Draft genome sequence of fastidious pathogen Ceratobasidium theobromae, which causes vascular-streak dieback in Theobroma cacao

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Abstract

Background: Ceratobasidium theobromae, a member of the Ceratobasidiaceae family, is the causal agent of vascular-streak dieback (VSD) of cacao, a major threat to the chocolate industry in the South-East Asia. The fastidious pathogen is very hard to isolate and maintain in pure culture, which is a major bottleneck in the study of its genetic diversity and genome.

Result: This study describes for the first time, a 33.90 Mbp de novo assembled genome of a putative C. theobromae isolate from cacao. Ab initio gene prediction identified 9264 protein-coding genes, of which 800 are unique to C. theobromae when compared to Rhizoctonia spp., a closely related group. Transcriptome analysis using RNA isolated from 4 independent VSD symptomatic cacao stems identified 3550 transcriptionally active genes when compared to the assembled C. theobromae genome while transcripts for only 4 C. theobromae genes were detected in 2 asymptomatic stems. De novo assembly of the non-cacao associated reads from the VSD symptomatic stems uniformly produced genes with high identity to predicted genes in the C. theobromae genome as compared to Rhizoctonia spp. or genes found in Genbank. Further analysis of the predicted C. theobromae transcriptome was carried out identifying CAZy gene classes, KEGG-pathway associated genes, and 138 putative effector proteins.

Conclusion: These findings put forth, for the first time, a predicted genome for the fastidious basidiomycete C. theobromae causing VSD on cacao providing a model for testing and comparison in the future. The C. theobromae genome predicts a pathogenesis model involving secreted effector proteins to suppress plant defense mechanisms and plant cell wall degrading enzymes.

Keywords: Ceratobasidiaceae, Chocolate, Rhizoctonia, RNA-Seq, VSD
Background

*Ceratobasidium theobromae* (P.H.B. Talbot & Keane) Samuels & Keane (syn. *Oncobasidium theobromae*, *Thanatephorus theobromae*) is a basidiomycete fungus that causes Vascular-streak dieback (VSD), the second most important disease of *Theobroma cacao* (cacao) in Southeast Asia after *Phytophthora* pod rot (black pod) [1]. While the global loss due to black pod is around 400,000 tons annually, VSD is responsible for approximately 30,000 tons of cacao crop loss, similar to losses due to frosty pod, another disease restricted to South and Central America [1]. Established as the causal pathogen during the 1960s [2], cacao and avocado are the only known hosts of this new encounter disease [3]. The short-lived, wind-borne spores of *C. theobromae* infect the soft young leaves at the branch tip and colonize the leaf xylem. From the leaf, the hyphae reach the stem through the petiole and spread via the xylem vessels and cause vascular necrosis and dieback of the branches, sometimes killing young trees (Fig. 1). The pathogen is endemic to most cacao growing areas of Melanesia and Southeast Asia, causes severe losses locally and contributes to declining productivity and farmers abandoning cacao cultivation. Due to the limited availability of morphological and genetic diversity information and its fastidious nature, *C. theobromae* has been very difficult to study. Currently, the pathogen is identified based on PCR/sequencing of ITS regions [1]. In a recent report, Oberwinkler et al. [4] suggested that the number of nuclei in the hyphal cells was inadequate as a taxonomically defining character and hence, *C. theobromae* should be renamed *Rhizoctonia theobromae*. Within the *Ceratobasidiaceae* family, *Rhizoctonia* spp. are the most widely studied due to their wide host range and global distribution. Though there are at least 5 publicly available genomes of multinucleate *R. solani* isolates with various anastomosis groups (AG1-1A, AG1-1B, AG2-2IIB, AG3 and AG8) [5–9], there are no similar studies on binucleate *Rhizoctonia* spp. (teleomorphs: *Ceratobasidium* spp.).

