Intercellular cooperation in a fungal plant pathogen facilitates host colonization

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Cooperation is the process by which biological units, or modules, incur a cost to provide a benefit to the group they belong to (1). The emergence of cooperation typically is associated with division of labor—the specialization of several modules (cells, organs, or whole organisms), each devoted to specific tasks. Cooperation also generally involves the exchange of molecules between specialized modules, a process called “resource allocation” in the case of cytoplasmic metabolite exchange between cells (2–5). The efficiency with which a group of cells exploits resources of their environment may increase as a consequence of cooperation. For instance, in some syntrophic microbial communities, the secretion of digestive enzymes by individual specialized cells changes local resource availability, which in turn can benefit neighboring cells (6, 7). Cooperation is associated with major transitions in the evolution of life through the emergence of new functions at a higher level of organization, such as multicellularity and sociality (2, 8). It is the basis of many complex traits in nature, including virulence in some pathogenic species. In bacterial pathogens, cooperative behaviors such as biofilm formation, swarming motility, and the secretion of virulence factors are crucial to virulence and controlled by quorum sensing, the release and perception of small signal molecules (9–11). Distinct developmental and functional stages, interdependent for nutrition, were documented in the mycelium of wood-decay fungi in response to local environmental changes within host tissues (12–14). In the fungal meningitis pathogen Cryptococcus gattii, cooperation has also been associated with the emergence of outbreak strains that proliferate rapidly within host cells (15). Imaging and modeling approaches emphasized the importance of intrahyphal nutrient translocation for growth in Ascomycete and Basidiomycete fungi (14, 16). However, since the formulation of the cell theory in the 1830s, the eukaryotic cell has often been considered as an autonomous life unit (17). A number of processes such as the circadian clock, embryonic cell fate determination, and immunity can indeed be cell-autonomous in multicellular organisms (18). Furthermore, disorders such as cancers and neurodegenerative diseases result from the proliferation of one cell type detrimental to the whole organism (4). The extent to which cooperation contributes to complex traits, such as host colonization in pathogens with diverse infection strategies, remains largely unknown.

Natural selection shapes genes, cells, and organisms to promote their own evolutionary success at the expense of their competitors. In this context, the emergence of cooperation is favored only when specific conditions are met (8). Genetic proximity between individuals, also designated as kin selection, is a well-established condition that increases the likelihood of cooperative phenotypes to evolve (1). Cooperation is also expected to prevail if cooperative modules are segregated in space and interact preferentially with specific neighbors (19, 20). Spatial theory suggests that the spatial structure of interacting populations can promote cooperation (21). Experiments in synthetic yeast colonies showed that spatial expansion promotes the evolution of cooperation locally (22). Variation, in the form of mutations, phenotypic changes, or local environment properties, is another factor favoring task specialization and cooperation.

**Significance**

Cooperation between specialized cells and organisms supports complex biological functions, from the colonization of unfavorable environments to the formation of organs and sociality. Some bacterial pathogens are known to rely on cooperation between individuals and species for efficient colonization of their host and the onset of disease. We examined the regulation of genes in cells from different parts of a fungal plant pathogen and found evidence for cooperation between these fungal cells. We further show that cooperation between fungal cells is particularly important for the efficient colonization of resistant plants. These findings establish cooperation as a mechanism supporting disease caused by fungal pathogens that should be taken into account in the design of disease management strategies.

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(23). For instance, the random and progressive introduction of metabolic auxotrophies in yeast populations leads to the spontaneous establishment of metabolically cooperating communities (5).

The white mold fungus *Sclerotinia sclerotiorum* has a devastating impact on crops such as rapeseed and soybean, threatening food security worldwide (24). Like other fungal pathogens, it derives energy from its hosts for growth and reproduction. To do so, it produces a mycelium consisting of interconnected linear hyphae that colonize host tissues intercellularly and secrete proteins and metabolites that modify host cells’ physiology (25, 26). *S. sclerotiorum* has a nonotrophic lifestyle and rapidly causes host cell death and host tissue maceration at the center of the infected area, while the margin of the mycelium spreads through living plant cells (27). The morphology of *S. sclerotiorum* hyphae also differs between the margin and the center of the colony during plant infection (28). These hyphae are exposed to a heterogeneous and variable environment, with the apex of the mycelium growing through intact host tissues while its center is surrounded by a host-derived substrate with distinct chemical and physical properties. *S. sclerotiorum* therefore presents several properties favorable to the establishment of cooperative phenotypes that prompted us to test whether virulence is associated with intercellular cooperation. Here we show that compartments of *S. sclerotiorum* invasive hyphae cooperate through resource allocation and division of labor to facilitate host colonization, supporting a role for intercellular cooperation in fungal virulence.

**Results**

Transcriptome Analyses Reveal the Spatial Organization of *S. sclerotiorum* Invasive Hyphae. To test experimentally for spatial organization in *S. sclerotiorum* invasive hyphae, we analyzed the global transcriptome of *S. sclerotiorum* mycelium central and apical areas by RNA sequencing (RNA-seq). We harvested in triplicate separate areas corresponding to the central and apical zones of fungal mycelium grown on solid medium in vitro and on *Arabidopsis thaliana* plants (Fig. 1A). The first two principal components of the global gene expression dataset accounted for ∼77.6% of the total variation in read count (Fig. 1B). The major driver of differential gene expression was growth conditions (in vitro versus *in planta*, accounting for 53.7% of total variance), followed by the mycelium area (central versus apical, accounting for 23.9% of the total variance); these effects greatly exceeded the effects from biological replicates. The amplitude of principal component 2, which separates samples based on colony areas, was largely determined by the samples collected *in planta*, suggesting that the transcriptional differentiation of mycelium center and apex was stronger *in planta*. We used gene expression in *S. sclerotiorum* grown in liquid medium determined by RNA-seq as a reference to determine gene induction folds. We identified a total of 1,133 genes (10.2% of the genome) induced fourfold either *in planta* on solid medium in vitro (Fig. 1C and Dataset S1). Only 54 genes (4.7%) were induced in all four conditions, and 218 genes (19.2%) were induced both in vitro and *in planta*.

