

# Complete genome of the Medicago anthracnose fungus, Colletotrichum destructivum, reveals a mini-chromosome-like region within a core chromosome

Nicolas Lapalu, Adeline Simon, Antoine Lu, Peter-Louis Plaumann, Joëlle Amselem, Sandrine Pigné, Annie Auger, Christian Koch, Jean-Félix Dallery, Richard J O'Connell

# ▶ To cite this version:

Nicolas Lapalu, Adeline Simon, Antoine Lu, Peter-Louis Plaumann, Joëlle Amselem, et al.. Complete genome of the Medicago anthracnose fungus, Colletotrichum destructivum, reveals a minichromosome-like region within a core chromosome. Microbial Genomics, 2024, 10 (8). hal-04683720

# HAL Id: hal-04683720 https://hal.inrae.fr/hal-04683720v1

Submitted on 2 Sep 2024

**HAL** is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers. L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.



Distributed under a Creative Commons Attribution 4.0 International License

# <sup>1</sup> Complete genome of the Medicago anthracnose fungus,

- 2 *Colletotrichum destructivum*, reveals a mini-chromosome-like
- <sup>3</sup> region within a core chromosome
- 4 Nicolas Lapalu<sup>1§</sup>, Adeline Simon<sup>1§</sup>, Antoine Lu<sup>1</sup>, Peter-Louis Plaumann<sup>2</sup>, Joëlle Amselem<sup>3</sup>,
- 5 Sandrine Pigné<sup>1</sup>, Annie Auger<sup>1</sup>, Christian Koch<sup>2</sup>, Jean-Félix Dallery<sup>1#</sup>, Richard J. O'Connell<sup>1#</sup>
- 6 <sup>1</sup> Université Paris-Saclay, INRAE, UR BIOGER, 91120 Palaiseau, France
- 7 <sup>2</sup> Division of Biochemistry, Department of Biology, Friedrich-Alexander-Universität Erlangen-
- 8 Nürnberg, 91058 Erlangen, Germany
- 9 <sup>3</sup> Université Paris-Saclay, INRAE, URGI, 78000 Versailles, France
- 10 <sup>§</sup> These authors contributed equally.

**\* Corresponding authors:** Richard J. O'Connell, richard.oconnell@inrae.fr; Jean-Félix Dallery jean felix.dallery@inrae.fr.

- **Keywords:** fungal genomics; accessory chromosome; chromosome rearrangements; segmental
- 14 duplication; phytopathogenic fungus, *Medicago truncatula*.
- 15 **Repositories:** GEO GSE246592; NCBI BioProject PRJNA1029933.

16 Abbreviations: AT: acyltransferase domain; AR: accessory region; BDBH: bidirectional best hit; BGC: biosynthetic genes cluster; CAT: conidial anastomosis tubes; CAZyme: carbohydrate active enzyme; 17 18 CDS: coding sequence; CE: carbohydrate esterase; Chr: chromosome; DNA: deoxyribonucleic acid; GH: 19 glycoside hydrolase; GO; gene ontology; HCT: horizontal chromosome transfer; HPI: hours post-20 inoculation; KS: ketosynthase domain; LINE: long interspersed nuclear element; LTR: long terminal 21 repeats; MITE: miniature inverted-repeat transposable element; NRPS: non-ribosomal peptide 22 synthetase; PCA: principal component analysis; PCP: peptidyl carrier protein domain; PCR: polymerase 23 chain reaction; PFGE: pulsed-field gel electrophoresis; PKS: polyketide synthase; PL: polysaccharide 24 lyase; RBH: reciprocal best hit; RFP: red fluorescent protein; RNA: ribonucleic acid; SD: segmental 25 duplication; SMKG: secondary metabolism key gene; SMRT: single molecule real time; TE: transposable element; TIR: terminal inverted repeat; TPM: transcript per million. 26

## 27 <u>Abstract</u>

28 Colletotrichum destructivum (Cd) is a phytopathogenic fungus causing significant economic losses on

29 forage legume crops (*Medicago* and *Trifolium* species) worldwide. To gain insights into the genetic 1

This work is licensed under a CC-BY 4.0 license. <u>https://creativecommons.org/licenses/by/4.0/</u> A CC-BY public copyright license has been applied by the authors to the present document and will be applied to all subsequent versions up to the Author Accepted Manuscript arising from this submission, in accordance with the grant's open access conditions. 30 basis of fungal virulence and host specificity, we sequenced the genome of an isolate from *M. sativa* 31 using long-read (PacBio) technology. The resulting genome assembly has a total length of 51.7 Mb and 32 comprises 10 core chromosomes and two accessory chromosomes, all of which were sequenced from 33 telomere to telomere. A total of 15,631 gene models were predicted, including genes encoding 34 potentially pathogenicity-related proteins such as candidate secreted effectors (484), secondary 35 metabolism key enzymes (110) and carbohydrate-active enzymes (619). Synteny analysis revealed 36 extensive structural rearrangements in the genome of Cd relative to the closely-related Brassicaceae 37 pathogen, C. higginsianum. In addition, a 1.2 Mb species-specific region was detected within the 38 largest core chromosome of Cd that has all the characteristics of fungal accessory chromosomes 39 (transposon-rich, gene-poor, distinct codon usage), providing evidence for exchange between these 40 two genomic compartments. This region was also unique in having undergone extensive intra-41 chromosomal segmental duplications. Our findings provide insights into the evolution of accessory 42 regions and possible mechanisms for generating genetic diversity in this asexual fungal pathogen.

## 43 Impact statement

44 Colletotrichum is a large genus of fungal phytopathogens that cause major economic losses on a wide 45 range of crop plants throughout the world. These pathogens vary widely in their host specificity and 46 may have either broad or narrow host ranges. Here, we report the first complete genome of the alfalfa 47 (Medicago sativa) pathogen, Colletotrichum destructivum, which will facilitate the genomic analysis of 48 host adaptation and comparison with other members of the Destructivum species complex. We 49 identified a species-specific 1.2 Mb region within chromosome 1 displaying all the hallmarks of fungal 50 accessory chromosomes, which may have arisen through the integration of a mini-chromosome into a 51 core chromosome and could be linked to the pathogenicity of this fungus. We show this region is also 52 a focus for segmental duplications, which may contribute to generating genetic diversity for adaptive 53 evolution. Finally, we report infection by this fungus of the model legume, Medicago truncatula, 54 providing a novel pathosystem for studying fungal-plant interactions.

## 55 Data summary

All RNA-seq data were submitted to the NCBI GEO portal under the GEO accession GSE246592. *C. destructivum* genome assembly and annotation are available under the NCBI BioProject PRJNA1029933 with sequence accessions CP137305-CP137317.

59 Supplementary data (genomic and annotation files, genome browser) are available from the INRAE 60 BIOGER Bioinformatics platform (https://bioinfo.bioger.inrae.fr/). Transposable Elements consensus 61 sequences are also available from the French national data repository, research.data.gouv.fr with doi 62 10.57745/TOO1JS.

## 63 Introduction

- 64 The ascomycete fungal pathogen *Colletotrichum destructivum*, causes anthracnose disease on lucerne
- 65 (alfalfa, *Medicago sativa*) and *Trifolium* species and is responsible for significant economic losses on
- these forage legumes [1, 2]. Despite being isolated most frequently from members of the Fabaceae, *C*.
- 67 destructivum has occasionally been recorded from genera of the Asteraceae (Helianthus, Crupina),
- 68 Poaceae (*Phragmites*) and Polygonaceae (*Rumex*) [3, 4]. It has a worldwide distribution that includes
- 69 the USA, Canada, Argentina, Italy, Netherlands, Greece, Serbia, Morocco, Saudi Arabia, and Korea. C.

*destructivum* is a haploid fungus with no known sexual stage [3]. Previous reports of a sexual stage
 (*Glomerella glycines*) for soybean isolates of *C. destructivum* [5, 6] were based on incorrect
 identification of the soybean pathogen, which was recently shown to be *C. sojae* [7].

Over the last decade, the application of multi-locus molecular phylogeny approaches has revealed that 73 74 C. destructivum belongs to the Destructivum species complex, which contains 17 accepted taxa [3, 8]. 75 All these plant pathogenic species show distinct host preferences, spanning phylogenetically diverse 76 botanical families. An increasing number of species in the Destructivum complex have now been 77 genome sequenced, namely C. higginsianum [9, 10], C. tanaceti [11], C. lentis [12] and C. shisoi [8], 78 which cause disease on Brassicaceae, Tanacetum (Asteraceae), Lens (Fabaceae) and Perilla 79 (Lamiaceae), respectively. The clade therefore provides excellent opportunities for comparative 80 genomic studies on the genetic determinants of host adaptation.

- The availability of complete genome sequences is crucial not only for the analysis of large gene clusters, such as secondary metabolism biosynthetic gene clusters, but also for understanding fungal genome evolution. Complete or near-complete genome sequences have enabled the structure and dynamics of accessory mini-chromosomes to be analyzed in several *Colletotrichum* species [9, 13, 14]. The importance of mini-chromosomes for virulence on plant hosts has been demonstrated in several fungal pathogens including *Fusarium oxysporum* f.sp. *lycopersici* [15], *Magnaporthe oryzae* [16], *C. lentis* [12] and *C. higginsianum* [17].
- Here, we present the complete genome sequence and gene annotation of C. destructivum strain LARS 88 89 709, hereafter called Cd709, based on long-read sequencing with PacBio Single Molecule, Real-Time 90 (SMRT) Sequel technology. The resulting high-quality chromosome-level assembly allowed us to 91 perform comparative genomics with the close sister species, C. higginisianum, highlighting gene 92 content specificity and extensive genomic rearrangements. In particular, the genome showed evidence 93 of multiple segmental duplications, as well as the likely integration of a mini-chromosome into one 94 core chromosome. Although the origin of this integrated region remains to be determined, it displays 95 all the hallmarks of fungal mini-chromosomes. We also show for the first time that C. destructivum is 96 pathogenic, and completes its life-cycle, on the model plant Medicago truncatula, providing a new 97 tractable pathosystem in which both partners have been genome-sequenced.

