Colletotrichum higginsianum effectors exhibit cell to cell hypermobility in plant tissues and modulate intercellular connectivity amongst a variety of cellular processes

Mina Ohtsu†, Joanna Jennings†, Matthew Johnston¹, Xiaokun Liu¹, Nathan Hughes², Kara Stark¹, Richard J. Morris², Jeroen de Keijzer¹‡, Christine Faulkner¹*

¹Crop Genetics, John Innes Centre, Norwich Research Park, UK
²Computational and Systems Biology, John Innes Centre, Norwich Research Park, UK
‡Present address: Developmental and Stem Cell Biology, Duke University, Durham, North Carolina, USA
ªPresent address: Cluster Plant Developmental Biology, Wageningen University, Droevendaalse steeg 1, 6708 PB, Wageningen, The Netherlands
†These authors contributed equally to this work
*Corresponding author: christine.faulkner@jic.ac.uk, +44 (0)1603 450000

Summary

Multicellular organisms exchange information and resources between cells to co-ordinate growth and responses. In plants, plasmodesmata establish cytoplasmic continuity between cells to allow for communication and resource exchange across the cell wall. Some plant pathogens use plasmodesmata as a pathway for both molecular and physical invasion. However, the benefits of molecular invasion (cell-to-cell movement of pathogen effectors) are poorly understood. To begin to investigate this and identify which effectors are cell-to-cell mobile, we performed a live imaging-based screen and identified 15 cell-to-cell mobile effectors of the fungal pathogen Colletotrichum higginsianum. Of these, 6 are “hypermobile”, showing cell-to-cell mobility greater than expected for a protein of its size. We further identified 3 effectors that can indirectly modify plasmodesmal aperture. Transcriptional profiling of plants expressing hypermobile effectors implicate them in a variety of processes including senescence, glucosinolate production, cell wall integrity, growth and iron metabolism. However, not all effectors had an independent effect on virulence. This suggests a wide range of benefits to
infection gained by the mobility of *C. higginsianum* effectors that likely interact in a complex way during infection.
Cell-to-cell communication is essential for multicellularity and there are a variety of mechanisms by which cells exchange information and resources. Plant cells are surrounded by cell walls but have tunnel-like structures called plasmodesmata which cross the cell wall and directly connect the cytoplasm of adjacent cells to establish the symplast. Small molecules such as sugars, metabolites and hormones can all move between cells through plasmodesmata (Stahl and Simon, 2013; Cheval and Faulkner, 2018; Liu and Chen, 2018), driven by advection and diffusion. Further, larger proteins can pass between cells via mechanisms that involve active components such as protein unfolding and refolding or possible intercellular trafficking motifs/domain (Kragler et al., 1998; Taoka et al., 2007; Xu et al., 2011; Chen et al., 2013; Chen et al., 2014).

The regulation of plasmodesmata is a critical component of plant-microbe interactions. Many plant immune responses are triggered by cell autonomous recognition of pathogen molecules, but we and others have shown that plasmodesmata dynamically respond to immune signals. We previously found that both fungal and bacterial molecules induce plasmodesmal closure in Arabidopsis thaliana. Chitin (from fungal cell walls) and flg22 (from bacterial flagellin) both trigger plasmodesmal closure, regulated by LYSM-CONTAINING GPI-ANCHORED PROTEIN 2 (LYM2) and CALMODULIN-LIKE 41 respectively (Faulkner et al., 2013; Xu et al., 2017). Observations that plasmodesmal function influences infection outcomes identify that plasmodesmal responses are key to ultimate defence success (Lee et al., 2011; Faulkner et al., 2013; Caillaud et al., 2014; Xu et al., 2017). This suggest two possibilities: plasmodesmal regulation is important for the execution of plant immune responses and/or it impairs infection mechanisms deployed by the invading pathogens. The latter implicates pathogen access to the symplast as a critical component of infection.

Plasmodesmata present a route by which microbes can access non-infected cells and tissues. Indeed, there are several examples of pathogens directly using plasmodesmata to facilitate their passage between host cells in a growing infection. The best understood examples of this are viral infections such as Cucumber mosaic virus (CMV) and Tobacco mosaic virus (TMV) (Heinlein and Epel, 2004). As obligate parasitic pathogens that infect intracellularly, viruses actively target and modify plasmodesmata to translocate their genomes between cells to establish an infection. Interestingly, it has also been revealed that some hemi-biotrophic fungal pathogens, including the rice blast pathogen Magnaporthe oryza, pass from cell to cell.
at plasmodesmal pitfields (Kankanala et al., 2007). These demonstrate that plant pathogens across different kingdoms have acquired the capacity to recognize and exploit plasmodesmata as sites of connection between cells to enable the spread of infection.