Fig. 1. Global transcriptome analyses reveal the spatial organization of fungal cells *in planta*. (A) Sampling strategy for the analysis of spatial differentiation of *S. sclerotiorum* global transcriptome in vitro and *in planta* on *A. thaliana* leaves. (B) Individual factor map of a PCA on *S. sclerotiorum* top differentially expressed genes. Ellipses show 95% confidence interval for each sample, calculated based on shuffling gene expression 100 times between three biological replicates. (C) Venn diagram showing the distribution of genes induced at least fourfold compared with in vitro liquid medium growth condition. (D) Quantitative RT-PCR validation of the expression profile for three genes specifically expressed at the mycelium apex *in planta* (plain green boxes) and three genes specifically expressed at the center of mycelium *in planta* (plain brown boxes). Error bars show SD of the mean for three biological replicates. (E) Enrichment analysis for PFAM domain annotations among differentially expressed genes in each sample type. The x axis shows enrichment in vitro versus *in planta* and the y axis shows enrichment in mycelium center versus apex. Annotations are colored according to the minimum enrichment P value (P value < 0.018 are shown), and sized according to the cumulated number of normalized sequence reads.
A total of 288 genes (25.5%) were induced in vitro only, and 627 (55.4%) were induced in planta only, indicating that plant colonization requires extensive specific transcriptional reprogramming. A total of 553 genes (48.8%) were induced both at the center and at the apex of colonies, indicating that a majority of transcriptional reprogramming is area-specific. Among genes induced in the mycelium center only (306 in total), there were only four (1.3%) induced both in vitro and in planta, further illustrating specific and local transcriptional reprogramming of *S. sclerotiorum* during *A. thaliana* colonization.

To confirm patterns of local cellular specialization in *S. sclerotiorum* during plant colonization, we analyzed the expression pattern of selected genes by quantitative RT-PCR. We found that the expression of some fungal genes was restricted to cells of the apex or the center of the mycelium (Fig. 1D). The Scsle1g002960 peptidase and the Scsle1g006490 and Scsle0g007010 oxidoreductases showed, respectively, a 21.8-, 10.2-, and 23.5-fold higher expression at the apex of *S. sclerotiorum* mycelium in planta than in any other tested condition. Conversely, the Scsle1g010740 aldo-keto reductase, the Scsle0g006420 CO-esterase, and the Scsle0g005560 aminotransferase showed, respectively, 8.1-, 13.5-, and 7.3-fold higher expression at the center of *S. sclerotiorum* mycelium in planta than in any other tested condition. This validates the specific and local activation of some *S. sclerotiorum* genes during host colonization. To document the nature of specific and local transcriptional programs, we grouped induced genes according to their predicted function. We analyzed in which condition these functions were predominantly activated by testing for enrichment in planta compared with in vitro and at the margin compared with the center of the mycelium (Fig. 1E). We found that a number of key metabolic functions for host colonization were performed locally. Functions strongly enriched at the apex of the mycelium include pectinesterases and polygalacturonases (GH28) that degrade pectic homogalacturonans, β-1,4-endoglucanases (GH12) and β-1,4-xylosides (GH11), serine proteases (peptidases S10, S28, and prokumamolysin), and cytochrome P450, notably involved in fungal toxin biosynthesis (29) (SI Appendix, Fig. S1). Functions strongly enriched at the center of the mycelium include arabinofuranosidases, β-1,4-xylosides (GH43), and α-1,4-galactosidase that cleave carbohydrate backbones, cutinases, muconate lactonizing enzyme (MR_MLE) involved in the breakdown and assimilation of aromatic compounds, and dihydrolipoamide dehydrogenase (DHDPS), a key enzyme in lysine biosynthesis. We conclude that host colonization triggers division of labor in *S. sclerotiorum* invasive hyphae with specific gene expression patterns at the apex and the center of the mycelium.

**Division of Labor Drives Metabolic Heterogeneity Along *S. sclerotiorum* Invasive Hyphae.** To assess quantitatively the impact of division of labor on major cell functions in *S. sclerotiorum* hyphae, we performed a flux balance analysis (FBA) using experimentally determined transcriptomes. To this end, we first reconstructed a genome-scale metabolic model (GEM) of *S. sclerotiorum* (Fig. 2A and Dataset S2). Our model includes a metabolic module with 1,277 reactions of the central metabolism, nutrient uptake pathways, degradation pathways, and the biosynthesis pathways of the major components of the fungal biomass. The model also includes a plant cell-wall degradation module with 218 reactions catalyzed by secreted plant cell-wall-degrading enzymes. Altogether, the *S. sclerotiorum* GEM encompasses 1,495 unique reactions associated with 1,039 genes, comparable to validated yeast and filamentous fungi models (30, 31). To evaluate the performance of this GEM, we determined experimentally *S. sclerotiorum* growth on 288 metabolites by using Biolog phenotype microarrays and compared it with model predictions (Fig. 2B and Dataset S3). The model correctly predicted the observed behavior on 252 metabolites (73 used as a C or N source and 179 not used), reaching 88% accuracy, 83% sensitivity, and 78% precision for axenic growth.