## 98 Materials and Methods

## 99 Fungal and plant materials

The *C. destructivum* strains used in this study were originally isolated from *M. sativa* in Saudi Arabia (CBS 520.97, LARS 709) and Morocco (CBS 511.97, LARS 202) [2], and are hereafter called *Cd*709 and *Cd*202. The *C. higginsianum* strains used for comparative genome and chromosome analyses were IMI 349063A and MAFF 305635 [10, 17, 18], hereafter called *Ch*63 and *Ch*35, respectively. The fungi were cultured as described previously [18].

Seeds of nine *M. truncatula* accessions (Table S1) were provided by the INRAE Centre de Ressources Biologiques Medicago truncatula (UMR 1097, Montpellier, France), while *M. sativa* seeds were purchased from Germ'line SAS (France). *M. truncatula* seeds were first abraded with sandpaper and imbibed with water for 1 h before sowing in seed compost (Floragard Vertriebs-GmbH, Oldenburg, Germany), while *M. sativa* seeds were sown directly in the same compost. All plants were grown in a controlled environment chamber (23°C day, 21°C night, 12-h photoperiod, PPFR 110 µmol·m<sup>-2</sup>·s<sup>-1</sup>).

#### 111 Infection assays and microscopy

- 112 To test the susceptibility of *M. truncatula* accessions to *Cd*709, intact plants (17-days-old) were
- 113 inoculated by first immersing the above-ground parts in a solution of 0.01 % (v/v) Silwet to wet the
- leaves, then by immersion in a suspension of *C*. *destructivum* spores ( $2 \times 10^6$  ml<sup>-1</sup>). The inoculated plants
- were incubated in a humid box inside a controlled environment chamber (25°C, 12-h photoperiod, PPFR 40  $\mu$ mol m<sup>-2</sup> s<sup>-1</sup>). For microscopic examination, pieces of infected tissues were cleared with a 1:3
- 117 mixture of chloroform:ethanol for 1h, then with lactophenol for 30 min, before mounting on a
- 118 microscope slide in 70 % glycerol and imaging with a Leica DM5500 light microscope. Symptoms were
- 119 recorded at 4 dpi.

#### 120 Pulsed-field gel electrophoresis (PFGE) and Southern blotting

- 121 The plugs containing the conidial protoplasts for PFGE were prepared as previously described [17].
- 122 Pulsed-field gel electrophoresis (Bio-rad CHEF-DR II system) was performed using the following
- 123 conditions: Runtime 260 hours; Switch time 1200 s to 4800 s; 1.5 V / cm; 0.75 x TBE at 8°C. Yeast
- 124 chromosomal DNA served as size marker (BioRad; 200 kb 2 Mb).
- 125 Southern blotting was conducted using standard protocols [19]. A digoxigenin labeled probe was 126 generated by PCR following the manufacturer's instructions (PCR DIG Probe Synthesis Kit,Roche). The 127 993 bp probe (Cd709 chr1, position 6,711,095 to 6,712,088) was specific to mini-chromosome-like 128 sequences at the right arm of chromosome 1 in Cd709. Hybridization was performed in DIG Easy Hyb 129 buffer at 42°C overnight. The membrane was then extensively washed with low and high stringency 130 buffers and subsequently blocked with buffer B2 (1% Blocking powder [Roche] in buffer B1 [100 mM 131 Maleic acid, 150 mM NaCl, pH 7.5]). The blocking solution was then replaced with antibody solution 132 (buffer B2 containing DIG-antibody 1:26,000 (Roche)). The membrane was washed with buffer B1 133 containing 0.3% Tween20. The membrane was subsequently equilibrated in buffer B3 (100 mM Tris
- 134 pH 9.5, 100 mM NaCl, 50 mM MgCl<sub>2</sub>) and developed with chemiluminescence (CDP-Star, Roche).

#### 135 Genome data, assembly, rearrangements and duplications

136 The genomic DNA of Cd709 was used to prepare a size-selected library (20kb) prior to sequencing with 137 a PacBio Sequel sequencer (kit 2.1, Keygene N.V., Wageningen, The Netherlands) on two SMRT cells, 138 yielding raw data with approximately 224 X genome coverage (1.474.759 reads, N50 10.837 bp). 139 Genome assemblies were generated from several runs of the Hierarchical Genome-Assembly Process 140 version 4 (HGAP4) and Canu [20] assemblers. The draft genome was polished with the Arrow algorithm 141 and the completeness of the assembly was evaluated with BUSCO using the Ascomycota gene set as 142 evidence [21]. Telomeres were validated by the presence of at least three repeats of the 143 TTAGGG/CCCTAA motif at the end of assembled contigs [22]. The polished assembly was aligned with 144 nucmer against the Ch63 and Ch35-RFP genomes to visualize chromosome rearrangements. 145 SDDetector [9] was used to detect segmental duplications in combination with Bedtools and BWA-146 MEM for validation. The Cd709 mitochondrial genome was assembled with Organelle\_PBA [23] (Table 147 S2).

#### 148 Transcriptome data and analysis

149 RNA sequencing was performed on samples of mRNA from undifferentiated mycelium grown 150 axenically and two different stages of plant infection, 48 and 72 h after inoculation, corresponding to 151 the biotrophic and necrotrophic phase, respectively. Mycelium was grown for three days in potato 152 dextrose liquid medium (PDB, Difco) at 25°C with shaking (150 rpm) and harvested by filtration. Seedlings of *M. sativa* (eight days old) were inoculated by placing a droplet (10 µl) of *Cd*709 spore 153 154 suspension (7 x  $10^5$  spores/ml) onto the surface of each cotyledon and the plants were then incubated as described for *M. truncatula*. Discs of infected cotyledon tissue were harvested using a cork borer (4 155 156 mm diameter). After grinding the tissues in liquid nitrogen, total RNA was extracted using the RNeasy 157 plant mini kit (Qiagen). Libraries were then prepared from each sample type using the TruSeq Paired-158 end Stranded mRNA Kit and sequenced (100 bp reads) using a HiSeq4000 sequencing platform 159 (IntegraGen Genomics, Evry, France). RNA-Seq paired reads were cleaned and trimmed using Trimmomatic [24] and then mapped to the genome assembly of Cd709 using STAR [25]. A genome-160 161 guided transcript assembly was obtained from mappings with StingTie v1.3.4. Assembled raw 162 transcripts were then filtered based on the TPM distribution per transcript per library.

### 163 Genome annotation

164 Transposable elements (TE) were searched in the *C. destructivum* genome sequence using the REPET

- package [26, 27]. Consensus sequences identified with the TEdenovo pipeline were classified using the
- 166 PASTEC tool [28], based on the Wicker hierarchical TE classification system [29], and then manually
- 167 filtered and corrected. The resulting library of consensus sequences was used to annotate TE copies in
- 168 the whole genome using the TEannot pipeline.
- Protein-coding genes were annotated using the Eugene [30] and FunGAP [31] tools. Predicted genes
   were filtered out when 10% of their CDS overlapped a Transposable Element predicted by the REPET
- package. Filtered predicted genes from Eugene and FunGAP were clustered together based on their
- 172 CDS coordinates (overlap of one base required) with no strand consideration. The Annotation Edit
- 173 Distance (AED) [32] was computed with transcript and protein evidence for each transcript and the
- predicted model with the best score was retained at each locus. Mitochondrial genomes were
- annotated with MFannot [33] and MITOS2 [34]. Results were manually inspected and in case of
- 176 divergence between the predictions, the longer gene model was retained.
- The synteny between *C. destructivum* and *C. higginsianum* proteomes was analysed with SynChro [35] which detects ortholog proteins with Reciprocal Best Hit (RBH), based on 40% similarity and a length ratio of 1.3. Colinear orthologs were then grouped in syntenic blocks, according to a delta threshold = 1 (very stringent mode). Non-syntenic blocks were extracted when five or more consecutive nonsyntenic genes were found. Proteome similarities with other *Colletotrichum* spp. were performed with Blast 2.2.28+ and the results filtered with a cut-off of 30% identity and 50% query coverage. Proteome
- 183 synteny and associated figures were obtained using Clinker [36].

## 184 Functional annotation of predicted genes

- Functional annotations of genes obtained using Interproscan 5.0 [37] and Blastp (e-value <1e-5) [38]</li>
   against the NCBI nr databank (September 2019) were then used to perform Gene Ontology [39]
   annotation with Blast2GO [40]. Carbohydrate active enzymes (CAZymes) were annotated with dbCAN2
- 188 [41] launching HMMER, Diamond and Hotpep against dedicated databases. Genes were considered as
- 189 CAZymes when at least 2 of the three tools provided a positive annotation.
- Genes encoding potential secreted proteins were predicted with a combination of SignalP v4.1 [42],
   TargetP v1.1 [43] and TMHMM v2.0 [44] results. The secretome was defined as the union of SignalP
   and TargetP provide and then interpreted with TMUMM results (0 or only 1 transmembrane domain)

Proteins smaller than 300 amino acids were then extracted and considered as Small Secreted Proteins (SPPs). In parallel, EffectorP v2.0 [45] was applied to the predicted secretome to identify putative effector proteins. Finally, the intersection of EffectorP and SPPs results was retained to establish a list of potential effectors.