In order to manipulate host plant systems, plant pathogens secrete an arsenal of proteins, called effectors (Le Fevre et al., 2015; Toruño et al., 2016). Plasmodesmata allow the spread of soluble molecules, and while effectors can act in the host cell to which they are delivered, soluble effectors also have the potential to move between cells via plasmodesmata. Indeed, *M. oryzae* produces the PWL2 and BAS1 effectors that can move into non-infected cells (Khang et al., 2010). This suggests that pathogens exploit plasmodesmata to access and manipulate non-infected cells ahead of the infection front. Further implicating the symplast in infection, the *Fusarium oxysporum* effector Six5 was found to enable cell-to-cell translocation of its co-transcribed effector Avr2 via plasmodesmata (Cao et al., 2018). Moreover, both the *Pseudomonas syringae* effector HopO1-1 and the *Phytophthora Brassicaceae* effector RxLR3 target and modify plasmodesmata (Aung et al., 2020; Tomcynska et al., 2020).

It is not yet fully understood what a microbe gains by accessing the host symplast, or how common cell-to-cell mobility is within effector repertoires. To address these questions, we characterized the cell-to-cell mobility of candidate effectors from the hemi-biotrophic fungal pathogen *Colletotrichum higginsianum*. We used a live imaging-based screen to identify candidate effectors that move cell to cell in plant tissues and identified effectors that are cell-restricted (immobile), move cell to cell to a degree expected for a protein of that size (mobile) and move further than expected (hypermobile). Within the hypermobile effectors, we identified effectors that modify plasmodesmata, consistent with enhanced mobility, and one with a signature of active translocation. Expression of hypermobile effectors in host tissue identified that these three effectors have a differential effect on pathogen virulence and the host transcriptome. The latter identified processes associated with nutrient uptake and defence that illustrate what *C. higginsianum* gains by molecular invasion of host tissues. We conclude that pathogen access to the host symplast is a complex component of the infection strategy of *C. higginsianum* allowing it to manipulate a variety of host processes ahead of the infection front.
Material and Methods

Plant material

*Nicotiana benthamiana* plants were grown at 23 °C under long day conditions (16 h: 8 h, light: dark). *Arabidopsis thaliana* were grown on soil at 22 °C under short day conditions (10 h: 14 h, light: dark) or on MS media under short day conditions (10 h: 14 h, light: dark).

DNA Constructs

Constructs for plant expression were assembled using the Golden Gate cloning method (Engler *et al.*, 2008) and all module information is in Table S1. The coding sequences of effector candidates (without predicted signal peptides) were domesticated to remove BpiI and BsaI sites and synthesised as Golden Gate-compatible Level 0 vectors. For subcellular localisation analysis, effector sequences were fused to the N-terminus of GFP and expressed from the CaMV 35S promoter. For the cell-to-cell mobility assay and generating Arabidopsis stable lines, multi-component binary vectors were assembled as outlined in Fig. S1 and Table S1.

Plant transformation

Transient gene expression in *N. benthamiana* was performed as described (Cheval *et al.*, 2020). *Agrobacterium tumefaciens* (GV3101) carrying binary plasmids were infiltrated into *N. benthamiana* leaves at OD$_{600nm}$ = $1.0 \times 10^{-2}$ to check subcellular localisation and at OD$_{600nm}$ = $1.0 \times 10^{-5}$ to generate single cell transformation events for the mobility assay. Samples were imaged 72 h post infiltration. Stable transgenics were made by floral dipping Arabidopsis Col-0.

Microscopy

Leaf tissue was imaged by confocal microscopy (Zeiss LSM800) with a 20x water-dipping objective (W N-ACHROPLAN 20x/0.5; Zeiss). GFP was excited with a 488 nm solid-state laser and collected at 509-530 nm and dTomato was excited by a 561 nm solid-state laser and collected at 600-640 nm using sequential scanning.

Image and data analysis

To quantify effector-GFP mobility we recorded the number of fluorescent cells around the transformed cell, identified by NLS-dTomato fluorescence. Data analysis was performed in R statistical computing language v4.0.3 (R Core Team, 2020). The standard curve was generated
with data obtained from mobility of GFP, YFP\textsubscript{N}-GFP, YFP\textsubscript{C}-GFP and 2xGFP using a quasi-Poisson generalised linear model with a log link function and a Bonferroni corrected p-value < 1x \times 10^{-5}. Effectors were determined to be significantly mobile by the equivalent of a t-test for Poisson distributions (an exact Poisson test) in which the mobility (cell count at 3dpi) is significantly different to the standard curve (p <1x10^{-5}). For analysis of NLS-dTomato movement in the presence of different effectors we tested the null hypothesis that mobility of NLS-dTomato is 2.5 cells (as observed for the set of standard constructs) and independent of effector mobility using an exact Poisson test.