The xylem-infecting pathogen grows from infected tissue onto water agar or Corticium Culture Medium (CCM) [10], but cannot be maintained in pure culture [11]. Due to its slow growth, other fungi present in the stems, petioles or leaves often overgrow *C. theobromae* during isolation attempts. When isolated in an apparent pure culture, the fungus dies out with repeated attempts at subculturing. The absence of pure culture biomass is the major constraint in obtaining good quality DNA required for molecular studies of *C. theobromae*. After multiple prolonged attempts, we obtained putative mycelia of *C. theobromae* and have carried out whole genome sequencing of the fungus for the first time. *C. theobromae* being fastidious in nature, we used the transcriptome sequence of symptomatic and asymptomatic cacao tissue to confirm the causal agent of VSD and identify its transcriptionally active genes.

Result and discussion

VSD, caused by the fastidious basidiomycete *C. theobromae*, is a serious threat to the cacao industry in South East Asia [1]. To isolate the VSD causing pathogen, we collected stems and petioles showing symptoms of VSD from the Soppeng District of South Sulawesi Province, Indonesia. One isolate (CT2) from these tissues was suspected to be *C. theobromae* based on its mycelial morphology [11] on water agar, and transferred to CCM media. After 3 months of growth, mycelia were harvested and DNA extracted and subjected to ITS sequencing. BLASTn searches of the ITS sequence showed 99.85% similarity to a *C. theobromae* strain South Sulawesi 2 (GenBank: HQ424246). Strains of multinucleate *R. solani* have been assigned to 13 anastomosis groups (AG1 to AG-13). Whereas, strains of binucleate *Rhizoctonia* spp. with *Ceratobasidium* telemorphs have been grouped into 21 anastomosis groups designated as AG-A to AG-U [12]. Publicly available ITS sequences were obtained from GenBank for different *Rhizoctonia* spp. and *Ceratobasidium* spp. representing 30 of the anastomosis groups mentioned above. A phylogenetic study involving the ITS sequence of the *C. theobromae* isolate CT2 along with isolate previously reported from Indonesia and the 30 anastomosis groups shows that *C. theobromae* forms a separate clade and the closest related anastomosis groups are AG-A and AG-K (Fig. 2). Similar results were also reported by Samuels et al. [11] who also confirmed the binucleate nuclear condition of *C. theobromae* mycelia and observed hyphal anastomoses. Using a nested PCR assay, 7 VSD symptomatic cacao stem samples from the Luwu District of South Sulawesi Province, Indonesia were confirmed to be infected by *C. theobromae*, and alignment of the ITS sequences from the 7 samples showed no variation (result not shown). There was one base pair difference between the ITS sequences of CT2 and the set of samples from Luwu (result not shown), indicating possible regional variation among the *C. theobromae* strains. Among the confirmed *C. theobromae* infected VSD symptomatic stem samples, 4 were subjected to RNA-Seq analysis along with 2 asymptomatic control samples.

