000; Lanyon-Hogg et al., 2010). Barley \textit{HvCAT1} and \textit{HvCAT2}, each contains a typical PTS1 signal, PNM or PSM, respectively (Supplementary Figure 4; Mhamdi et al., 2012), and both are localized to the peroxisomes of barley cells in a transient expression analysis (Figure 5). Although different mechanisms may account for the specific re-localization of \textit{HvCAT1} upon co-expression with CSEP0027, one scenario can be that CSEP0027 interacts with \textit{HvCAT1} but not \textit{HvCAT2} in the cytoplasm thus interferes with the interaction of PTS1 signal of \textit{HvCAT1} with the peroxisomal receptor. However, how \textit{HvCAT1} is specifically regulated by CSEP0027 is not yet clear. Further investigation of the subcellular localization, trafficking, and post-translational modification of \textit{HvCAT1} will help to better understand the functions of the catalase and the virulence strategies of the biotrophic fungus.

MATERIALS AND METHODS

Plant and Fungal Materials
Barley (\textit{Hordeum vulgare} L.) cultivars (cv) in this study include Golden Promise and “P01” (isogenic line from cv Pallas containing \textit{Mla1}). Barley seedlings were grown in a growth chamber at 20°C with 16 h light and 8 h dark cycles. \textit{N. benthamiana} plants were grown in greenhouse at 24 ± 1°C with a long-day cycle (16 h light/8 h dark).

The barley powdery mildew (\textit{B. graminis} f.sp. \textit{hordei} [\textit{Bgh}]) isolates A6 (\textit{AvrMla6}, \textit{AvrMla10}, and \textit{virMla1}) and K1 (\textit{AvrMla1}, \textit{virMla6}, and \textit{virMla10}) used in this study were maintained on Golden Promise.

Cloning and Plasmid Construction for CSEP Genes
Total RNA was extracted from P01 barley leaves inoculated with \textit{Bgh} isolate A6 using Trizol solution (Invitrogen; 15596-026) and the cDNA was synthesized using reverse transcriptase M-MLV (Invitrogen; C28025). Candidate CSEP sequences excluding the signal peptide (ΔSP) were amplified using the specific primer
pairs (Supplementary Table 2) and subcloned into pGR107 vector through restriction enzyme digestion and ligation for agroinfiltration in N. benthamiana (Wang et al., 2011), all candidates confirmed by sequencing.

Quantitative Real-Time Polymerase Chain Reaction (qRT-PCR)
The analysis of CSEP0027 expression profile was performed as previously described (Ahmed et al., 2015). In brief, total RNA was isolated from P01 leaves at 0, 3, 6, 12, 24, and 48 hpi inoculated with virulent isolate A6. The epiphytic Bgh tissues and the remaining leaf tissues containing Bgh haustoria were separately collected at 24 and 48 hpi. The epiphytic tissues were collected from leaf surfaces by dipping the Bgh-infected leaves into 10% cellulose acetate according to previously described (Ahmed et al., 2015). A quantitative real-time PCR (qRT-PCR) was performed on Applied Biosystems step-one real time PCR system with indicated primers (Supplementary Table 2). Bgh glyceraldehyde 3-phosphate dehydrogenase was used as the reference gene. The statistical significance was evaluated by Student's t test. The assays were repeated two times with three replicates each time.

Yeast Invertase Secretion Assay
The yeast invertase secretion was previously described (Gu et al., 2011). Briefly, the predicted SP sequence of CSEP0027 and Avr1b, or the first 25 amino acids of Magnaporthe oryzae Mg87 were fused in frame with the yeast invertase lacking its own SP in the vector pSUC2. The pSUC2-derived constructs were fused in frame with the yeast invertase lacking its own signal peptide. The pSUC2 constructs were transformed into the invertase secretion-deficient yeast strain YTK12, and yeasts were then placed on CMD-W medium (0.67% yeast N base without amino acids, 0.075% tryptophan dropout YTK12, and yeasts were then placed on CMD-W medium (0.67% yeast N base without amino acids, 0.075% tryptophan dropout

Y2H Analysis
Yeast two-hybrid screening was performed according to the protocols of the manufacturer (Clontech; PT4048-1). In total, 5 × 10^7 transformants were screened. In brief, yeast strain Y2HG0ld expressing pGBK7T-CSE0027 (ASP) was used for mating with yeast strain Y187 harboring a cDNA prey library derived from Bgh-infected barley leaves and placed onto SD-Leu-Trp-His-Ade plates at 30°C. After 35 days, the resistant clones were selected for further verification.