**Protein extraction and Western blotting**

Protein extraction was performed as described (Adachi et al., 2019). *N. benthamiana* leaves transiently expressing effectors were collected tissue for protein extraction 3 days post infiltration. Leaf material was homogenised in ice-cold extraction buffer (10% glycerol, 25 mM Tris pH 7.5, 1 mM EDTA, 150 mM NaCl, 2% w/v PVPP, 10 mM DTT, protease inhibitor cocktail (Sigma), 0.2% Igepal (Sigma)) and the soluble fraction was collected by centrifugation at 12,000 x g for 10 min. Proteins were separated by SDS-PAGE and transferred PVDF membrane (BioRad). Proteins were detected with 1/5,000-diluted anti-GFP conjugated to HRP (sc-9996; Santa Cruz Biotechnology).

**Microprojectile bombardment**

Microprojectile bombardment assays were performed as described (Faulkner et al., 2013). Four- to six-week-old expanded leaves of relevant Arabidopsis lines were bombarded with 1 nm gold particles (BioRad) coated with pB7WG2.0-mRFP, using a Biolistic PDS-1000/He particle delivery system (BioRad). Bombardment sites were imaged 24 h post bombardment by confocal microscopy (Zeiss LSM800) with a 10x (EC Plan-NEOFLUAR 10× 0.3; Zeiss) or 20x dry objective (Plan-APochromat 20x/0.8; Zeiss). Data were collected from at least 2 independent bombardment events, each of which consisted of leaves from at least 3 individual plants. The median normalised mobility (#cells) for different lines was analysed in R statistical computing language v4.0.3 (R Core Team, 2020) by a bootstrap method (Johnston and Faulkner, 2020).

**C. higginsianum infection**

*C. higginsianum* spores (5 × 10^6 spores/mL) were drop-inoculated on detached leaves of 4- to 5-week-old Arabidopsis plants on 2% water agar plates. Plates were sealed with parafilm
and left for 6 days under short day conditions (10 h: 14 h, light: dark; 25 °C). The area of necrotic lesions was measured in Fiji (Schindelin et al., 2012). Lesions were measured for at least 30 plants (2 leaves per plant) and across 3 independent experiments. The mean lesion area for different lines was compared using a bootstrapping method (Johnston and Faulkner, 2020).

**RNAseq analysis and GO term analyses**

Leaves 7 and 8 of 4-week-old MS grown Arabidopsis was harvested for RNA extraction. Leaves from 3 plants were pooled for a single replicate and 3 replicates were collected for each genotype. RNA was extracted using RNAeasy Mini Kit (Qiagen) followed by DNase treatment (TurboDNase, ThermoFisher Scientific). Library preparation and Illumina sequencing (PE150, Q30>80%) with 10M paired reads was performed by Novogene. Sequencing reads quality was evaluated using FastQC v0.11.9 (Andrews, 2010) and multiqc v1.9 (Ewels et al., 2016). After quality control, trimomatic v0.39 was used to remove Illumina sequence adapters and low-quality reads. Processed reads were re-assessed with FastQC v0.11.9 and mapped to the Arabidopsis thalina TAIR10 release 37 genome assembly using hisat2 v2.2.0 and samtools v1.11 (Li et al., 2009).

Differential expression analysis was performed with DESeq2 v1.20.0 (Love et al., 2014) and the R statistical computing language v4.0.3 (R Core Team, 2020). Analysis using the methods described by Love et al., 2014. were used to calculate normalised expression values for each gene across all samples. Normalised expression values were compared for all expressed genes in Col-0 to those in the effector expressing lines using a hypergeometric test with the Benjamini and Hochberg False Discovery Rate (FDR) correction. Differentially expressed genes were defined by |log2[fold change]|≥1 and FDR corrected p-value < 0.01. Differentially expressed genes were analysed by GoTermFinder (Boyle et al., 2004) with a hypergeometric test with a Bonferroni correction to identify biological processes enriched in the samples (adjusted p-value>0.01). CirGO (Kuznetsova et al., 2019) was used to visualise the results.

**Results**

**Identification of candidate cell-to-cell mobile *Colletotrichum higginsianum* effectors**

To establish a candidate list of putative *C. higginsianum* effectors (hereafter referred to as effectors) exhibiting cell-to-cell mobility, we mined published transcriptome data covering
different infection stages (O’Connell et al., 2012; Dallery et al., 2017). Many effectors are secreted from pathogens into host plant tissues (Lo Presti et al., 2015), and therefore we limited our candidate cell-to-cell mobile effectors as those that encode conventionally secreted proteins with a predicted signal peptide. Gene expression data in O’Connell et al. (2012) defines expression profiles during the following stages of growth and infection: in vitro appressoria (VA), in planta appressoria (PA), biotrophic phase (BP) of infection and necrotrophic phase (NP) of infection. We reasoned that cell-to-cell mobile effectors would be primarily relevant during the penetration (PA) and biotrophic phases (BP) of infection (i.e. when the host tissue is alive) and thus defined candidate cell-to-cell mobile effectors as those that have enhanced expression in PA and BP phases relative to the other 2 stages (i.e. PA/VA, BP/VA, PA/NP, and BP/NP > 2). We further limited candidates to those for which PA reads > 50 and BP reads > 20. These criteria produced a list of 46 candidate effectors.