197 To detect secondary metabolism biosynthetic gene clusters (BGCs), predicted genes were submitted 198 to antiSMASH (Antibiotics and Secondary Metabolite Analysis Shell) v5 [46]. Only core biosynthetic 199 genes (commonly known as secondary metabolism key genes, SMKGs) were considered for further 200 analysis. Presence/absence patterns of SMKGs were based on reciprocal best hits with Ch63 and Ch35, 201 and then manually inspected. Among the newly predicted secondary metabolism key genes (SMKGs), 202 those encoding polyketide synthases (PKS) and non-ribosomal peptide synthases (NRPS) were checked 203 for the presence of the minimal expected set of enzymatic domains, namely KS and AT domains for 204 PKS, and A and PCP domains for NRPS. Terpene synthases and dimethylallyltryptophan synthase 205 (DMATS) genes were manually inspected and retained if they had RNA-seq or protein support. Those 206 Cd709 genes not predicted as SMKGs by antiSMASH, but orthologous to a C. higginsianum SMKG were 207 also included. For example, antiSMASH failed to annotate six terpene synthase (TS) that are present in 208 both species.

### 209 Codon usage analysis

- 210 Codon usage was computed for predicted gene coding sequences (CDS) on each chromosome or region 211 using the EMBOSS tool 'cusp'. The resulting codon usage matrix (i.e. the fraction of each codon in a 212 given amino acid) was subjected to Fisher's exact tests (with a Bonferroni correction for multiple 213 testing) to address the statistical significance of differences between the core and mini-chromosomes. 214 The matrix was also subjected to a Principal Component Analysis (PCA) and the results were projected 215 onto the first two principal components. To analyse the GC percentage of the three letters of each 216 codon, the 'cusp' tool was run individually on each CDS of each chromosome or region and the results 217 were represented as density plots. The corresponding figures were generated using R (v. 4.0.5) and
- the libraries ggplot2 (v. 3.3.3), cowplot (v.1.1.1) and ggbeeswarm (v. 0.6.0), all available from the CRAN
- 219 repository (https://cran.r-project.org/).

## 220 <u>Results</u>

## 221 A novel Colletotrichum destructivum - Medicago truncatula pathosystem

222 The cell biology of infection of M. sativa by C. destructivum isolate 709 (Cd709) was previously 223 described [2]. Here, we report infection of the model plant *M. truncatula* (barrel medic) by this species. 224 Five out of the nine tested M. truncatula accessions, including the genome-sequenced accession 225 ESP074-A [47], were found to be susceptible to C. destructivum in two independent infection assays 226 (Fig. 1, Table S1). At 4 days post inoculation (dpi), necrotic water-soaked lesions were visible on the 227 trifoliate leaves of the susceptible accessions (Fig. 1). In contrast, the leaves of resistant accessions 228 presented only small necrotic flecks or no visible symptoms. The genome-sequenced accession R108-229 C3, which is widely used for *M. truncatula* functional genomics [48], was resistant to *C. destructivum* 230 in these infection assays.

On cotyledons of the susceptible *M. truncatula* accession ESP155-D, *Cd*709 spores germinated to form
 melanized appressoria, which by 48 hpi had penetrated host epidermal cells to form bulbous,

233 intracellular biotrophic hyphae that were confined to the first infected cell (Fig. 2a). Thinner 234 necrotrophic hyphae started to emerge from the tips of the biotrophic hyphae at 60 hpi (Fig. 2b), and 235 after 72 hpi the fungus had completed its asexual cycle by producing sporulating structures (acervuli) 236 on the surface of the dead tissues (Fig. 2c). On cotyledons of the resistant accession ESP163-E, 237 appressoria formed abundantly on the leaf surface but penetrated host epidermal cells very 238 infrequently (Fig. 2d, e). Groups of dead epidermal cells underlying the appressoria appeared yellow-239 brown in colour and had granulated contents, suggesting they had undergone a hypersensitive cell 240 death response. Rarely, small hyphae were visible in epidermal cells beneath appressoria but they

- 241 developed only a short distance into the dead cells and most remained smaller than the appressorium.
- 242 Acervuli were never observed on plants of accession ESP163-E.

#### 243 Genome assembly and structural annotation

244 Long-read data allowed us to generate a complete genome assembly for Cd709, with a total length of 245 51.75 Mb in which all 12 chromosomes were sequenced from telomere to telomere (Fig. 3), together 246 with the circular mitochondrial genome (34 kb). Annotation of transposable elements revealed a total 247 of 49 consensus sequences, representing all the possible TEs in the Cd709 genome. Classification of 248 the TEs (Table S3) showed that the genome contains 18 different families of retrotransposons, 249 including eleven LTR (Long Terminal Repeats) and seven LINE (long interspersed nuclear element), 28 250 DNA transposons, including 25 TIR (terminal inverted repeat), one helitron and two MITE (Miniature 251 Inverted-Repeat Transposable Elements), as well as three unclassified repeated elements. The library 252 of 49 consensus sequences was then used to annotate TE copies in the Cd709 genome. Overall, TEs 253 covered 6.2 % of the genome assembly by length. The Class I LTR Gypsy superfamily was the most 254 abundant in terms of coverage and number of copies, whereas the Class I TIR Tc1-Mariner was the 255 most abundant in terms of full-length copies. Two Gypsy transposons (R172 and G87) resemble the 256 most abundant TE family in C. higginsianum, namely the LTR transposon family RLX\_R119 [9]. Looking 257 at the distribution of TE families along the chromosomes, we found that the telomeres of all twelve C. 258 destructivum chromosomes were associated with a single copy of a TE belonging to the helitron family 259 (G103).

260 To annotate the protein-coding genes, a genome-guided assembly of RNA-Seq reads provided 16,122, 261 13,901 and 15,081 transcripts for axenic mycelium, 48 hpi and 72 hpi libraries, respectively (Table S4), 262 with 1.88 TPM, 9.38 TPM and 4.90 TPM as minimum expression levels, respectively (Fig. S1). Assembled transcripts were then used to predict gene models in conjunction with Colletotrichum and 263 264 Ascomycota protein databanks. The results of EuGene and FunGap were combined and filtered to 265 generate the Cd709 gene set comprising 15,631 complete gene models, of which 11,853 had transcript 266 support and 15,172 resembled Ascomycota predicted proteins. Features of the gene annotation are 267 summarized in Table S5. The completeness of this annotation was confirmed by comparison to the 268 BUSCO Ascomycota set (1,315 genes), with 1,309 complete genes predicted and only one missing. 269 Functional annotation assigned InterPro entries to 10,298 genes, among which 7,475 had at least one 270 GO term and 1,105 were potential enzymes (annotated with an Enzyme Code). Based on Blast2GO 271 descriptions, 12,192 predicted genes (78%) had a predicted function, i.e. a description other than 272 "hypothetical protein" (Table S6 tab 'All'). The mitochondrial genome of Cd709 was annotated with 29 273 tRNAs, 2 rRNAs (small and long subunit) and 21 genes.

#### 274 Plant interaction-related genes

275 A total of 619 Cd709 genes were annotated to encode CAZymes, among which 410 were assigned to the Glycoside Hydrolase (GH), Carbohydrate Esterase (CE) and Polysaccharide Lyase (PL) CAZyme 276 277 classes (Table S6 tab 'CAZyme'). The proportion of genes in each CAZyme class closely resembled that 278 previously found in Ch63 [49], and 98% (400/410) of Cd709 CAZyme genes were also detected in the 279 Ch63 genome. In silico analysis of the Cd709 secretome revealed a total of 2,608 potential extracellular 280 secreted proteins, including 1,118 small proteins (<300 amino acids). Among these, 484 genes were 281 retained as putative effectors because they were also present among 508 genes identified by EffectorP. 282 Comparing these to the effector repertoire of Ch63, a total of 127 putative effectors (26.2%) were 283 unique to Cd709, having no Reciprocal Best Blast Hit in Ch63 (Table S6 tab 'Predicted effectors'). A total 284 of 110 secondary metabolism key genes (SMKGs) were detected in the Cd709 genome using the fungal 285 version of antiSMASH and were manually curated. These C. destructivum SMKGs were compared to 286 the 105 C. higginsianum SMKGs [9]. Overall, 78 % (94 out of 120) of the SMKGs were present in both 287 species (Table S6 tab 'Secondary metabolism', Fig. S2). A total of 17 C. destructivum SMKGs, distributed 288 over eight BGCs, were not detected in C. higginsianum.

#### 289 Chromosome structure comparison

290 Complete chromosome-level assemblies are available for two different C. higginsianum strains, 291 namely IMI 349063A (Ch63) [9] and MAFF 305635-RFP (Ch35-RFP), a transformant of MAFF 305635 292 (Ch35) expressing red fluorescent protein which lacks both mini-chromosomes 11 and 12 [10, 17]. The 293 genetic proximity of C. destructivum and C. higginsianum allowed us to align assemblies to observe 294 chromosome structural variations. This generated 38 Mb of C. destructivum alignments (>10 kb) with 295 each C. higginsianum strain, ranging from 88 to 96.7% identity. Thus, C. destructivum shared 296 approximately 73.6 % of its total genome length with C. higginsianum. At the chromosome scale, 297 alignments revealed that five chromosomes of C. destructivum (chr1, 2, 3, 5 and 9) were not involved 298 in any large rearrangements, five others (chr4, 6, 7, 8 and 10) showed inter-chromosomal 299 rearrangements, while the two mini-chromosomes (chr11 and 12) lacked large regions of conserved 300 sequences and appear to be species specific (Fig. 4A).