Next, we used FlexFluxOmics (32) to assess metabolic fluxes through *S. sclerotiorum* GEM based on global gene expression determined experimentally in vitro and in planta. We normalized flux distributions by fixing the biomass production fluxes measured experimentally for each condition. In *S. sclerotiorum* cells growing in vitro, 536 to 555 reactions supported nonnull fluxes, covering 79 to 86 distinct metabolic pathways. During *A. thaliana* colonization, 592 to 610 reactions supported nonnull fluxes in *S. sclerotiorum* cells, covering 101 to 103 distinct metabolic pathways (Dataset S4). Among these pathways, 16 supported significantly higher fluxes in apex compared with center cells during *A. thaliana* colonization, including detoxification pathways (cyanide, reactive oxygen species, and nitrile detoxification), biosynthesis of organic acids (oxalate and acetate metabolism and pH homeostasis), carbohydrate metabolism (glycolate and citric acid cycles and C1 metabolism), and secondary metabolite trafficking (peroxisomal and toxin transport) (Fig. 2C). Conversely, six pathways supported significantly lower fluxes in apex compared with center cells during *A. thaliana* colonization, related to energy and reduced cofactor production (glycolysis and pentose phosphate pathways with the respective upstream pathways, pyruvate metabolism, and arabinose degradation). By contrast, there was no pathway supporting significantly higher fluxes in apex compared with center cells during in vitro growth. This observation is consistent with apex and center cells performing complementary metabolic functions during plant colonization. In this analysis, predictions for pathways known to be regulated posttranscriptionally should be taken with caution as these regulation events are not included in the model. For instance, we identified higher fluxes through glycogen metabolism in center cells, but we could not determine whether biosynthesis or degradation of glycogen was activated since these pathways are controlled by phosphorylation events. To further characterize metabolic heterogeneity, we calculated total carbon metabolic uptake and carbon secretion for major virulence functions (oxalic acid production and protein secretion) by *S. sclerotiorum* apex and center cells (Fig. 2D). During in vitro growth, carbon metabolic uptake was similar in apex and center cells (20.0 ± 0.5 and 20.4 ± 1.9 mmol·g−1·h−1), as well as carbon secretion for virulence functions (0.80 ± 0.05 and 0.67 ± 0.35 mmol·g−1·h−1) in apex and center cells, respectively, *P* = 0.56 (Dataset S4). The total carbon metabolic uptake was also similar in apex and center cells during plant colonization (19.9 ± 0.7 and 18.4 ± 3.4 mmol·g−1·h−1, respectively, *P* = 0.52). However, the flux of carbon secretion for virulence was 8.5-fold lower in center cells (0.34 ± 0.06 mmol·g−1·h−1) than in apex cells (2.89 ± 0.34 mmol·g−1·h−1, *P* = 0.0047), suggesting that center cells may store carbon and transfer it toward apex cells, as observed in some filamentous fungi (14, 16, 33, 34). Metabolic exchange between center and apex cells prompted us to test whether the continuity between hyphal cells is required for successful host colonization.

**Continuity Between Central and Apical Compartments of *S. sclerotiorum* Hyphae Facilitates Host Colonization.** Molecules such as glucose can be transported over long distance in fungal hyphae (33, 34). After the ablation of subapical hyphal compartments, the growing hyphal tip is rapidly isolated from damaged cells by Woronin bodies and continues to grow in isolation (35, 36). To test whether the continuity between hyphal cells plays a role in host colonization, we inoculated *A. thaliana* host plants with *S. sclerotiorum* and measured the hyphal linear growth in planta (Fig. 3A). The average growth rate was 0.37 mm·h−1 between 9 and 13 h postinoculation (hpi). In untouched disease lesions, the hyphal growth rate remained unchanged between 13 and 17 hpi. When we performed at 13 hpi a circular ablation of the center of the lesion (and fungal hyphae it contains), the hyphal growth rate decreased on average by 38% (Student’s t test, *P* = 2.7e−59). To exclude the negative
Division of labor drives metabolic heterogeneity along *S. sclerotiorum* hyphae during host colonization. (A) Overview of reactions and metabolites included in *S. sclerotiorum* GEM highlighting the major cellular compartments and the extracellular compartment. The table describes features of GEM with the number of reactions and genes associated with the metabolic module and plant cell wall (PCW) degradation modules. (B) Experimental evaluation of the performances of *S. sclerotiorum* GEM using Biolog phenotype microarrays. (B, Top) Contingency table of the modeled (rows) and experimentally determined (columns) metabolic capacities for *S. sclerotiorum*. The squares are sized according to the number or carbon sources; correct predictions are shown in green and incorrect predictions in red. (B, Bottom) Bars show performance statistics. (C) Radar plots showing metabolic pathways supporting significantly different carbon fluxes in center and apex cells during *A. thaliana* colonization. Shaded areas show mean fluxes calculated by FBA on three biologically independent global transcriptome sequencings, with values for individual replicates shown as dots.

Division of Labor and Resource Allocation Increase Fungal Invasive Growth Synergistically. Previous work reported a role for multicellularity in fungal growth in vitro (37). Our results demonstrate that intercellular cooperation in *S. sclerotiorum* hyphae promotes host colonization. To estimate the relative impact of intercellular cooperation on *S. sclerotiorum* fitness in various environments, we modeled invasive growth as a proxy for fitness (38, 39) in simulations with environments of various resistance levels. For this, we developed a biophysical multicell model of the growing hyphae during host infection. Following observations on fungal model systems (40, 41), we modeled the hyphae as a stack of cells with a single dividing apical cell (Fig. 4I). In light of our FBA results, we considered that energetic costs associated with the production of virulence factors decreased upon the death of the host cells, corresponding to the ability of the fungus to overcome host resistance (detoxification of host environment, degradation of host-derived compounds, and host manipulation). The diffusion of metabolites through the hyphae tends to minimize the gradient of metabolites between cells, according to the second law of thermodynamics. We considered the difference of energetic costs associated with virulence factors in apical and central cells of the hyphae as a measure of the degree of division of labor, and a proxy for the relative level of host susceptibility (from 0.0 fully resistant to 1.0 fully susceptible). We considered the rate of cytoplasmic diffusion between hyphal cells as a proxy for the degree of resource allocation among hyphal cells. The
model allowed simulating hyphal growth according to (i) the relative susceptibility of host compartments, (ii) the degree of resource allocation between hyphal cells, and (iii) division of labor in the hyphae. We simulated hyphal length (Fig. 4A, y axis) in hosts with contrasted susceptibility levels between 0 and 1,000 iterations of the model (x axis) to monitor invasive growth. On a susceptible host (susceptibility 0.9; Fig. 4A, Left), the virtual hyphae grows rapidly, reaching a length >100 in 1,000 iterations. There is no apparent impact of suppressing division of labor in the model (dotted blue line). When host susceptibility is lower (0.21; Fig. 4A, Right), hyphal growth is reduced, reaching a length of 18 in 1,000 iterations, and can be completely suppressed in the absence of division of labor (dotted blue line).