Plant proteins that are known to be cell-to-cell mobile are typically soluble within the cytoplasm or the nucleus (Kim et al., 2002; Gallagher et al., 2004; Gallagher and Benfey, 2009; Chen et al., 2013) and we made the assumption that C. higginsianum cell-to-cell mobile proteins would have similar properties. Thus, to further refine our list of candidate cell-to-cell mobile effectors, we cloned these 46 candidate effectors as fusions to GFP and screened for nucleocytoplasmic and nuclear subcellular localisations by transient transformation of N. benthamiana. Of these 46 effector-GFP fusions, 25 showed nucleocytoplasmic localisation but none showed a specific nuclear localisation.

Live cell screening for cell-to-cell mobility

To assay the cell-to-cell mobility of candidate effectors, we performed a live cell imaging-based screen using transient transformation of single epidermal cells in N. benthamiana. For this we used Golden Gate modular cloning (Engler et al., 2008) to assemble effector-GFP fusions and a cell transformation marker in a single vector as a dual expression cassette vector (Fig. 1a). For a cell transformation marker, we used dTomato fused to a nuclear localisation sequence (NLS-dTomato), reasoning that the dimerization properties of dTomato would make a protein complex too large to move from cell to cell.

To confirm the utility of NLS-dTomato as a cell transformation marker, we generated constructs that express GFP or 2xGFP with NLS-dTomato (pICH4723.GFP.NLS-dTomato and pICH4723.2xGFP.NLS-dTomato respectively). Agrobacterium infiltration of N. benthamiana leaves demonstrated that both fluorophores were expressed in the transformed cell. While GFP
was frequently seen to move freely from the transformed cell, both NLS-dTomato and 2xGFP were mostly retained within the transformed cell (Fig. 1 and S1).

Cell-to-cell mobility via plasmodesmata is dependent upon the size of the molecule (Oparka et al., 1999; Crawford and Zambryski., 2001). To determine the relationship between size and mobility in *N. benthamiana* leaves in more detail, we assayed the mobility of four proteins of different sizes: GFP (26 kDa), YFPC-GFP (37.2kDa), YFPn-GFP (45 kDa) and 2x GFP (52 kDa) (Fig. S1). We generated single cell transformation sites by low OD600 Agrobacterium infiltration of *N. benthamiana* leaves and counted the number of cells which exhibited GFP surrounding a transformed cell for each construct (marked by NLS-dTomato) (Fig. 1). We fit a quasi-Poisson function through the data which makes minimal assumptions, obtaining a smooth fit through the data that allows for the infrequent mobility of the larger standard proteins (Fig. S1). However, we point out that the data itself does not exclude the existence of a SEL which could be anywhere upwards from YFPC-GFP. This model, with a Bonferroni corrected confidence interval of the mean ($p < 1x10^{-5}$), defined a standard curve against which to compare mobility of effectors of varying sizes.

**Cell-to-cell mobility screen identifies mobile and hypermobile effectors**

To characterize the cell-to-cell mobility of nucleocytoplasmic effectors, we cloned each of the 25 effectors in a dual expression cassette vector as described (i.e. pICH4723.Effector-GFP.NLS-dTomato) (Fig. 1a). Assaying for cell-to-cell mobility, we observed varying degrees of cell-to-cell mobility for the effectors (i.e. Fig. 1, lower panels); 10 effectors were restricted to the transformed cell (i.e. Fig. 1b, middle panels) in *N. benthamiana* and 15 exhibited cell-to-cell mobility (Table S2).

To determine whether an effector’s mobility was as expected for a soluble protein of that size, we compared effector mobility to the standard curve (Fig. S1b; Fig. S2). An exact Poisson test identified a subset of 7 effectors that had greater than expected mobility, and we defined these as ‘hypermobile’ (Fig. 2). To confirm that this mobility did not arise from cleavage of the effector-GFP fusion (to produce a smaller and thus more mobile protein), we assayed the protein size of the GFP-fused 7 hypermobile effectors expressed in *N. benthamiana* by protein extraction and Western blot analysis. Two strong bands for ChEC130-GFP were observed, suggesting this effector is cleaved in host cells (Fig. S3). However, the other 6 effectors showed little evidence of significant degradation or cleavage (Fig. S3). Thus, we concluded that ChEC123, ChEC124, ChEC125, ChEC128, ChEC129 and ChEC132 are hypermobile.
Mobile effectors can modify plasmodesmal function