301 One rearrangement involved chr7 and chr8 of Cd709 resulting in chr4 and chr10 of Ch63. The break-302 points in chr7 and chr8 were associated with TEs in Cd709 (Fig. S3 A and B). A similar rearrangement 303 was found relative to Ch35-RFP, albeit with different break-points in both species that were not 304 associated with TEs (Fig. S3 F and G). A second rearrangement involved chr4 and chr10 of Cd709 such 305 that their left and right arms result in chr9 and chr7 of Ch63, respectively (Fig. S3 C and D). Interestingly 306 this rearrangement was not found relative to Ch35-RFP, suggesting that it is specific to particular 307 C. higginsianum strains, as was noted previously [10]. A third inter-chromosomal rearrangement 308 concerned 121 kb at the 5' extremity of Cd709 chr6 coming from chr4 and contig\_1 of Ch63 and Ch35-309 RFP, respectively. In C. destructivum, this break-point is surrounded by TEs and non-syntenic regions 310 (Fig. S3 E). Remarkably, a specific rearrangement of 42 kb between chr11 of Cd709 and contig 11 of 311 Ch35-RFP (Fig. S3 H) corresponds to a region that is absent from the Ch63 genome assembly and which 312 encodes highly variable effectors (having  $\leq$  90% alignment coverage) and secondary metabolism-313 related proteins [10]. In addition, several short stretches (2 to 5 kb in length) from chr11 of Cd709 were 314 present at the extremities of chromosome 6 in Ch63 and the corresponding region of Ch35-RFP (contig 315 \_9) (Fig. 4).

316 A notable feature of the *C. destructivum* genome assembly is the unusually large size of chr1 (7.3 Mb), 317 which is 0.9 Mb longer than the largest chromosome in C. higginsianum (6.4 Mb). Genome alignments 318 highlighted a near-complete synteny between chr1 of Cd709 and chr2 of Ch63 except for a 1.2 Mb 319 subtelomeric region (coordinates chr1:6076875-7282542), for which no similarity was found in C. 320 higginsianum (Fig. 4). Synteny between the genes of Cd709 and those of Ch63 was investigated using 321 SynChro. With stringent settings, 400 syntenic blocks were identified based on 12,135 Reciprocal Best 322 Hits. A total of 1,083 genes were found in 47 non-syntenic blocks composed of at least five consecutive 323 Cd709-specific genes (Tables S7 & S8). The largest non-syntenic block, corresponding to the 1.2 Mb 324 region specific to Cd709 on chr1, contained 305 genes. Mini-chromosome chr12 contained one non-325 syntenic block of 170 genes, while chr11 was divided into seven non-syntenic blocks, the largest 326 containing 106 genes. Although only 356/1,083 genes inside non-syntenic blocks could be annotated 327 with a GO term, GO enrichment tests revealed that the Cd709-specific genes were enriched in protein 328 kinases, protein phosphorylation activity and secondary metabolism process (Table S9). Likewise, 329 effector genes were found to be enriched in non-syntenic blocks whereas CAZymes were depleted

330 (Table S10).

### 331 Validation of the 1.2 Mb non-syntenic region in *C. destructivum* chromosome 1

To verify the large non-syntenic region identified within chr1, we first checked for potential errors in the sequence assembly of this region by manually inspecting long reads spanning the two junctions (Fig. S4). Secondly, to obtain an assembly-independent validation, pulsed-field gel electrophoresis (PFGE) and a Southern hybridization were performed (Fig. 5A, B). A 993 nt probe (coordinates chr1: 6,711,095 to 6,712,088) was designed within the 1Mb non-syntenic region to target a unique locus that avoided TEs (Fig. 5B). This probe is 83.5% identical to the gene CH63R\_14488 located on chromosome 11 of *Ch*63 that was used as a hybridization control.

339 Chromosomes of two C. destructivum isolates (Cd709 and Cd202) and two C. higginsianum isolates (Ch63 and Ch35) were separated by PFGE and analysed by Southern hybridization (Fig. 5C, D). For both 340 341 C. destructivum isolates, the probe hybridized to molecules with high molecular weight that could correspond to the largest chromosome, consistent with a location on chr1 (Fig. 5C, D). The high 342 343 molecular weight signals were absent in the C. higginsianum blots, and instead hybridization signals 344 were detected at a position corresponding to mini-chromosome 11, although these were weak, as 345 expected for a probe with only 83.5% identity to the target. Overall, our findings validate that a non-346 syntenic region is embedded within chr1 of C. destructivum. Hereafter, we refer to the syntenic and 347 non-syntenic portions as chr1A and chr1B, respectively, and their distinct properties were explored 348 further in the following analyses.

### 349 Region chr1B shows the characteristic features of fungal accessory chromosomes

In many aspects, the region chr1B of Cd709 resembled the mini-chromosomes 11 and 12. All three 350 351 compartments were more AT-rich than the core genome. Region chr1B was also highly enriched with 352 TEs, having 32.8 % coverage with TE copies by length, similar to chr11 and chr12 (32.3 and 35.1 %, 353 respectively), whereas the core chromosomes (excluding chr1B) had only 3 to 6.2 % TE coverage (Table 354 1, Fig. 3, Table S11). Moreover, the distribution of TE families in region 1B and the two mini-355 chromosomes differed markedly from the core chromosomes in that they were all enriched with LINE 356 retrotransposons (44 %, 19 % and 34 % coverage, respectively), compared to only 7 % in the core 357 genome (Table 3). LINE TEs are also present in C. higginsianum on mini-chromosomes 11 and 12, but their expansion was less striking in this species (7 % and 2 % coverage, respectively) (Fig. S5) than in *Cd*709 [9].

360 Examination of the gene content of region chr1B revealed that, similar to the mini-chromosomes, it 361 was overall depleted in protein-coding genes (2-fold less than the core chromosomes), contained a 362 significantly larger proportion of genes encoding proteins of unknown function (i.e. annotated as hypothetical proteins), and had fewer expressed genes (RNA-seq transcript evidence) compared to the 363 364 core genome (Table 1). Considering categories of potentially pathogenicity-related genes, no CAZyme 365 genes or SMKGs were detected in either region chr1B or chr 12, although eight SMKGs were present 366 on chr11 (Table S6, Tab 'Secondary metabolism'), all of which had RNA-seq transcript support. 367 Moreover, 38 effectors were found in chr1B and the two mini-chromosomes. Remarkably, 36 of these 368 were absent from C. higginsianum (had no RBH in Ch63), of which 20 were expressed in planta (Table S6 tab 'Predicted effectors). With 15 and 10 effectors respectively, the mini-chromosomes 11 and 12 369 370 were significantly enriched in putative effectors compared to the core chromosomes whereas no 371 enrichment was observed for the 13 effectors of the chr1B (Table 1). Remarkably, the most highly 372 expressed effectors during the biotrophic phase (48 hpi), namely CDEST\_01870 (chr1B) and 373 CDEST\_15472 (chr12), were located on mini-chromosome-like regions. This raises the possibility that 374 genes carried in such regions are important for virulence.

#### 375 Codon usage in region chr1B and the mini-chromosomes differs from the core chromosomes

376 Analyses of codon usage were used previously to detect differences between the core and accessory 377 chromosomes or lineage-specific compartments of other plant pathogenic fungi [15, 50, 51]. We 378 therefore computed the codon usage of CDS located on the core chromosomes, mini-chromosomes 379 and the chr1B region of Cd709. Based on a principal component analysis, codon usage on the core 380 chromosomes was very homogeneous, whereas that of the mini-chromosomes and region chr1B 381 clustered together and separately from the core chromosomes (Fig. 6A). To illustrate this in greater detail, we plotted the codon usage for each amino acid and for each chromosome or region (Fig. S6, 382 383 representative examples are given for 3 amino acids in Fig. 6B). For these analyses, we excluded the two amino acids (Trp and Met) that are encoded by a single codon. Based on Fisher's exact tests for 384 385 each of the remaining 59 codons, almost all the codon usages were different between the core 386 chromosomes on one hand and chr1B, chr11 or chr12 on the other hand. In striking contrast, there 387 were only three differential codon usages between chr1B and chr11 and one between chr1B and chr12. 388 However, chr11 and chr12 were most different from each other with 15 differential codons (Table S12; 389 adjusted *P* < 0.001).

#### 390 **Region chr1B is a hotspot for segmental duplications**

The genome of *Cd*709 was inspected for segmental duplications, as described previously for *C. higginsianum* [9]. A total of 48 duplications involving genes were detected on four chromosomes (chr1, chr6, chr11 and chr12). Among them, 12 duplications were larger than 10 kb (Fig. 7) of which only three were inter-chromosomal (all involving chr12). Similar to *C. higginsianum* [9], these interchromosomal duplications were all associated on at least one side with TEs, supporting a potential role of TEs in duplication (Fig. S7). However, in contrast to *C. higginsianum*, these duplications did not take place preferentially near telomeres. 398 A remarkable feature of region chr1B was that it showed a strong intra-chromosome duplication 399 pattern, with some regions replicated up to three times (Fig. 7). Assembling large duplications can be 400 difficult even with long-read sequences [52]. To check for possible bias during assembly, the eight 401 largest intra-chromosome duplications on chr1B were inspected manually (Table S13). Due to the 402 problem of multiple reads mapping to duplicated regions, we considered only uniquely mapped reads. 403 Consequently, the read-coverage of these eight regions was on average 2-fold lower than the non-404 duplicated regions. No other regions of chr1B showed a significant decrease in coverage, and the 405 extremities of the SD regions were well-anchored to chr1B. Reads were identified spanning the two 406 smallest duplications, SD1B-2 (10 reads) and SD1B-6 (22 reads), but other duplicated regions were too 407 large (>16 kb) to be spanned by single PacBio reads. Finally, the short-read RNA-seq data used to 408 annotate the genome were also employed to detect mutations within the duplicated genes. Mutations 409 were detected in all the duplicated regions, albeit with support from only few reads in most cases. 410 Taken together, these results support the reliability of the observed duplications in region chr1B.