For Y2H assay, the corresponding bait and prey vectors were co-transformed into yeast strain Y2HG0ld and plated onto SD-Leu-Trp plates. The positive interactions were detected by placing the strains onto SD-Leu-Trp-His-Ade plates at 30°C.

Luciferase Complementation Imaging Assays
Luciferase complementation imaging assays were performed according to previously described by Chen et al. (2008). Briefly, the coding region of CSEP0027 (ASP) and HvCAT1 were subcloned into vectors pCAMBIA-Cluc or pCAMBIA-Nluc, respectively, to generate constructs for expressing Cluc-CSEP0027 and Cluc-HvCAT1, or CSEP0027-Nluc and HvCAT1-Nluc. The Nluc-/Cluc-derivative constructs were transformed into the A. tumefaciens strain GV3101. The overnight agrobacteria cultures were resuspended with infiltration buffer (2% sucrose, 0.5% MS, 100 µM acetosyringone, and 10 mM MES) into OD600 = 1.0. Equal volume of agrobacteria resuspension carrying the nLUC and cLUC derivative constructs were mixed and co-infiltrated into the N. benthamiana leaves. The infiltrated area was examined for the luciferase activity 40–50 h post agroinfiltration with a cooled charge-coupled device (CCD) imaging apparatus. For each pair of constructs, at least 10 leaves were co-infiltrated in one experiment, and three independent replicates were conducted.

Glutathione S-Transferase(GST) Pull-Down and Co-immunoprecipitation (Co-IP) Assays
Pull-down assays were performed according to previously described with some modifications (Chang et al., 2013). Briefly, 500 ng of GST-CSEP0027 and GST proteins purified from Escherichia coli were incubated with 150 µl of Glutathione Sepharose 4B beads for 1 h at 4°C, then, beads were washed with 100 µg BSA for 1 h and incubated with 1.0 g crude protein extracted from N. benthamiana leaves expressing HvCAT1-HA. After incubation for 2 h, the beads were washed five times with RB buffer, then resuspended with 30 µl of 2 × Laemmli buffer, and loaded for sodium dodecyl-sulfate polyacrylamide gel electrophoresis (SDS-PAGE) and immunoblotting with anti-HA antibody. GST-CSEP0027 and GST proteins were detected by Ponceau staining.
For Co-IP assay, the total proteins extracted from N. benthamiana coexpressing GFP-Flag/CSEP0027-HA or HvCAT1-Flag/CSEP0027-HA were incubated with anti-FLAG antibody-coupled beads for 2 h, then washed five times with extraction buffer, proteins were further eluted from the beads using 0.5 mg ml⁻¹ 3 × Flag peptide and used for immunoblotting with anti-HA antibody, or anti-Flag antibody.

Confocal Laser Scanning Microscopy and Localization Analysis

For subcellular localization analysis, the coding sequences of CSEP0027ΔSP, HvCAT1 and HvCAT2 were subcloned into vectors pUBI-mYFP-GW and pUBI-GW-CFP to generate pUBI-CSEP0027 ΔSP-CFP, pUBI-mYFP-HvCAT1, and pUBI-mYFP-HvCAT2 constructs. A pair of constructs was delivered into barley leaf epidermal cells by the particle bombardment for coexpression of the indicated fusion proteins, and confocal imaging was conducted at 48 h post-particle delivery. Laser illumination was set at 405 nm for CFP, 488 nm for YFP, and 561 nm for RFP using a Nikon confocal microscope. This assay was repeated three independent times and at least 20 cells were examined for each coexpression.