The identification of both mobile and hypermobile effectors suggests that *C. higginsianum* might access the host symplast via different mechanisms. Hypermobility suggests three possibilities: that the shapes or Stokes radius of such effectors allows for more efficient translocation through plasmodesmata; that effectors exploit an active translocation mechanism; or effectors modify PD (directly or indirectly) which allows their passage. To address the question of whether mobile and hypermobile effectors modified plasmodesmal function, we exploited the unexpected low-level mobility of NLS-dTomato observed post-collection when the image contrast was adjusted (Fig. S4). NLS-dTomato clearly marked the transformed cell in all cases, but upon increasing the contrast of the images it was observed in an average of 2.5 cells surrounding the brightest transformed cell in size-standard controls (Fig. S4a; Fig. S5). We quantified NLS-dTomato movement when co-expressed with each mobile and hypermobile effector and observed variation in the spread of NLS-dTomato (Fig. 3a; Fig. S4b). To determine if hypermobility is associated with a general increase in mobility that would indicate plasmodesmal regulation, we compared relative mobility (Mob_r = Mob_{observed}/Mob_{expected}) to the mobility of NLS-dTomato. An exact Poisson test of this data, using the null hypothesis that mobility of NLS-dTomato is 2.5 cells for all values of Mob_r, revealed that ChEC127 and ChEC8 both significantly increase NLS-dTomato mobility (Fig. 3a). This result suggests that ChEC127 and ChEC8 modify plasmodesmal function. Curiously, while ChEC127 is a hypermobile effector, ChEC8 is not, suggesting that despite modifying plasmodesmal function, the effector itself does not have increased translocation. This phenomenon might be explained if ChEC8 binds other proteins in plant cells to increase its effective size.

NLS-dTomato is targeted to the nucleus and therefore has a limited pool available in the cytoplasm for cell-to-cell movement. Further, NLS-dTomato is expected to dimerise to form a complex that we expect has reduced mobility as a consequence of increased size. Thus, mobility of this protein is a low sensitivity marker for changes to plasmodesmal function detecting only large changes to plasmodesmal aperture. Therefore, we extended our analysis of plasmodesmal function in the presence of hypermobile mobile effectors to examine the mobility of cytoplasmic mRFP in Arabidopsis, a native host plant of *C. higginsianum*. For this, we generated Arabidopsis lines that stably express fluorescent protein fusions of the hypermobile effectors ChEC123, ChEC127 and ChEC132 (Fig. S6). We performed microprojectile bombardment assays on expanded leaves of two independent lines of each
effector (Faulkner et al., 2013) and quantified spread of mRFP from transformed cells one day post-bombardment. This data showed that mRFP diffusion in ChEC123-expressing lines was similar to Col-0, while ChEC127 and ChEC132 expressing lines showed increased movement of mRFP relative to the Col-0 control (Fig. 3b). Thus, this data indicates that Ch132 also modifies plasmodesmal function like ChEC127 and ChEC8, and that ChEC123 mediates its translocation by a possible active mechanism that does not involve plasmodesmal modification.

**Heterologous expression of ChEC127 promotes virulence**

Pathogen effectors are assumed to positively regulate virulence, facilitating infection success. To assess whether the hypermobile effectors ChEC123, ChEC127 and ChEC132 independently promote virulence, we infected plants constitutively expressing these effectors with *C. higginsianum* and measured the size of disease lesions 6 days post inoculation. These assays showed that both independent transgenic lines that express ChEC127 develop larger disease lesions (Fig. 4), identifying that expression of ChEC127 promotes virulence. Plants expressing ChEC123 or ChEC132 showed no increase in susceptibility (Fig. 4), suggesting that in these conditions neither effector independently promotes virulence.

**ChEC132 and ChEC123 transcriptionally perturb a variety of host processes**

To explore the function of hypermobile effectors, and thus ask what a pathogen might gain from these during infection, we assayed changes to the plant transcriptome induced by ChEC123, ChEC127 and ChEC132. For this, we performed RNAseq analysis of leaf tissue of plants constitutively expressing ChEC123-GFP, ChEC127-GFP and ChEC132-GFP relative to Col-0, defining differentially expressed genes as those for which we detected at least a 2-fold up or down regulation ($|\log_2[\text{fold change}]|\geq 1$) and an FDR corrected $p$-value $<0.01$ To identify processes that are perturbed by these effectors, we used GO Term Finder (Boyle et al., 2004) to identify biological processes that are significantly enriched within the differentially expressed genes for each effector.