411 To gain insight into the possible origin of region chr1B, we examined conservation of the 300 genes 412 contained within this region in the genomes of 23 other Colletotrichum species (Table S14). As 413 expected, given that C. destructivum and C. higginsianum belong to the same species complex [3], the 414 total proteome of Cd709 showed greatest similarity to that of Ch63 (14,372 conserved proteins). 415 Surprisingly however, the chr1B proteome shared most conserved proteins with a phylogenetically 416 distant species, namely C. truncatum (217 protein matches, compared to only 134 matches in C. 417 higginsianum) [53]. Almost half of the genes shared with C. truncatum were involved in segmental 418 duplications within the Cd709 chr1B. Remarkably, the region triplicated in SD1B-1, SD1B-3 and SD1B-419 7 was also found in a large duplicated region represented by two contigs within the C. truncatum 420 genome assembly (Fig. S8) [54], which may be located on a mini-chromosome due to their low GC 421 content (49.0%, compared to 51.2% in the longer contigs of C. truncatum). Other genes located within 422 the Cd709 SD1B-1 duplications had Blast matches that were mostly restricted to C. incanum, C. 423 spaethianum and C. tofieldiae (Spaethianum species complex), C. salicis and C. nymphaeae (Acutatum 424 species complex), C. fructicola (Gloeosporioides species complex), C. sublineola (Graminicola species 425 complex) and C. orchidophilum, which vary in their phylogenetic distance from C. destructivum [53]. 426 The absence of these gene sequences from the C. higginsianum genome was confirmed by Tblastn 427 searches against the NCBI wgs Colletotrichum database (266 genomes).

Examination of the gene content in duplicated regions of chr1B gave few clues to their possible role in the host interaction or the advantage for the fungus to maintain multiple mutated copies of these genes. One gene duplicated four times (CDEST\_01898, CDEST\_01949, CDEST\_02058 and CDEST\_02116) encoded a major facilitator superfamily transporter. The five genes duplicated between SD1B-2 and SD1B-6 comprised four FAD-binding domain-containing proteins and a patatin-like serine hydrolase.

## 434 Discussion

In this study, we present a chromosome-level reference assembly of the *C. destructivum* genome, a phytopathogen causing anthracnose disease principally on species of *Medicago* and *Trifolium* (Fabaceae). Among other members of the Destructivum species complex, which currently contains 17 recognised species [3], the genomes of *C. lentis, C. tanaceti* and *C. shisoi* were sequenced previously but the resulting assemblies were highly fragmented, containing 2980, 5242 and 36,350 contigs, respectively [8, 11, 12]. Using PacBio long-read sequencing, we were able to generate a gapless assembly of the *Cd*709 genome which, together with that of *Ch*63 [9], provides a second complete genome within the Destructivum species complex, facilitating future comparative genomic analyseswithin this important group of plant pathogens.

444 Alignment of the Cd709 genome assembly with those of C. higginsianum strains Ch63 and Ch35 445 revealed large-scale chromosome rearrangements between the two closely-related species. Some of 446 these rearrangements were potentially mediated by recombination between homologous regions 447 containing TEs, which flanked one or both of the breakpoints. Similar TE-mediated chromosome 448 rearrangements were previously reported at the intra-species level in C. higginsianum [10]. Our 449 analysis of synteny between the genomes of Cd709 and Ch63 also revealed the presence of a 1.2 Mb 450 species-specific region within Chr1 of Cd709, which we called Chr1B. This 'accessory region' (AR) 451 displays many of the hallmarks that characterize fungal mini-chromosomes, or 'accessory 452 chromosomes', in that it is AT-rich, transposon-rich, gene-poor and has a distinct codon usage [51, 55-453 57]. In all these respects, Chr1B resembles the mini-chromosomes Chr11 and Chr12 but is strikingly 454 different from the rest of Chr1 and other core chromosomes of Cd709. The TE enrichment observed in 455 Chr1B and both mini-chromosomes is largely caused by the specific expansion of LINE and TIR elements 456 in these compartments, unlike the core chromosomes where the Gipsy TE family predominates.

457 Using PFGE and Southern hybridization with a probe specific to Chr1B, we were able to confirm that 458 this AR is carried not only on Chr1 of Cd709 but also on the largest chromosome of Cd202, despite the 459 widely-separated geographical origins of these two isolates (Saudi Arabia and Morocco, respectively). 460 Analysis of a larger collection of C. destructivum isolates is now needed to determine the extent to 461 which Chr1B is conserved within this pathogen species. The presence of an AR embedded within a core 462 chromosome has been reported in other plant pathogenic fungi. For example, isolates of the T race of 463 Cochliobolus heterostrophus harbor an AR of about 1.2 Mb distributed between two core 464 chromosomes that contains the Tox1 locus producing the T-toxin polyketide [58, 59]. In Verticillium dahliae, Chr3 and Chr4 each harbor two ARs of ~300 kb [60], while in Fusarium poae a 204 kb block 465 466 with AR characteristics is inserted near one telomere of Chr3 [57]. However, it should be noted that in 467 these two examples the inserted AR blocks are 4- to 6-fold smaller than Chr1B of Cd709.

468 Our working hypothesis is that the AR Chr1B arose by the integration of a mini-chromosome into a 469 core chromosome of *C. destructivum*, but the mechanism by which this occurred is unclear. Despite 470 the subtelomeric position of Chr1B, its integration is unlikely to have resulted from the telomeric fusion 471 of a mini-chromosome with a core chromosome because it is flanked on both sides by portions of Chr1, 472 both of which are highly syntenic to Chr2 of C. higginsianum. A chromosome containing distinct regions 473 characteristic of core and accessory chromosomes was previously reported in the genome of C. 474 fructicola strain Nara gc5 [61]. In this case, the chimeric chromosome, called Nara\_c11, is smaller (2.8 475 Mb) than Cd709 chromosome 1 (7.3 Mb) and the TE-rich, gene-poor AR occupies most of the 476 chromosome (66 % by length), in contrast to Cd709 Chr1B, which occupies only 16 %. A further 477 difference to Cd709 Chr1B is that the AR of Nara\_c11 includes a telomere, suggesting that in this case 478 the chimeric chromosome arose through a different mechanism. Taken together, our findings provide 479 further evidence for genetic exchange between core and accessory genomic compartments in 480 Colletotrichum species [61]. In other fungi, chromosome breakage-fusion-bridge (BFB) cycles have 481 been invoked not only in the creation of accessory chromosomes from core chromosomes [62], but 482 also in their reintegration into core chromosomes [63].

483 A distinguishing feature of the Chr1B AR is that it has undergone extensive region-specific segmental duplications. Some inter-chromosomal SDs in Cd709 were associated with TEs at one or both of their 484 485 borders, as we found previously in Ch63 [9], but there was little evidence that the region-specific SDs 486 in Chr1B were mediated by TEs. Similarly, the AR of *C. fructicola* chromosome Nara\_c11 was found to 487 be implicated in numerous intra- and inter-chromosomal SDs but as in Cd709 these were not 488 consistently flanked by TEs [61]. Among fungal pathogens, SDs can play important roles in generating 489 genetic diversity and novel gene functions, either at the level of expression or coding sequence [64, 490 65]. A recent study on Fusarium strains infecting banana also highlighted the importance of SDs in 491 driving the evolution of ARs and the effector genes contained within them [66]. Although the C. 492 destructivum genome contains a complete Mat1-2-1 mating-type locus (Table S6, Tab MAT1-2-1), and 493 should therefore be capable of sexual reproduction, this has never been observed [67], [3]. In this 494 context, segmental duplication may therefore provide an important mechanism for generating genetic 495 diversity for host adaptation in this essentially asexual pathogen.

496 A remarkable finding was that some segmentally duplicated blocks of genes within Chr1B of C. 497 destructivum are conserved and syntenic with duplicated regions in the genome of C. truncatum, a 498 species that is phylogenetically very distant [53]. Given that these two taxa diverged ~60 million years 499 ago [68], soon after speciation in Colletotrichum, these SDs may be very ancient and have been 500 selectively retained in some species and lost in others. Alternatively, these duplicated regions may 501 have been acquired by horizontal chromosome transfer (HCT) from another species to a common 502 ancestor, or through independent transfers to C. destructivum and C. truncatum. HCT would be 503 consistent with the distinct codon bias in Chr1B and the taxonomic incongruity of many genes within 504 this region. The horizontal transfer of a mini-chromosome between vegetatively incompatible biotypes 505 of C. gloeosporioides was shown experimentally [69, 70], and it is well-documented that genetic 506 material can be exchanged following fusion between conidial anastomosis tubes of the same, or even 507 different, Colletotrichum species [71-73].

508 Chr1B contains a variety of genes with potential roles in fungal virulence, some of which were 509 expressed during infection. These include genes encoding 13 candidate secreted effector proteins, 8 protein kinases, 5 major facilitator superfamily membrane transporters, 5 heterokaryon 510 511 incompatibility (HET) proteins and 8 putative transcription factors (TFs) (Table S6). It is interesting to 512 note that, similar to Chr1B, the accessory 'pathogenicity chromosome' of Fusarium oxysporum f.sp. 513 lycopersici is enriched not only with effectors genes but also with genes encoding protein kinases, 514 membrane transporters, HET proteins and TFs, of which one TF was shown to regulate the expression 515 of plant-induced effector genes [74],[75]. TFs were also found to be enriched in the four lineage-516 specific ARs of V. dahliae [60]. Overall, the gene content of Chr1B suggests that it may contribute to C. 517 destructivum pathogenicity. This was demonstrated experimentally for ARs in two other members of 518 the Destructivum species complex, namely Chr11 of C. higginsianum (isolate Ch35) which was essential 519 for virulence on *A. thaliana* [17], and Chr11 of *C. lentis*, which was required for virulence on lentil [12]. 520 In the case of Cd709, it is noteworthy that the three most highly expressed and plant-induced effector 521 genes are all located in ARs, namely CDEST\_01870 on Chr1B, CDEST\_15404 on Chr11 and CDEST\_15472 522 on Chr12. These and other pathogenicity-related genes carried within these genomic compartments 523 will provide interesting candidates for future functional analysis.