To explore the effect of resource allocation and division of labor on hyphal growth in various environmental conditions, we calculated the gain of invasive growth as the ratio between the number of hyphal cells reached after 1,000 iterations of the model with and without resource allocation and division of labor (Fig. 4B and C, color scale). For a given host susceptibility, the gain of growth increased with the level of resource allocation (y axis). In the absence of division of labor (Fig. 4B), resource allocation conferred a significant gain on susceptible hosts, and there was no growth on hosts with susceptibility <0.3. Division of labor extended fungal host range by enabling growth on very resistant hosts (susceptibility <0.3) that would otherwise not be colonized (Fig. 4C). On resistant hosts, division of labor and resource allocation operated in synergy to provide high fitness gain. In S. sclerotiorum GEM, C fluxes toward virulence calculated during A. thaliana colonization corresponded to a level of susceptibility of 0.21. We estimated a 38 to 45% growth reduction when apex cells were disconnected from the center cells (Fig. 3 and SI Appendix, Fig. S3), corresponding to a resource allocation level of ~0.6. Based on our multicell model, we predict that division of labor was essential to allow A. thaliana colonization by S. sclerotiorum (Fig. 4C). A major prediction from the model is that the gain in fungal invasive growth conferred by cooperation is generally higher on resistant hosts than on susceptible hosts. In agreement, we found that the host plant Nicotiana benthamiana is more susceptible to S. sclerotiorum than A. thaliana (fungal growth rate 0.6 mm h⁻¹; SI Appendix, Fig. S3), and that ablation of central cells caused a stronger fungal growth penalty on A. thaliana (~44.9%) than on N. benthamiana (~19.4%; P = 0.007; SI Appendix, Fig. S3). We conclude that at the hyphae level the benefit from intercellular cooperation increases with the rate of resource allocation and the level of host resistance.

Discussion
Recent discoveries have considerably expanded the repertoire of molecular determinants known to underlie microbial virulence at the cellular level (42, 43). For instance, fungal pathogens secrete up to hundreds of effector proteins and small metabolites to facilitate host colonization (44). Several reports suggest that effector secretion occurs locally in invasive hyphae (45, 46). However, despite frequent spatial heterogeneity of diverse symptoms, evidence for a role of filamentous pathogen multicellular organization in virulence is lacking. Using systems biology and modeling approaches, our analyses demonstrate how local transcriptional programs in fungal cells favor the cooperative growth of multicellular hyphae in hostile host environments. Complex processes such as division of labor require mathematical modeling to generalize conclusions drawn from experiments (47, 48). Our study provides a framework combining experimental and modeling approaches to test for the adaptive significance of division of labor in many diseases caused by filamentous microbes. Fungal processes contributing to intercellular cooperation could also be exploited as novel targets to control diseases.

The division of tasks among specialized group members is an important aspect of major transitions in evolution that occurred many times independently (49, 50). Despite this, the evolutionary and ecological drivers leading to the establishment of division of labor remains an active area of research. Several approaches have identified several factors favorable to the evolution of division of labor and cooperation. At the population level, genetic relatedness, reciprocity, and group selection can lead to the emergence of cooperation (8). At the organism or multicellular level, the position of modules within an organism, existence of a fitness trade-off between two tasks, and synergistic interaction between modules favor division of labor (20). An experimental prisoner’s dilemma game in synthetic budding yeast colonies further demonstrated that spatial expansion is sufficient to drive enrichment in cooperators (22). Our analyses support the view that “colony growth alone” (14, 22), and the heterogeneity it creates in the environment, can promote cooperation in microbes. Pathogenic interactions with diverse hosts, increasing the range of environment encountered at the population level, could have favored the evolution of division of labor in fungi (13, 14). Synergistic interaction between metabolic pathways was shown to be favored when a pathway is of high metabolic burden or toxicity, highly complex, or relies on numerous extracellular steps (51). Several of these criteria are fulfilled in our study: S. sclerotiorum virulence involves the biosynthesis of complex toxins and secondary metabolites and the degradation of plant cell wall involves multiple extracellular enzymes (26, 52). In a previous study, we reported that fungal pathogens with a broad host range secrete more numerous and complex proteins than specialized fungi, likely increasing the metabolic burden of virulence in these species (53). These

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Fig. 3. Optimal growth of S. sclerotiorum during A. thaliana colonization requires central hyphal compartments. (A) Experimentally measured outward of semicircular ablation of the lesion center at 13 hpi (red). Lesion growth was mostly unaffected in untouched lesions (green), after transverse cut (yellow), and after distal cut (orange). Significance was determined by pairwise Welch t tests (P < 0.01). (B) Variation in the speed of lesion radial growth between 13 and 17 hpi by S. sclerotiorum measured outward of semicircular ablation of the lesion center (red), semicircular cut (yellow), and on the symmetric, untouched half of the same lesions (blue; see leaf diagrams). Significant differences between groups were determined by pairwise Welch t tests (***P < 0.01).

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observations suggest that selection for the emergence of division of labor would be higher in broad-host-range fungi and may be related to fungal lifestyles.