Despite positively regulating virulence, ChEC127 differentially regulated the expression of only 13 genes (11 up-regulated and 2 down-regulated; Table S4). By contrast, expression of ChEC123 and ChEC132 caused differential expression of 176 and 217 genes respectively (Table S3; S5). GO term analysis indicates ChEC123 up-regulated genes associated with leaf senescence (Fig. 5c) and down regulates genes associated with glycosyl compound catabolism and iron starvation responses and transport (Fig. 5b). Constitutive expression of ChEC132 also induced down regulation of genes associated with iron transport and responses, in addition to
cell wall modification, growth, syncytium formation (5 EXPANSIN genes also represented in the cell wall loosening GO term), and lipid metabolism (Fig. 5a).

Discussion

Host cell-to-cell connectivity is increasingly identified as a component of plant immunity and pathogen infection. This suggests that pathogen access to non-infected cells is important for the infection strategies of a range of pathogens. Previous studies have identified that specific effectors move cell-to-cell in host tissues (Khang et al., 2010; Cao et al., 2018), and it was recently suggested that cell-to-cell mobility is a property common to many proteins within an effector repertoire (Li et al., 2020). To investigate this further, we performed a screen for cell-to-cell mobility of effectors from the hemi-biotrophic pathogen C. higginsianum. We generated a candidate list of secreted effectors from publicly available data and used live-cell imaging to identify a subset of 25 that have a nucleocytoplasmic localisation similar to many known cell-to-cell mobile molecules. To identify mobile effectors, we established a live cell imaging-based screen for cell-to-cell mobility and identified that 60% of nucleocytosolic effectors are cell-to-cell mobile (15/25), with 24% (6/25) showing greater than expected mobility (hyper-mobility) (Fig. 2; Fig. S2; Table S2).

In addition to identifying that cell-to-cell mobility is possible for a range of effectors, our data suggests that effectors from C. higginsianum can move through plasmodesmata by different mechanisms. Firstly, we identified a subset of effectors that move ‘passively’ from cell to cell, i.e. they move through plasmodesmata at a rate expected for soluble molecules of the same size (Fig. 2). Like the endogenous plant transcription factor LEAFY (Wu et al., 2003), these molecules can be considered to have no mechanism for active translocation and simply move from cell to cell as soluble, freely diffusive molecules. Secondly, we identified hypermobile effectors that modify plasmodesmal function such as ChEC127 and ChEC132. These effectors trigger plasmodesmata opening to a degree such that the effector itself (Fig. 2), as well as other soluble molecules (as observed for mRFP; Fig. 3b), can move faster and further to neighbouring cells. Thirdly, we identified one hypermobile effector, ChEC123, that has no general effect on plasmodesmal function (Fig. 2; Fig. 3). This effector could therefore be considered to have an active and specific mode of translocation to surrounding cells similar to endogenous plant transcription factors such as SHORT-ROOT (SHR) (Nakajima et al. 2001), KNOTTED1(KN1) (Lucas et al., 1995) or Dof family proteins (Chen et al., 2013).
This data identifies different mechanisms by which individual effectors can access the symplast, but this data must be considered in the context of infection. Thus, the presence of multiple effectors that modify plasmodesmata during infection raises the possibility that there is a general increase in plasmodesmal aperture that might allow cell-to-cell mobility of effectors that we identified here as immobile.

Our screen revealed that three nucleocytoplasmic effectors can modify plasmodesmal function (ChEC8, ChEC127 and ChEC132), identifying that plasmodesmal regulation might occur during infection via indirect mechanisms. While effectors are expected to positively contribute to virulence, this is not always observable as an independent contribution; effectors may act in concert with other effectors or environmental conditions. In our study, we saw that only ChEC127 significantly and positively contribute to virulence independently as evidenced by increased susceptibility of Arabidopsis plants that express the effector (Fig. 4). Infection is a complex process and interacts with an array of host and environmental variables. Thus, it may not be surprising that excess of a single effector does not promote infection.

To identify host processes that are modulated in a non-cell autonomous way during infection, we performed an RNAseq analysis of plants constitutively expressing the hypermobile effectors ChEC123, ChEC127 and ChEC132 (Table S3-S5). Despite regulating pathogen virulence, ChEC127 induced differential expression of only 13 genes (Table S4), suggesting the mechanism by which it promotes virulence does not involve significant perturbation of gene expression. By contrast, ChEC123 and ChEC132, which did not independently promote virulence, did induce significant changes in gene expression (Table S3; Table S5). ChEC123 downregulated genes associated with glucosinolate production (glycosyl compound catabolism) and iron starvation and transport, indicating it may regulate defence and nutritional processes (Fig. 5b). The same effector up-regulated genes associated with leaf senescence which might contribute to the transition of \textit{C. higginsianum} to the necrotrophic lifestyle (Fig. 5c). ChEC132 down regulated genes associated with iron metabolism (Fig. 5a) but most significantly perturbed processes associated with plant cell wall modification and growth. This raises the possibility that ChEC132 perturbs growth processes, possibly to limit resource consumption by the host. Overall, transcriptional analysis identifies that different host processes can be targeted by cell-to-cell mobile effectors. How manipulation of these processes in cells ahead of the infection front contributes to infection success requires further investigation.
Our screen for cell-to-cell mobility of *C. higginsianum* effectors has identified both mobile and hypermobile effectors. Further, we found evidence that some of these cytoplasmic effectors, both mobile and hypermobile, indirectly regulate plasmodesmata to increase their functional aperture. These observations identify that *C. higginsianum* has complex strategies for accessing the symplast which allows it to perturb processes associated with defence, nutrition and cell structure ahead of hyphal invasion. Evidence of hypermobility in the *C. higginsianum* effector repertoire identifies that exploiting plasmodesmata and cell to cell connectivity to extend pathogen reach is critical for infection.