524 Finally, we show here that *Cd*709 can complete its life cycle not only on its original host, *M. sativa*, but 525 also on the widely-studied model legume, *M. truncatula*. Until now, the only other *Colletotrichum* 

- 526 species known to attack *M. truncatula* was *C. trifolii*, which belongs to the phylogenetically distant
- 527 Orbiculare species complex and uses a different infection process where the biotrophic phase extends
- to many host cells [76, 77]. With complete genome assemblies and high-quality gene annotations available for both partners, together with abundant genetic tools and resources on the plant side, the
- 530 *C. destructivum M. truncatula* interaction could provide a tractable new model pathosystem for
- 531 studying hemibiotrophic fungal interactions with Fabaceae hosts. Our identification of susceptible and
- 532 resistant *M. truncatula* accessions also raises the possibility that natural variation among accessions
- 533 could be exploited to analyse the genetic basis of resistance to *C. destructivum* [78].

# 534 Conflicts of interest:

535 The authors declare that there are no conflicts of interest.

# 536 <u>Funding:</u>

- 537 This work was partly supported by funding from the Agence Nationale de la Recherche (ERA-CAPS grant
- 538 ANR-17-CAPS-0004-01) to R.J.O. The BIOGER unit benefits from the support of Saclay Plant Sciences-
- 539 SPS (ANR-17-EUR-0007). The Funders had no role in the study design, data analysis, data interpretation
- 540 or decision to publish.

# 541 <u>Author contributions:</u>

- 542 Conceptualization: NL, AS, CK, JFD, RJO; Investigation: PLP, SP, AA, RJO; Formal analysis: NL, AS, AL, JA,
- 543 JFD; Visualization: NL, AS, PLP, JA, CK, JFD, RJO; Writing Original Draft: NL, AS, PLP, JA, CK, JFD, RJO;
- 544 Writing Review & Editing: NL, AS, CK, JFD, RJO; Supervision: NL, CK, RJO; Funding acquisition: RJO.

# 545 Acknowledgements:

546 We are grateful to the following bioinformatics platforms and partners for providing help and/or 547 computing and/or storage resources: Genotoul bioinformatics platform Toulouse Occitanie (Bioinfo 548 Genotoul, doi: 10.15454/1.5572369328961167E12), CATI BARIC (https://www.cesgo.org/catibaric/), 549 and INRAE-LIPME Bioinfo (Sébastien Carrère).We also thank Dr Alexander Wittenberg (KeyGene 550 N.V.,The Netherlands) for help with sequencing, and the INRAE Centre de Ressources Biologiques 551 *Medicago truncatula* for providing seeds.

# 552 <u>References</u>

- Frayssinet S. Colletotrichum destructivum: a new lucerne pathogen in Argentina. Australasian
   Plant Disease Notes 2008 3:1 2008;3:68–68.
- Latunde-Dada AO, Bailey JA, Lucas JA. Infection process of *Colletotrichum destructivum* O'Gara
   from lucerne (Medicago sativa L.). *European Journal of Plant Pathology* 1997;103:35–41.
- Damm U, O'Connell RJ, Groenewald JZ, Crous PW. The *Colletotrichum destructivum* species
   complex hemibiotrophic pathogens of forage and field crops. *Studies in Mycology* 2014;79:49–
   84.
- Sun HY, Liang Y. First report of anthracnose on sunflower caused by *Colletotrichum destructivum* in China. *Plant Disease* 2018;102:245.

- 562 5. Manandhar JB. Colletotrichum destructivum, the anamorph of Glomerella glycines.
  563 Phytopathology 1986;76:282.
- 564 6. **Tiffany LH, Gilman JC**. Species of *Colletotrichum* from Legumes. *Mycologia* 1954;46:52–75.
- Damm U, Sato T, Alizadeh A, Groenewald JZ, Crous P. The Collectotrichum dracaenophilum, C. magnum and C. orchidearum species complexes. Studies in mycology;92. 2019. DOI: 10.1016/J.SIMYCO.2018.04.001.
- Gan P, Tsushima A, Hiroyama R, Narusaka M, Takano Y, *et al. Colletotrichum shisoi* sp. nov., an
   anthracnose pathogen of *Perilla frutescens* in Japan: molecular phylogenetic, morphological and
   genomic evidence. *Scientific Reports*;9. 2019. DOI: 10.1038/s41598-019-50076-5.
- Dallery J-F, Lapalu N, Zampounis A, Pigné S, Luyten I, et al. Gapless genome assembly of
   *Colletotrichum higginsianum* reveals chromosome structure and association of transposable
   elements with secondary metabolite gene clusters. *BMC Genomics* 2017;18:667.
- Tsushima A, Gan P, Kumakura N, Narusaka M, Takano Y, et al. Genomic plasticity mediated by
   transposable elements in the plant pathogenic fungus Colletotrichum higginsianum. Genome
   biology and evolution 2019;11:1487–1500.
- Lelwala R V., Korhonen PK, Young ND, Scott JB, Ades PK, *et al.* Comparative genome analysis
   indicates high evolutionary potential of pathogenicity genes in *Collectotrichum tanaceti*. *PLOS ONE* 2019;14:e0212248.
- Bhadauria V, MacLachlan R, Pozniak C, Cohen-Skalie A, Li L, *et al.* Genetic map-guided genome
   assembly reveals a virulence-governing minichromosome in the lentil anthracnose pathogen
   *Collectorichum lentis. The New phytologist* 2019;221:431–445.
- 13. Wang Haoming, Huang Rong, Ren Jingyi, Tang Lihua, Huang Suiping, *et al.* The evolution of
   mini-chromosomes in the fungal genus *Colletotrichum. mBio* 2023;14:e00629-23.
- Flaumann P-L, Koch C. The many questions about mini chromosomes in *Colletotrichum spp*.
   *Plants*;9. 2020. DOI: 10.3390/plants9050641.
- 15. Ma LJ, Van Der Does HC, Borkovich KA, Coleman JJ, Daboussi MJ, *et al.* Comparative genomics
   reveals mobile pathogenicity chromosomes in *Fusarium*. *Nature 2010 464:7287* 2010;464:367–
   373.
- Liu S, Lin G, Ramachandran SR, Daza LC, Cruppe G, et al. Rapid mini-chromosome divergence
   among fungal isolates causing wheat blast outbreaks in Bangladesh and Zambia. New
   Phytologist. 2023. DOI: 10.1111/nph.19402.
- 17. Plaumann P-L, Schmidpeter J, Dahl M, Taher L, Koch C. A dispensable chromosome Is required
   for virulence in the hemibiotrophic plant pathogen *Colletotrichum higginsianum*. *Frontiers in microbiology* 2018;9:1005.
- 596 18. O'Connell R, Herbert C, Sreenivasaprasad S, Khatib M, Esquerré-Tugayé M-T, et al. A novel
   597 Arabidopsis-Colletotrichum pathosystem for the molecular dissection of plant-fungal
   598 interactions. Molecular plant-microbe interactions : MPMI 2004;17:272–82.
- 599 19. Stiehler F, Steinborn M, Scholz S, Dey D, Weber APM, *et al.* Helixer: cross-species gene
   annotation of large eukaryotic genomes using deep learning. *Bioinformatics* 2021;36:5291–5298.

- Koren S, Walenz BP, Berlin K, Miller JR, Bergman NH, et al. Canu: scalable and accurate long read assembly via adaptive k-mer weighting and repeat separation. *Genome research* 2017;27:722–736.
- Seppey M, Manni M, Zdobnov EM. BUSCO: Assessing genome assembly and annotation
   completeness. In: *Methods in Molecular Biology*. Humana Press Inc.; 2019. pp. 227–245.
- Rehmeyer C, Li W, Kusaba M, Kim Y-S, Brown D, et al. Organization of chromosome ends in the
   rice blast fungus, *Magnaporthe oryzae*. *Nucleic Acids Research* 2006;34:4685–4701.
- Soorni A, Haak D, Zaitlin D, Bombarely A. Organelle\_PBA, a pipeline for assembling chloroplast
   and mitochondrial genomes from PacBio DNA sequencing data. *BMC genomics* 2017;18:49.
- Bolger AM, Lohse M, Usadel B. Trimmomatic: a flexible trimmer for Illumina sequence data.
   *Bioinformatics (Oxford, England)* 2014;30:2114–20.
- Dobin A, Davis CA, Schlesinger F, Drenkow J, Zaleski C, *et al.* STAR: ultrafast universal RNA-seq
   aligner. *Bioinformatics* 2013;29:15–21.
- Flutre T, Duprat E, Feuillet C, Quesneville H. Considering transposable element diversification in
   *de novo* annotation approaches. *PloS one* 2011;6:e16526.
- Amselem J, Lebrun M-H, Quesneville H. Whole genome comparative analysis of transposable
   elements provides new insight into mechanisms of their inactivation in fungal genomes. *BMC genomics* 2015;16:141.
- Hoede C, Arnoux S, Moisset M, Chaumier T, Inizan O, *et al.* PASTEC: an automatic transposable
   element classification tool. *PLoS ONE* 2014;9:e91929.
- Wicker T, Sabot F, Hua-Van A, Bennetzen JL, Capy P, et al. A unified classification system for
   eukaryotic transposable elements. *Nature Reviews Genetics* 2007;8:973–982.
- Sallet E, Gouzy J, Schiex T. EuGene: An automated integrative gene finder for eukaryotes and
   prokaryotes. Humana, New York, NY; 2019. pp. 97–120.
- Min B, Grigoriev I V, Choi I-G. FunGAP: fungal genome annotation pipeline using evidence-based
   gene model evaluation. *Bioinformatics* 2017;33:2936–2937.
- 627 32. Eilbeck K, Moore B, Holt C, Yandell M. Quantitative measures for the management and
   628 comparison of annotated genomes. *BMC bioinformatics* 2009;10:67.
- 33. Valach M, Burger G, Gray MW, Lang BF. Widespread occurrence of organelle genome-encoded
   5S rRNAs including permuted molecules. *Nucleic acids research* 2014;42:13764–77.
- Bernt M, Donath A, Jühling F, Externbrink F, Florentz C, et al. MITOS: improved de novo
   metazoan mitochondrial genome annotation. *Molecular Phylogenetics and Evolution* 2013;69:313–319.
- Brillon G, Carbone A, Fischer G. SynChro: a fast and easy tool to reconstruct and visualize
   synteny blocks along eukaryotic chromosomes. *PLoS ONE* 2014;9:e92621.
- 636 36. Gilchrist CLM, Chooi YH. clinker & clustermap.js: automatic generation of gene cluster
   637 comparison figures. *Bioinformatics (Oxford, England)* 2021;37:2473–2475.