Pathogens' ability to colonize a host and trigger disease is determined largely by the virulence factors they express (54). The biosynthesis of virulence factors and their delivery to their site of action imposes a metabolic burden to pathogen cells. The existence of a trade-off between virulence and transmission, or virulence and within-host growth, is central to many theories on the evolution of parasites (55, 56). Trade-offs between virulence and transmission are also apparent in a growing number of experimental studies (57–59). Our results points toward division of labor as a strategy to mitigate virulence–transmission trade-offs in fungal pathogens through the optimization of resource usage and dampening toxicity of some fungal virulence factors and host defense compounds. The requirement for virulence factor biosynthesis comes in addition to the need for pathogen cells to grow and multiply to outcompete cooccurring microbes and disseminate efficiently. Cooccurring microbes and defective mutants may indeed benefit from secreted virulence factors produced by virulent pathogens manipulating host physiology in a “public-good” manner (60–62). In pathogenic public-good producers, reducing virulence, and thereby the ability to benefit from cooperation, may switch selection toward favoring cheats at high microbial density (63). The uneven distribution of nutrients made available by fungal pathogens may affect differently the growth of microbes forming the leaf microbiome. Consequently, disease management strategies exploiting virulence reduction may have undesired effects due to the cooperative nature of fungal virulence. Indeed, weakly virulent strains can alleviate rate–efficiency trade-offs on resource use and result in higher prevalence of virulent strains in natural populations (62). In addition, the silencing of secreted virulence factors may prove inefficient due to compensation through cooperative hyphal organization, increased natural selection on cooperative resource allocation, and resistance to host defense compounds, ultimately displacing natural population balance toward more aggressive strains.

**Materials and Methods**

**Plant and Fungus Material and Growth Conditions.** *A. thaliana* plants of accession Col-0 were grown in jiffy pots at 22 °C with a 9-h light period under a light intensity of 120 μmol·m⁻²·s⁻¹ for 4 wk before infection. *S. sclerotiorum* strain 1980 was subcultured on potato dextrose agar plates at 22 °C. For inoculations, a 5-mm-wide agar plug containing actively growing *S. sclerotiorum* mycelium was placed on the adaxial surface of leaves and plants were maintained at 80% humidity in Percival AR-41L3 chambers under the same day/night condition as for plant growth, in trays closed with plastic wrap to control for humidity.

**Transcriptomic Analyses.** *S. sclerotiorum* gene expression was analyzed in six growth conditions: (i) mycelium cultured in potato dextrose broth (PDB) as described in ref. 64, (ii) mycelium grown on minimum agar medium as described in ref. 64, (iii and iv) the center and margin of mycelium (colony diameter 25 mm) on potato dextrose agar plates, and (v and vi) the center and margin of disease lesion (lesion diameter 25 mm) on *A. thaliana* (Col-0 genotype). Samples were collected in three independent biological experiments.

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**Fig. 4.** Division of labor and resource allocation increase fungal invasive growth on resistant hosts in a multicell model. (A) Simulated fungal invasive growth (y axis) on hosts with contrasted susceptibility levels over 1,000 iterations of the model (Time, x axis). Simulations were run with division of labor (DOL, red line) and without DOL (blue dotted line) on a susceptible host (Left, level of susceptibility = 0.9) and a resistant host (Right, level of susceptibility = 0.21). Suppression of DOL abolished fungal growth on a host of susceptibility 0.21, which corresponds to observed susceptibility for *A. thaliana* Col-0. AU, arbitrary units. (B) Gain of fungal invasive growth (color scale) in the absence of DOL, defined as the ratio between the number of hyphal cells reached after 1,000 iterations of the model with and without division of labor (DOL, red line) and without DOL (blue dotted line) on a susceptible host (*A. thaliana*) observed. (C) Predicted growth gains are higher on resistant hosts (susceptibility between 0.1 and 0.3) than on susceptible hosts (green-yellow sector).
RNA extraction, sequencing, and reads mapping were performed as described in ref. 64. Normalized read count per gene and differential gene expression (DGE) was performed using the Bioconductor DESeq2 (version 1.8.2) package. Raw and normalized RNA-seq data have been deposited in the GEO database (accession nos. GSE108811 and GSE161194). Genes were considered significantly up-regulated when induced fourfold or greater compared with the control PDB condition, with a P value ≤0.01 (Dataset S1). Normalized read counts per gene for the top 500 most-expressed genes were shuffled between the three biological replicates 100 times for principal component analysis (PCA). PCA was performed using the factominR and factoextra R packages. For each PFAM domain, the number of differentially expressed genes was compared with the total number of genes in *S. sclerotiorum* v2 genome (S2) using a $\chi^2$ test implemented in R to determine enrichment P values.

Quantitative Real-Time PCR Analysis. RNA extraction was performed as described above. First-strand cDNA synthesis was performed using 1 μg of total RNA with an anchored oligo(dT) and the SuperScript III reverse transcriptase (Invitrogen), according to the manufacturer's instructions. Real-time PCR reactions were performed on a Light Cycler 480 thermocycler (Roche) at 60 °C annealing temperature using 1:10 diluted cDNAs and LightCycler 480 SYBR Green I Master (Roche) in a final reaction volume of 7 μL with primer pairs given in SI Appendix, Table S1. Transcript levels were normalized to the reference gene *A. thaliana* (At2g33370) and calculated using the 10−ΔΔCt method. The relative expression levels were calculated as the relative change in the expression level of the target gene compared to the reference gene. The fold change was calculated using the formula 2^−ΔΔCt.