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**Authors Contributions**

MO, JJ, XL, KS, and JdK generated materials and performed experiments; MO, JJ, MJ, RJM and NH analysed data; CF managed the project; and MO, JJ and CF wrote the paper with support from all co-authors.

**Data Availability**

Raw or normalised cell counts from image analysis is available in Table S1. DESeq2 analysis of RNA sequencing experiments is available in Tables S3-S5.
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Figure Legends

**Fig. 1** GFP fusions allow detection of cell-to-cell mobility screening of *C. higginsianum* effectors. (a) Schematic of the dual expression module binary vectors used to screen for cell-to-cell mobility. The cartoon represents the patterns of localisation for each module’s gene product expected for a mobile effector-GFP fusion. (b) Projections of confocal z-stacks (8-20 individual focal planes, taken at 5.66 μm intervals) of tissues expressing GFP alone and effector-GFP fusions. Mobility is visible when cells surrounding the transformed cell, marked by NLS-dTomato, show GFP fluorescence. ChEC133-GFP (middle row) shows an example of a cell-restricted, immobile effector while ChEC125-GFP (bottom row) shows an example of mobility to surrounding cells as observed for GFP alone (top panel). Arrows indicate NLS-dTomato fluorescence in the transformed cell, stars represent the transformed cells and arrowheads represent cells the GFP has moved into. Scale bar is 100 μm.

**Fig. 2** Mobility quantification for nucleocytoplasmic effectors reveals mobile and hypermobile effectors. Mobility (number of cells to which GFP has moved around a transformed cell) of effector-GFP fusions. The standard curve is a quasi-Poisson generalised linear model with a log link function and the Bonferroni corrected confidence interval of the mean (*p* < 1x10^{-5}) (red ribbon). Effectors were determined to be significantly mobile (purple squares) by an exact Poisson test indicating the rate of movement is significantly different to the standard curve (*p* < 1x10^{-5}). Data are means ± standard error (*n* >30, *p* < 1 x 10^{-5})

**Fig. 3** Plasmodesmal regulation by mobile and hypermobile effectors. (a) Mobility of NLS-dTomato plotted against the relative mobility (Mobr) of a co-expressed effector in *N. benthamiana* leaf epidermal cells. The standard curve represents NLS-dTomato mobility in the presence of GFP variants of different sizes with Bonferroni corrected confidence interval of the mean (*p* < 1x10^{-5}red ribbon). NLS-dTomato movement was analysed by an exact Poisson test, identifying that ChEC8 and ChEC127 increase NLS-dTomato mobility (b) Mobility of mRFP (number of cells) in Arabidopsis leaf tissue assayed by microprojectile bombardment assays. Independent transgenic lines expressing ChEC127-GFP lines and ChEC132-GFP showed increased mobility of mRFP relative to Col-0. Boxes signifies the upper and lower quartiles, and the whiskers represent the minimum and maximum within 1.5 × interquartile range. The number of bombardment sites (*n*) counted is ≥ 92. Data was analysed by bootstrap analyses and asterisks indicate statistical significance compared with Col-0 plants (***p* < 0.01 and *p* < 0.05)
Fig. 4 *C. higginsianum* lesion areas are larger in ChEC127 expressing Arabidopsis. Detached leaves from 4-5-week-old Arabidopsis were inoculated with *C. higginsianum* spores and lesion areas were measured 6 dpi (n > 60) and bootstrap analysis of the lesion area means identified that lesions are larger in ChEC127 expressing plants.