- 37. Jones P, Binns D, Chang H-Y, Fraser M, Li W, *et al.* InterProScan 5: genome-scale protein
   function classification. *Bioinformatics* 2014;30:1236.
- 640 38. Camacho C, Coulouris G, Avagyan V, Ma N, Papadopoulos J, *et al.* BLAST+: architecture and
   641 applications. *BMC Bioinformatics* 2009;10:421.
- 642 39. Gene Ontology Consortium. The Gene Ontology (GO) database and informatics resource. *Nucleic* 643 *Acids Research* 2004;32:258D 261.
- 644 40. Götz S, García-Gómez JM, Terol J, Williams TD, Nagaraj SH, *et al.* High-throughput functional
   645 annotation and data mining with the Blast2GO suite. *Nucleic Acids Research* 2008;36:3420.
- 41. Zhang H, Yohe T, Huang L, Entwistle S, Wu P, *et al.* dbCAN2: a meta server for automated
   carbohydrate-active enzyme annotation. *Nucleic Acids Research* 2018;46:W95–W101.
- 42. Petersen TN, Brunak S, von Heijne G, Nielsen H. SignalP 4.0: discriminating signal peptides from
   transmembrane regions. *Nature Methods* 2011;8:785–786.
- 43. Emanuelsson O, Nielsen H, Brunak S, von Heijne G. Predicting subcellular localization of proteins
   based on their N-terminal amino acid sequence. *Journal of Molecular Biology* 2000;300:1005–
   1016.
- 44. Krogh A, Larsson B, von Heijne G, Sonnhammer ELL. Predicting transmembrane protein
  topology with a hidden markov model: application to complete genomes. *Journal of Molecular Biology* 2001;305:567–580.
- 45. Sperschneider J, Dodds PN, Gardiner DM, Singh KB, Taylor JM. Improved prediction of fungal
  effector proteins from secretomes with Effector P 2.0. *Molecular plant pathology* 2018;19:2094–
  2110.
- 46. Blin K, Shaw S, Steinke K, Villebro R, Ziemert N, *et al.* antiSMASH 5.0: updates to the secondary
   metabolite genome mining pipeline. *Nucleic Acids Research* 2019;47:W81–W87.
- 47. Zhou P, Silverstein KAT, Ramaraj T, Guhlin J, Denny R, *et al.* Exploring structural variation and
  gene family architecture with *de novo* assemblies of 15 Medicago genomes. *BMC Genomics*2017;18:261.
- 48. Moll KM, Zhou P, Ramaraj T, Fajardo D, Devitt NP, *et al.* Strategies for optimizing BioNano and
   Dovetail explored through a second reference quality assembly for the legume model, *Medicago truncatula. BMC Genomics* 2017;18:578.
- 667 49. O'Connell RJ, Thon MR, Hacquard S, Amyotte SG, Kleemann J, *et al.* Lifestyle transitions in plant
   668 pathogenic *Colletotrichum* fungi deciphered by genome and transcriptome analyses. *Nature* 669 *Genetics* 2012;44:1060–1065.
- 50. Coleman JJ, Rounsley SD, Rodriguez-Carres M, Kuo A, Wasmann CC, *et al.* The genome of
   *Nectria haematococca*: contribution of supernumerary chromosomes to gene expansion. *PLOS Genetics* 2009;5:e1000618.
- 51. Hu J, Chen C, Peever T, Dang H, Lawrence C, et al. Genomic characterization of the conditionally
  dispensable chromosome in *Alternaria arborescens* provides evidence for horizontal gene
  transfer. *BMC genomics* 2012;13:1–13.

- 52. Vollger MR, Dishuck PC, Sorensen M, Welch AE, Dang V, et al. Long-read sequence and
   assembly of segmental duplications. *Nature Methods* 2019;16:88–94.
- 53. Liu F, Ma ZY, Hou LW, Diao YZ, Wu WP, *et al.* Updating species diversity of *Colletotrichum*, with
   a phylogenomic overview. *Studies in Mycology*;101.
- 680 54. Rogério F, Boufleur TR, Ciampi-Guillardi M, Sukno SA, Thon MR, *et al.* Genome sequence
   681 resources of *Colletotrichum truncatum*, *C. plurivorum*, *C. musicola*, and *C. sojae*: four species
   682 pathogenic to soybean (*Glycine max*). *Phytopathology* 2020;110:1497–1499.
- 55. Langner T, Harant A, Gomez-Luciano LB, Shrestha RK, Win J, *et al.* Genomic rearrangements
   generate hypervariable mini-chromosomes in host-specific lineages of the blast fungus. *bioRxiv* 2020;2020.01.10.901983.
- 56. Langner T, Harant A, Gomez-Luciano LB, Shrestha RK, Malmgren A, *et al.* Genomic
   rearrangements generate hypervariable mini-chromosomes in host-specific isolates of the blast
   fungus. *PLOS Genetics* 2021;17:e1009386.
- 57. Vanheule A, Audenaert K, Warris S, van de Geest H, Schijlen E, *et al.* Living apart together:
   crosstalk between the core and supernumerary genomes in a fungal plant pathogen. *BMC Genomics* 2016;17:1–18.
- 692 58. Condon BJ, Leng Y, Wu D, Bushley KE, Ohm RA, *et al.* Comparative genome structure, secondary
   693 metabolite, and effector coding capacity across *Cochliobolus* pathogens. *PLoS Genetics* 694 2013;9:e1003233.
- 59. Yang G, Turgeon BG, Yoder OC, Bronson CR, Yoder OC, *et al.* Toxin-deficient mutants from a toxin-sensitive transformant of *Cochliobolus heterostrophus*. *Genetics* 1994;137:751–7.
- 60. Klosterman SJ, Subbarao K V., Kang S, Veronese P, Gold SE, *et al.* Comparative genomics yields
  insights into niche adaptation of plant vascular wilt pathogens. *PLOS Pathogens*2011;7:e1002137.
- 61. Gan P, Hiroyama R, Tsushima A, Masuda S, Shibata A, *et al.* Telomeres and a repeat-rich
   chromosome encode effector gene clusters in plant pathogenic *Colletotrichum* fungi.
   *Environmental Microbiology* 2021;23:6004–6018.
- 62. Croll D, Zala M, McDonald BA. Breakage-fusion-bridge cycles and large insertions contribute to
   the rapid evolution of accessory chromosomes in a fungal pathogen. *PLOS Genetics* 2013;9:e1003567.
- 63. Bertazzoni S, Williams AH, Jones DA, Syme RA, Tan K-C, et al. Accessories make the outfit:
   accessory chromosomes and other dispensable DNA regions in plant-pathogenic fungi. *Molecular Plant-Microbe Interactions* 2018;31:779–788.
- Francis A, Ghosh S, Tyagi K, Prakasam V, Rani M, *et al.* Evolution of pathogenicity-associated
   genes in *Rhizoctonia solani* AG1-IA by genome duplication and transposon-mediated gene
   function alterations. *BMC Biology* 2023;21:1–19.
- Fraser JA, Huang JC, Pukkila-Worley R, Alspaugh JA, Mitchell TG, *et al.* Chromosomal
  translocation and segmental duplication in *Cryptococcus neoformans*. *Eukaryotic Cell*2005;4:401–406.