Ablation Assays and Linear Hyphal Growth Measurements. Detached leaves of 4-wk-old *A. thaliana* plants (Col-0 accession) and *N. benthamiana* plants were inoculated with a 3-mm-wide agar plug containing actively growing *S. sclerotiorum* 1980. Leaves were maintained on wet paper towel in Petri dishes for the duration of the experiment. Pictures were taken every 30 min between 10 and 20 hpi. At 13 hpi, a circular area colonized by the fungus of 3- to 4-mm radius was removed with a scalpel from a subset of leaves (circular ablation). To control for wounding effects, transverse cuts through the colonized area or in distal parts of the leaves were applied with a scalpel to another subset of leaves (transverse and distal cuts). For each leaf and at each time point, the radius of colonized area was measured along four orthogonal directions with ImageJ version 1.51. This assay was repeated three times with similar results. The radial hyphal growth rate (in planta) was computed as the slope of the linear regression along measured radii. Growth rates were calculated during the 9- to 13-hpi interval and during the 13- to 17-hpi interval.

GEM Reconstruction and FBA. A draft model for *S. sclerotiorum* 1980 was obtained first by transfer from already reconstructed models of the Ascomycete fungi *Aspergillus fumigatus* CBS 513.88 (66) and *Aspergillus niger* CBS 513.88 (66) based on gene homology; see Peyraud et al. (58) for detailed information. Briefly, reactions from available models including genes with orthologs (>30% identity and >50% coverage) in *S. sclerotiorum* 1980 genome (S2) were collected. Gene orthology was further verified using HMMer (Peyraud et al., Fig. S2). Transcript levels were normalized to the reference gene *A. thaliana* (At2g33370) and calculated using the 10−ΔΔCt method. The relative expression levels were normalized using experimentally measured growth rates.

Phenotype Microarrays. Phenotype microarrays analysis was performed using Biolog plates PM1-3. To produce homogeneous fractions of *S. sclerotiorum* mycelium grown in PDB at 24 °C for 24 h at 180 rpm then filtered on 100 μm. The filter membranes containing fractions that grew above 100 μm were washed with minimal medium with no carbon source, and the resulting mycelium suspension was diluted to 62% transmittance. α-glucose at 20 mM was added for the inoculation of plates PM3. Plates were inoculated following the manufacturer’s instructions with the following modifications: the volumes of mycelium suspensions used to obtain a final 24 mL of inoculation solution were 1.5 mL for PM1 and 2, 0.3 mL for PM9, and 3 mL for the other plates. Measurements were recorded on an Omnilog reader (Biolog) every 30 min for 7 d. Six biological replicates were run for each plate. The data were analyzed using the R package OPM (Dataset S3).

Multicell Hyphal Model. A coarse-grain model was designed to assess the spatiotemporal effect of resource allocation and division of labor. The hyphae were modeled as a growing cylinder composed of a row of cells of identical volume. Every cell of the hyphae produced virulence factors from carbon uptake. Turgor pressure in cells was assumed constant and did not constrain growth. Under these conditions, growth depends only on the speed of cell-wall biosynthesis (73, 74), which occurs only in the apical cell (40). In the same manner, the colonized host was modeled as a stack of cells of identical volume containing carbon. Under the latter assumptions, the net variation of carbon uptake for every cell i different from the apical one was:

$$\frac{dN_{fungi}}{dt} = \frac{dN_{host-fungi}}{dt}$$

whereas in the apical cell the production rate of growth factor was:

$$\frac{dN_{growth-aphial}}{dt} = \frac{dN_{fungi}}{dt}$$

with $N_{fungi}$ the quantity of nutrients uptaken by the cell i, $N_{growth-aphial}$ the quantity of virulence factor produced by the cell i, and $N_{growth-aphial}$ the quantity of cell wall polysaccharides produced by the apical cell. $dN_{host-fungi}/dt$ and $dN_{fungi}/dt$ were the speeds at which carbon was extracted from host and exchanged with neighboring cell. They were modeled by Fick's law:

$$\frac{dN_{fungis}}{dt} = D_{host-fungi} (N_{host} - N_{fungis})$$

with $D_{host-fungi}$ and $D_{fungis}$ coefficients of diffusion (in [0,1/2]). Growth of the hyphae was discretized. When carbon quantity in the apical cell overcame a threshold ($N_{growth}$), a new cell was produced. The resistance of a host cell was assumed to vanish when a certain amount of virulence
factor was secreted by the associated cell of the pathogen. When the resistance of a host cell vanished, nutrients were free to diffuse to the hyphae and the local metabolism of the hyphae switched from apical to central. On the basis of results provided by metabolic reconstruction, the rate of production of virulence factors was assumed proportional to the carbon uptake of the cell:

$$\frac{dN_v}{dt} = \mu_i \left( \frac{dN_{host,fungal}}{dt} + \frac{dN_{host,apical}}{dt} \right).$$

[5]

The $\mu_i$ parameter had two values depending on the cell location (center or apex). The parameter $\mu_i$ was introduced to describe the bias in resource allocation in the production of virulence factors between center and apex of the colony:

$$\mu_{i\text{ center}} = \mu_{i\text{ periphery}} < \mu_i < 1.$$  

[6]

**Simulation of Hyphal Invasive Growth.** We assumed an infinite quantity of carbon in the host cell which involved a constant speed of uptake $(N_{\text{host,fungal}})$ on the basis of results provided by GEM analysis, we assumed constant resistance of a host, identical for cells in center and in apical region of the hyphae, and not dependent on host resistance. The speed of uptake was set to 1 mmol per simulation time unit. Increase in host resistance was modeled by the increase in the bias of resources allocation in the production of virulence factor $\mu_i$. $\mu_{i\text{ center}}$ was assumed constant and set to 0.03. Low host resistance was associated with $\mu_i = 1$, whereas high host sensitivity was associated with $\mu_i$ close to 0 (Eq. 6). The level of resource allocation between cells was modeled by the dimensionless number

$$\alpha = \frac{\mu_{\text{host,fungal}}}{0.5} \in [0,1].$$

[7]

The sensitivity of hyphal growth to resource allocation was varying by $\alpha$. The sensitivity of hyphal growth to division of labor was tested by varying $\frac{1}{\mu_i}$ to the same extent as $\alpha$. In every test set, simulations were initiated with hyphae composed of an apical cell. Simulations were computed over 1,000 iterations and length of the modeled hyphae was reported. Gain for a given host susceptibility $S$ was computed as

$$G_i(S) = \frac{\Delta L_{\alpha L}}{\Delta L_{\alpha_0}} - 1 = \frac{\mu_{\text{host,apical}} - \mu_{\text{host,fungal}}}{\mu_{\text{host,apical}}} \in [0,1].$$

[8]

The model implemented in Python 2.7 is available at https://github.com/QIPteam/intercellularcooperation.