Barley Stripe Mosaic Virus (BSMV)-Mediated Gene Silencing in Barley

Barley stripe mosaic virus-mediated gene silencing in barley was performed as previously described (Yuan et al., 2011). Briefly, an antisense fragment of HvCAT1 was cloned into the pCaBS-γbLIC vector to create pCaBS-γb-HvCAT1 construct with indicated primers (Supplementary Table 2). pCaBS-α, pCaBS-β, and pCaBS-γb-HvCAT1 constructs were transformed into A. tumefaciens strain EHA105, respectively. The agrobacteria were resuspended in infiltration buffer to OD₆₀₀ = 1.0 and mixed at 1:1:1 ratio to infiltrate N. benthamiana. After 12 days, N. benthamiana leaf sap was extracted to inoculate 10-day-old barley leaves. After 2–3 weeks of inoculation, the newly emerged leaves with virus caused symptoms were used for Bgh infection, and microcolony scoring was done at ~60–72 hpi. For each treatment, at least four barley leaves were chosen for analysis, and three independent replicates were conducted. The statistical significance was evaluated by Student's t test.

Agroinfiltration Mediated Transient Gene Expression in N. benthamiana

Agrobacterium tumefaciens-mediated transient gene expression in N. benthamiana assays were performed as described previously (Wang et al., 2011). A. tumefaciens strain GV3101 was transformed with indicated constructs. Agrobacteria were cultured overnight at 28°C, at 200 rpm, then resuspended in 10 mM MgCl₂ to a final OD₆₀₀ = 0.5 and infiltrated into 4-week-old N. benthamiana leaves. The cell death symptoms were photographed at 5 days post-infiltration. For trypan blue staining, the leaves were boiled in a 1:1 mixture of ethanol and staining solution for 5 min as described before (Bai et al., 2012). The leaves were de-stained with chloral hydrate solution (2.5 g ml⁻¹) for 2 days.

DATA AVAILABILITY STATEMENT

The original contributions presented in this study are included in the article/Supplementary Material, further inquiries can be directed to the corresponding author.

AUTHOR CONTRIBUTIONS

HY and Q-HS designed the research. HY, CJ, and HP performed the experiments with helps from XL, JL, and WH. HY, CJ, LZ, XL, RF, and Q-HS, analyzed the data. WL and RF provided the reagents. Q-HS, HY, and LZ wrote the 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.2021.733237/full#supplementary-material

Supplementary Figure 1 | The MLA13/AVR₉₁₃ triggers cell death in N. benthamiana. Expression of MLA13/AVR₉₁₃ and GFP in N. benthamiana. The experimental procedure used here was the same as that in Figure 1A.

Supplementary Figure 2 | CSEP0027 induces H₂O₂ production in N. benthamiana. DAB staining was performed at 2 days after infiltration to detect H₂O₂ accumulation in the areas infiltrated with agrobacteria transformed with a corresponding construct.

Supplementary Figure 3 | Bgt homologs of CSEP0027 do not trigger cell death in N. benthamiana. (A) Protein sequence alignment of CSEP0027 and its Bgt homologs. Alignment was performed using the DNAMAN software. The red box indicates signal peptides, blue box indicates the Y/FxC motif, and green box
and HvCAT1-CT (402–492) used in the Y2H assay. Catalase core domain and catalase-related immune responsive domain (catalase-reI) are indicated. (B) Y2H analysis of the interaction between NT- or CT-fragments of HvCAT1 and CSEP0027. Yeast was transformed with the indicated bait and prey constructs. Serial dilutions from cell suspension of yeast expressing bait and prey constructs are shown. Growth on SD-Leu-Trp plates indicates yeast clone carrying bait and prey constructs. Interactions were detected on SD-Leu-Trp-His-Ade plates.

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Supplementary Figure 4 | Sequence alignment of amino acids of HvCAT1 and HvCAT2.

Supplementary Figure 5 | CSEP0027 N-terminus interacts with HvCAT1 in yeast. (A) Schematic diagram of HvCAT1 constructs, and HvCAT1-NT (1–401) and HvCAT1-CT (402–492) used in the Y2H assay. Catalase core domain and catalase-related immune responsive domain (catalase-reI) are indicated. (B) Y2H analysis of the interaction between NT- or CT-fragments of HvCAT1 and CSEP0027. Yeast was transformed with the indicated bait and prey constructs. Serial dilutions from cell suspension of yeast expressing bait and prey constructs are shown. Growth on SD-Leu-Trp plates indicates yeast clone carrying bait and prey constructs. Interactions were detected on SD-Leu-Trp-His-Ade plates.

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