Fig. 5 Hypermobile effectors can alter host plant gene expression patterns. CirGO visualisation of GO Terms enriched amongst (a) genes down-regulated by ChEC132, (b) genes down-regulated by ChEC123, and (c) genes upregulated by ChEC123. Slice size represents the proportion of DEGs attributed to this GO Term. The inner ring slices represent ‘parent’ GO terms and the labelled outer ring slices represent ‘child’ GO terms.
Supplementary Figure Legends

Fig. S1 Mobility of GFP variants of known sizes to generate a standard curve. Binary vectors with different sized GFP fusions and NLS-Tomato were transiently expressed in 5-week-old N. benthamiana and imaged by confocal microscopy 3 dpi: GFP (26 kDa), YFPc-GFP (37.2 kDa), YFPn-GFP (45 kDa) and 2xGFP (52 kDa). (a) Arrows indicate NLS-dTomato fluorescence in the nuclei of transformed cells and arrowheads indicate examples of GFP movements and stars indicate the transformed cell. Each image is a maximum projection of a z-stack comprising 8-20 individual focal planes acquired at 4.61/5.66 μm intervals. Scale bars represent 100 μm. (b) Observed mobility for the various GFP-fusions plotted against their molecular weight. This data was used to define a standard curve with a quasi-Poisson generalised linear model with a log link function. The standard error (purple ribbon) and the Bonferroni corrected confidence interval of the mean (p < 1x10^{-5}, red ribbon) was calculated. The point density shows the number of replicates at that value.

Fig. S2 Effector-GFP movement was dependent on effector size. Raw data of mobility assay showing data spread (summarised in Figure 2) for all effector-GFP fusions. The dot gray level indicates the number of replicates at that value.

Fig. S3 Stability of hypermobile effector-GFPs in N. benthamiana leaves Five-week-old N.benthamiana leaves expressing free GFP and effector-GFP fusions were harvested 3dpi. Total proteins were extracted from harvested leaves, separated by SDS-PAGE and were detected using anti GFP antibody. Protein loading was monitored by Coomassie Brilliant Blue (CBB) staining of bands corresponding to the ribulose-1,5-bisphosphate carboxylase large subunit (RBCL).

Fig. S4 Mobility of NLS-dTomato in effector expressing tissues (a) Mobility of NLS-dTomato was detected when image display settings were adjusted post-collection. The image on the left shows the imaging data under unsaturated black/white display levels and on the right when the brightness and contrast were enhanced. The image represents a maximum projection of a z-stack comprising 11 individual focal planes. Scale bars represent 50 μm. (b) NLS-dTomato was detected in surrounding cells (arrows) when co-expressed with a variety of effector-GFP fusions. Stars identify the transformed cell. Images are maximum projections of z-stacks comprising 8-20 individual focal planes acquired at an interval of 5.66 μm. Scale bars represent 100 μm.
Fig. S5 NLS-Tomato moves an average of 2.5 cells irrespective of GFP fusion size. Binary vectors encoding different sized GFP fusions (from 26kDa to 52kDa) and NLS-Tomato were transiently expressed in *N. benthamiana* leaves and imaged by confocal microscopy 3 dpi. The number of cells the NLS-dTomato had moved was counted and a line of best fit generated. The standard error (purple ribbon) and the Bonferroni corrected confidence interval of the mean (*p* < 1x10^-5) (red ribbon) was calculated for the data. The dot gray level indicates the number of replicates at that value.

Fig. S6 Expression and localisation of the hypermobile effector in Arabidopsis stable lines

Confocal micrographs of the epidermis of mature leaves of two independent transgenic Arabidopsis lines that express ChEC123, ChEC127 and ChEC132 fused to a fluorescent protein. Each image is a maximum projection of a z-stack comprising 8-20 individual focal planes acquired at a plane interval of 3 µm. Scale bars are 100 µm.

Supplemental Table legends

**Table S1** Plasmids used and constructed in this study

**Table S2** Nucleocytoplasmic effectors screened for cell-to-cell mobility

**Table S3** DESeq2 analysis of ChEC123-expressing *A. thaliana* compared to Col-0

**Table S4** DESeq2 analysis of ChEC127-expressing *A. thaliana* compared to Col-0

**Table S5** DESeq2 analysis of ChEC132-expressing *A. thaliana* compared to Col-0

**Table S6** Raw data from mobility assays presented in Fig 2, Fig 3 and Fig S2.
Fig. 1 GFP fusions allow detection of cell-to-cell mobility screening of *C. higginsianum* effectors. (a) Schematic of the dual expression module binary vectors used to screen for cell-to-cell mobility. The cartoon represents the patterns of localisation for each module’s gene product expected for a mobile effector-GFP fusion. (b) Projections of confocal z-stacks (8-20 individual focal planes, taken at 5.66 μm intervals) of tissues expressing GFP alone and effector-GFP fusions. Mobility is visible when cells surrounding the transformed cell, marked by NLS-dTomato, show GFP fluorescence. ChEC133-GFP (middle row) shows an example of a cell-restricted, immobile effector while ChEC125-GFP (bottom row) shows an example of mobility to surrounding cells as observed for GFP alone (top panel). Arrows indicate NLS-dTomato fluorescence in the transformed cell, stars represent the transformed cells and arrowheads represent cells the GFP has moved into. Scale bar is 100 μm.
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