- 66. van Westerhoven A, Aguilera-Galvez C, Nakasato-Tagami G, Shi-Kunne X, Martinez E, *et al.*Segmental duplications drive the evolution of accessory regions in a major crop pathogen. *bioRxiv* 2023;2023.06.07.544053.
- 67. Wilson AM, Lelwala R V., Taylor PWJ, Wingfield MJ, Wingfield BD. Unique patterns of mating
   pheromone presence and absence could result in the ambiguous sexual behaviors of
   *Colletotrichum* species. *G3 Genes*/*Genomes*/*Genetics*;11. 2021. DOI:
- 721 10.1093/G3JOURNAL/JKAB187.
- 68. Bhunjun CS, Phukhamsakda C, Jeewon R, Promputtha I, Hyde KD. Integrating different lines of
   evidence to establish a novel Ascomycete genus and family (*Anastomitrabeculia*,
   *Anastomitrabeculiaceae*) in Pleosporales. *Journal of Fungi 2021, Vol 7, Page 94* 2021;7:94.
- Manners JM, He C. Slow-growing heterokaryons as potential intermediates in supernumerary
   chromosome transfer between biotypes of *Colletotrichum gloeosporioides*. *Mycological Progress* 2011;10:383–388.
- 728 70. He C, Rusu AG, Poplawski AM, Irwin JAG, Manners JM. Transfer of a supernumerary
   729 chromosome between vegetatively incompatible biotypes of the fungus *Colletotrichum* 730 gloeosporioides. Genetics 1998;150:1459–1466.
- 731 71. Roca MG, Davide LC, Davide LMC, Mendes-Costa MC, Schwan RF, et al. Conidial anastomosis
   732 fusion between *Collectotrichum* species. *Mycological research* 2004;108:1320–1326.
- 733 72. Ishikawa FH, Souza EA, Shoji JY, Connolly L, Freitag M, *et al.* Heterokaryon incompatibility is
   r34 suppressed following conidial anastomosis tube fusion in a fungal plant pathogen. *PloS one*;7.
   r35 2012. DOI: 10.1371/JOURNAL.PONE.0031175.
- 736 73. Mehta N, Baghela A. Quorum sensing-mediated inter-specific conidial anastomosis tube fusion
   737 between *Colletotrichum gloeosporioides* and *C. siamense*. *IMA fungus*;12. 2021. DOI:
   738 10.1186/S43008-021-00058-Y.
- 739 74. Schmidt SM, Houterman PM, Schreiver I, Ma L, Amyotte S, *et al.* MITEs in the promoters of
   740 effector genes allow prediction of novel virulence genes in *Fusarium oxysporum*. *BMC genomics* 741 2013;14:1–21.
- 742 75. van der Does HC, Fokkens L, Yang A, Schmidt SM, Langereis L, *et al.* Transcription factors
   743 encoded on core and accessory chromosomes of *Fusarium oxysporum* induce expression of
   744 effector genes. *PLOS Genetics* 2016;12:e1006401.
- 745 76. Mould MJR, Boland GJ, Robb J. Ultrastructure of the *Colletotrichum trifolii-Medicago sativa*746 pathosystem. I. Pre-penetration events. *Physiological and Molecular Plant Pathology*747 1991;38:179–194.
- 77. Damm U, Cannon PF, Liu F, Barreto RW, Guatimosim E, *et al.* The *Colletotrichum orbiculare* species complex: Important pathogens of field crops and weeds. *Fungal Diversity* 2013;61:29–59.
- 750 78. C A-T, BB W, MS O, S D, H Z, et al. Identification and characterization of nucleotide-binding site rich repeat genes in the model plant *Medicago truncatula*. *Plant physiology*;146. 2008.
   752 DOI: 10.1104/PP.107.104588.
- 753

### **Table 1:** Characteristics of *C. destructivum* core and mini chromosomes.

	C. destructivum chromosomes			
	1-10 (except 1B)	1B region	11	12
Total length	48 456 982 bp	1 205 667 bp	1 275 594 bp	812 569 bp
G+C content	54.7 %	52.3 %	48.7 %	50.2 %
Number of protein-coding genes	14882	300	278	171
Proportion of genes by length	61.7 %	30.9 % ***	32.3 % ***	26.8 % ***
Proportion of genes with unknown function	21.3 %	42.0 % ***	28.4 % *	32.2 % *
Proportion of genes with RNA support	77.0 %	52.0 % ***	46.4 % ***	59.6 % <sup>*</sup>
Proportion of CAZyme genes	4.1 %	0.0 % ***	1.4 %	1.2 %
Proportion of effector genes	3.0 %	4.3 %	5.4 % *	5.8 % *
Proportion of SMKG	0.7 %	0.0 %	2.9 % **	0.0 %
Proportion of TE by length	4.4 %	32.8 % ***	32.3 % ***	35.1 % ***

755 Asterisks indicate that the data for chromosomes 1B, 11 or 12 differ significantly from the core chromosomes (Fisher's

756 exact test, \*\*\* P<0.001; \*\* P<0.01; \* P<0.05)

757



760 761 Colletotrichum destructivum LARS 709. Upper panel: Geographical distribution of M. truncatula in the 762 Mediterranean area according to GBIF (2019) and collection locations of the nine ecotypes used in this study. Lower panel: Symptoms produced on the trifoliate leaves of six *M. truncatula* accessions at 4 763 764 days post inoculation with spore suspension of C. destructivum LARS 709. Leaves of the susceptible accession DZA210-5 showed large necrotic lesions, while those of DZA327-7 and ESP155-D were 765 766 completely necrotic. Leaves of the resistant accessions ESP163-E, DZA016-F and R108-C3 showed small 767 necrotic flecks or no visible symptoms. Note that R108-C3 is considered to be *M. truncatula* ssp. 768 tricycla.



771 Figure 2: Microscopic analysis of Colletotrichum destructivum LARS 709 infecting cotyledon tissues 772 of Medicago truncatula. (A-C) Susceptible accession ESP155-D. At 48 hpi (A), melanized appressoria 773 (a) had formed on the plant surface and penetrated epidermal cells to form bulbous biotrophic hyphae 774 (bh). At 60 hpi (B), thin necrotrophic hyphae (nh) developed from the tips of biotrophic hyphae. At 72 775 hpi (C), acervuli erupted from the plant surface, consisting of a melanized, hair-like seta (s) and a mass 776 of conidia (c). (D,E) Resistant accession ESP163-E. At 72 hpi, few appressoria had penetrated cotyledon 777 epidermal cells, and groups of cells underlying the appressoria were pigmented yellowish brown with 778 granular contents (\*). Any hyphae (h) visible inside epidermal cells were typically smaller than the 779 appressorium. Scale bars =  $20 \mu m$ .



782 Figure 3: Schematic representation of the 12 chromosomes of *Colletotrichum destructivum* isolate

**709.** The distribution of genes and transposable elements (TE) across each chromosome are shown784 together with the corresponding genome statistics (inset table).



787 Figure 4: Whole-genome alignments between Colletotrichum destructivum LARS 709 (Cd709) and 788 two Colletotrichum higginsianum strains. Chromosomes of Cd709 (white bars) were aligned with (A) 789 the chromosomes of C. higginsianum IMI 349063 (Ch63, grey bars) or (B) the contigs of C. higginsianum 790 MAFF 304535-RFP (Ch35-RFP, grey bars). Syntenic regions (length >10 kb and percent identity > 88%) 791 were linked together using coloured arcs specific for each chromosome in the Cd709 genome 792 assembly. Red diamonds indicate interchromosomal rearrangements. Black diamonds indicate 793 chromosome breakpoints associated with separate contigs in the Ch35-RFP assembly only. The blue 794 track indicates gene blocks that are unique to Cd709. Note that region chr1B of Cd709 (highlighted in 795 green) has no alignments in either of the C. higginsianum isolates. The black arcs linking chr11 of Cd709 796 to the 3' end of chr6/contig\_9 in C. higginsianum (highlighted in pink) indicate regions with strong 797 sequence similarity (percent identity > 88%) that are smaller than 10 Kb. Asterisks indicate where 798 chromosome sequences were reverse-complemented for better visualization. Tick mark spacing = 1 799 Mb

800



Figure 5: Chromosome 1 of Colletotrichum destructivum has a bipartite structure. (A) Scheme of the 802 803 structure of Cd709 chromosome 1. The probe is specific to the mini-chromosome-like part of the 804 chromosome (chr1B). (B) Detailed scheme of the regions targeted by the 993 bp DIG-labelled probe in 805 Cd709 and in Ch63 (chr11: 493,380 to 494,373). Patterned boxes indicate sequence identity of the 806 target regions to the probe. (C) Pulsed-field gel electrophoresis of chromosomal DNA from C. 807 destructivum isolates LARS 202 (Cd202) and LARS 709 (Cd709) compared to C. higginsianum isolates 808 IMI349063 (Ch63) and MAFF305635 (Ch35). (D) Southern hybridisation. Numerals 1 to 4 indicate 809 signals corresponding to chromosomes displayed in (C).



Figure 6: Codon usage bias in the core and mini-chromosomes of C. destructivum. (A) Principal 812 component analysis (PCA) of codon usage for all amino acids on each chromosome. The region chr1B 813 814 was considered separately from the rest of chr1. The first two axes accounted for 95% of the variance. 815 (B) Plots showing codon usage bias for three amino acids (Alanine, Glutamine, Glycine) in genes located 816 on core chromosomes (1 to 10 excluding region 1B), mini-chromosomes 11 and 12 and region 1B. 817 Codon usage on chr11, chr12 and region chr1B differed significantly from that on core chromosomes 818 (Fisher's exact test, P < 0.001) for the 10 codons presented except GCG (all comparisons) and GGG 819 (chr12 vs core). Other amino acids are displayed in Fig. S6. The significance is reported for all the 820 codons in the Table S12.



823 Figure 7: Circos plot showing Colletotrichum destructivum segmental duplications larger than 10kb 824 found with SDDetector. The green and red tracks represent genes and transposable elements 825 respectively. The light green and orange arcs indicate intra-chromosomal and inter-chromosomal 826 duplications respectively. Duplicated genes are highlighted by blue arcs. The level of sequence 827 similarity along the duplications is shown by a line graph with a colour scale where green indicates 828 greater than 95% similarity, black between 95 and 90% similarity and red below 90% similarity. A sliding 829 window of 100 base pairs was used to calculate and display sequence similarity from large alignments. 830 The scale displayed on the graph ranges from 100% to 85% similarity.