**Data Availability.** Raw and normalized RNA-seq data have been deposited in the GEO database (accession nos. GSE106811 and GSE116194). Differentially expressed genes, GEM files, and phenotype microarray data are provided in Datasets S1-S6.

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1. Hamilton WD (1963) The evolution of altruistic behavior. *Am Nat* 97:354–366.
2. West SA, Fisher RM, Gardner A, Kiers ET (2015) Major evolutionary transitions in individuality. *Proc Natl Acad Sci USA* 112:10112–10119.
3. West SA, Cooper GA (2016) Division of labour in microorganisms: An evolutionary perspective. *Nat Rev Microbiol* 14:716–723.
4. Aktipis CA, et al. (2015) Cancer across the tree of life: Cooperation and cheating in bacteria. *Philos Trans R Soc B Biol Sci* 370:20140219.
5. Campbell K, et al. (2015) Self-establishing communities enable cooperative metabolic exchange in a eukaryote. *eLife* 4:e09943.
6. Koschwanzer JH, Foster KR, Murray AW (2011) Sucrose utilization in budding yeast as a model for the origin of undifferentiated multicellularity. *PLoS Biol* 9:e1001122, and erratum (2011) 9:e10.1371/annotation0903bbabd01d20–46a5b1231e2229ec6651.
7. Morris BEL, Henneberger R, Huber H, Moissi-Eichinger C (2013) Microbial syntrophy: Interaction for the common good. *FEMS Microbiol Rev* 37:384–406.
8. Nowak MA (2006) Five rules for the evolution of cooperation. *Science* 314:1560–1563.
9. Griffin AS, West SA, Buckland A (2004) Cooperation and competition in pathogenic bacteria. *Nature* 430:1040–1047.
10. Diggle SP, Griffin AS, Campbell GS, West SA (2007) Cooperation and competition in quorum-sensing bacterial populations. *Nature* 450:411–414.
11. Antunes LOM, Ferreira RBR, Buckner MMC, Finlay BB (2010) Quorum sensing in bacterial virulence. *Microbiology* 156:2271–2282.
12. Rayner ADM (1991) The phytopathological significance of mycelial individualism. *Annu Rev Phytopathol* 29:305–323.
13. Fricker MD, Heaton LLM, Jones NS, Boddy L (2017) The mycelium as a network. *The Fungal Kingdom*, eds Heismann J, et al. (American Society of Microbiology, Washington, DC), pp 355–367.
14. Voelz K, et al. (2014) ‘Division of labour’ in response to host oxidative burst drives a fatal Cryptococcus gattii outbreak. *Nat Commun* 5:5194.
15. Bebber DP, et al. (2007) Imaging complex nutrient dynamics in mycelial networks. *Fungi Environi* 2:31–32.
16. Balukta F, Volkman D, Barlow PW (2004) Eukaryotic cells and their cell bodies: Cell theory revisited. *Ann Bot* 94:9–32.
17. Randow F, MacMicking JD, James LC (2013) Cellular self-defense: How cell-autonomous immunity protects against pathogens. *Curr Opin Immunol* 29:253–265.
18. Baluska F, Volkmann D, Barlow PW (2004) Eukaryotic cells and their cell bodies: Cell theory revised. *Nat Rev Microbiol* 10:417–443.
19. Bleichrodt RJ, Hulsman M, Wösten HAB, Reinders MJT (2015) Switching from a uniaxial to a multicellular division of labor in fungal colonies. *Nat Commun* 6:325–335.
20. Diggle SP, Griffin AS, Campbell GS, West SA (2007) Cooperation and competition in quorum-sensing bacterial populations. *Nature* 450:411–414.
21. Antunes LOM, Ferreira RBR, Buckner MMC, Finlay BB (2010) Quorum sensing in bacterial virulence. *Microbiology* 156:2271–2282.
22. Rayner ADM (1991) The phytopathological significance of mycelial individualism. *Annu Rev Phytopathol* 29:305–323.
23. Fricker MD, Heaton LLM, Jones NS, Boddy L (2017) The mycelium as a network. *The Fungal Kingdom*, eds Heismann J, et al. (American Society of Microbiology, Washington, DC), pp 355–367.
24. Voelz K, et al. (2014) ‘Division of labour’ in response to host oxidative burst drives a fatal Cryptococcus gattii outbreak. *Nat Commun* 5:5194.
25. Bebber DP, et al. (2007) Imaging complex nutrient dynamics in mycelial networks. *Fungi Environi* 2:31–32.
26. Aktipis CA, et al. (2015) Cancer across the tree of life: Cooperation and cheating in bacteria. *Philos Trans R Soc B Biol Sci* 370:20140219.
27. Campbell K, et al. (2015) Self-establishing communities enable cooperative metabolic exchange in a eukaryote. *eLife* 4:e09943.
28. Nowak MA (2006) Five rules for the evolution of cooperation. *Science* 314:1560–1563.
29. Griffin AS, West SA, Buckland A (2004) Cooperation and competition in pathogenic bacteria. *Nature* 430:1040–1047.
30. Diggle SP, Griffin AS, Campbell GS, West SA (2007) Cooperation and competition in quorum-sensing bacterial populations. *Nature* 450:411–414.
31. Antunes LOM, Ferreira RBR, Buckner MMC, Finlay BB (2010) Quorum sensing in bacterial virulence. *Microbiology* 156:2271–2282.
32. Rayner ADM (1991) The phytopathological significance of mycelial individualism. *Annu Rev Phytopathol* 29:305–323.
33. Fricker MD, Heaton LLM, Jones NS, Boddy L (2017) The mycelium as a network. *The Fungal Kingdom*, eds Heismann J, et al. (American Society of Microbiology, Washington, DC), pp 355–367.
34. Voelz K, et al. (2014) ‘Division of labour’ in response to host oxidative burst drives a fatal Cryptococcus gattii outbreak. *Nat Commun* 5:5194.
35. Bebber DP, et al. (2007) Imaging complex nutrient dynamics in mycelial networks. *Fungi Environi* 2:31–32.
36. Aktipis CA, et al. (2015) Cancer across the tree of life: Cooperation and cheating in bacteria. *Philos Trans R Soc B Biol Sci* 370:20140219.
37. Campbell K, et al. (2015) Self-establishing communities enable cooperative metabolic exchange in a eukaryote. *eLife* 4:e09943.
38. Nowak MA (2006) Five rules for the evolution of cooperation. *Science* 314:1560–1563.
49. Szathmáry E (2015) Toward major evolutionary transitions theory 2.0. Proc Natl Acad Sci USA 112:10104–10111.
50. Simpson C (2012) The evolutionary history of division of labour. Proc R Sci 279:116–121.
51. Tsoi R, et al. (2018) Metabolic division of labor in microbial systems. Proc Natl Acad Sci USA 115:2526–2531.
52. Derbyshire M, et al. (2017) The complete genome sequence of the phytopathogenic fungus Sclerotinia sclerotiorum reveals insights into the genome architecture of broad host range pathogens. Genome Biol Evol 9:593–618.
53. Badet T, et al. (2017) Codon optimization underpins generalist parasitism in fungi. eLife 6:e22472.
54. Schulze-Lefert P, Panstruga R (2011) A molecular evolutionary concept connecting nonhost resistance, pathogen host range, and pathogen speciation. Trends Plant Sci 16:117–125.
55. Ebert D (1998) Experimental evolution of parasites. Science 282:1432–1435.
56. Alizon S, Hurford A, Mideo N, Van Baalen M (2009) Virulence evolution and the trade-off hypothesis: History, current state of affairs and the future. J Evol Biol 22:245–259.
57. Thrall PH, Burdon JJ (2003) Evolution of virulence in a plant host-pathogen metapopulation. Science 299:1735–1737.
58. Peyraud R, Cottret L, Marmiesse L, Gouzy J, Genin S (2016) A resource allocation trade-off between virulence and proliferation drives metabolic versatility in the plant pathogen Ralstonia solanacearum. PLoS Pathog 12:e1005939.
59. Bruns E, Carson ML, May G (2014) The jack of all trades is master of none: A pathogen’s ability to infect a greater number of host genotypes comes at a cost of delayed reproduction. Evolution 68:2453–2466.
60. Belhaj K, et al. (2017) Parallel evolution of the POQR prolyl oligopeptidase gene conferring plant quantitative disease resistance. PLoS Genet 13:e1007143.
61. Diard M, et al. (2013) Stabilization of cooperative virulence by the expression of an avirulent phenotype. Nature 494:353–356.
62. Lindsay RJ, Kershaw MJ, Pawlowska BJ, Talbot NJ, Gudelj I (2016) Harbouiring public good mutants within a pathogen population can increase both fitness and virulence. eLife 5:e18678.
63. Lindsay RJ, Pawlowska BJ, Gudelj I (2018) When increasing population density can promote the evolution of metabolic cooperation. ISME J 12:849–859.
64. Badet T, et al. (2017) Parallel evolution of the POQR prolyl oligopeptidase gene conferring plant quantitative disease resistance. PLoS Genet 13:e1007143.
65. Heavner BD, Smallbone K, Price ND, Walker LP (2013) Version 6 of the consensus yeast metabolic network refines biochemical coverage and improves model performance. Database (Oxford) 2013:bat059.
66. Pel HJ, et al. (2007) Genome sequencing and analysis of the versatile cell factory Aspergillus niger CBS 513.88. Nat Biotechnol 25:221–231.
67. Remm M, Storm CE, Sonnhammer EL (2001) Automatic clustering of orthologs and paralogs from pairwise species comparisons. J Mol Biol 314:1041–1052.
68. Thiele I, Palsson BO (2010) A protocol for generating a high-quality genome-scale metabolic reconstruction. Nat Protoc 5:93–121.
69. Cottret L, et al. (2010) MetExplore: A web server to link metabolomic experiments and genome-scale metabolic networks. Nucleic Acids Res 38:W132–W137.
70. Horton P, et al. (2007) WoLF PSORT: Protein localization predictor. Nucleic Acids Res 35:W585–W587.
71. Petersen TN, Brunak S, von Heijne G, Nielsen H (2011) SignalP 4.0: Discriminating signal peptides from transmembrane regions. Nat Methods 8:785–786.
72. de Oliveira Dal’Molin CG, Quek L-E, Palfreyman RW, Brumley SM, Nielsen LK (2010) AraGEM, a genome-scale reconstruction of the primary metabolic network in Arabidopsis. Plant Physiol 152:579–589.
73. Barbacci A, Lahaye M, Magne net V (2013) Another brick in the cell wall: Biosynthesis dependent growth model. PLoS One 8:e74400.
74. Ali O, Traas J (2016) Force-driven polymerization and turgor-induced wall expansion. Trends Plant Sci 21:398–409.