The initial genome assembly obtained was 44,646,720 bp and consisted of 105,861 contigs with a N50 length of 36,882 bp. Though the average GC content of the assembly was 49.25%, many of the smaller contigs had GC content greater than 60%, indicating the presence of bacterial DNA in the assembly, though the mycelial culture in liquid CCM appeared clean.
Although the bacterial contamination may have been random, another possibility is that the bacterial contaminant may have synergistic effects on the growth of *C. theobromae* possibly being necessary for its growth in vitro and/or contributing to the pathogens fastidious nature. A recent report on the presence of a bacterial endosymbionts in *R. solani* AG 2-2IIIB and its role in virulence [13], makes it relevant to resolve the presence of the bacterial genome along with *C. theobromae*. To remove the bacterial contamination from the genome assembly we combined two post-assembly cleaning approaches. A metagenome binning approach was used to filter out the fungal and bacterial genome. The second approach was based on GC-content and multiple similarity searches to detect potential contaminating sequences in the initial assembly. These post-assembly cleaning procedures have some advantages over pre-assembly cleanup [14].
Engaging MetaBAT2, an automated metagenome binning software tool to reconstruct single genomes [15], 105,861 contigs were grouped into 4 genome bins (Table 1). Bin 1 included 28.29 Mbp, with an average GC content of 49.33% and the closest representation was R. solani. Bin 2 and 3 were also represented as R. solani and the total sequence included was small (Table 1). Altogether, 613 contigs from the three bins correspond to 29.15 Mbp of sequence information and was considered as putative C. theobromae genome sequence. The fourth bin included 2.33 Mbp, with an average GC content of 72.25% and the closest representation was Actinomyces sp. The bacterial genome showed 83.5% BUSCO [16] completeness (Table 1). Further studies are needed to see if the same bacterium is always associated with C. theobromae and has any role in its virulence or fastidious nature. MetaBAT2 was unable to bin 104,801 contigs, which were manually processed based on GC-content and multiple similarity searches. For that, we first eliminated 96,369 smaller contigs (< 200 bp), comprising 7.6 Mbp sequence information as possible bacterial contamination or bad sequence (Fig. 3a). The remaining contigs were subjected to BLASTn searches against the NCBI nucleotide (nt) database (version December, 2018) and a closely related R. solani genome (GenBank: JATN00000000) with an e-value cutoff of 1e−5. In total, 1371 contigs corresponding to 822.5 Kbp of sequence information were identified as bacterial in origin based on BLASTn hits to bacterial genes or genomes. A BLASTn search against the Rhizoctonia spp. identified 2078 contigs corresponding to 2.25 Mbp of sequence information and were considered as additional putative C. theobromae genome sequence. Another 4983 contigs (corresponding to 2.49 Mbp) could not be attributed to either bacteria or Rhizoctonia spp. (Fig. 3a). As a benefit of the doubt, we considered those to be part of the C. theobromae genome. Comparison of GC content and contig

Table 1 Clustering of the initial 105,861 genome assembly contigs (44.64 Mbp) into genome bins using MetaBAT2

| Genome bins | Bin 1 | Bin 2 | Bin 3 | Bin 4 |
|-------------|-------|-------|-------|-------|
| Closest representation | R. solani | R. solani | Actinomyces sp. |
| Total contig length (bp) | 28,287,668 | 496,199 | 368,868 | 2,326,605 |
| Total contig number | 600 | 5 | 8 | 447 |
| BUSCO Completeness (%) | 93.4 | NA | NA | 83.5 |
| Max Contig size (bp) | 589,277 | 263,186 | 109,092 | 30,075 |
| Min Contig size (bp) | 2653 | 25,328 | 9627 | 1509 |
| Mean Contig size (bp) | 47,146 | 49,240 | 46,109 | 5205 |
| N50 Contig length (bp) | 93,535 | 263,186 | 81,760 | 6567 |
| Mean GC content | 49.33% | 49.23% | 50.41% | 72.25% |
| Max GC content | 54.13% | 49.61% | 51.48% | 79.16% |
| Min GC content | 44.57% | 47.88% | 49.44% | 64.56% |

* Based on BLASTx search of the contigs against Nr-database
length of the predicted bacterial and *C. theobromae* contigs show clear difference (Fig. 3b, c). Using BBMap version 37.58 [17], 333,747,598 short reads (92.65% of the initial reads) were aligned to the 7674 contigs considered to be of *C. theobromae* genome. These reads were re-assembled using SPAdes and resulted in a 33.90 Mbp genome sequence with approximately 930× coverage.

The estimated *C. theobromae* genome is smaller than closely related Ceratobasidiaceae like *R. solani* strains (56.02–36.9 Mbp) [5–9] and *Botryobasidium botryosum* (45.75 Mb) [18]. The assembly consisted of 6878 contigs with N50 length of 70,517 bp (Table 2). The overall GC content of the *C. theobromae* genome is 44.81%, which is close to different strains of *R. solani* (43.8–48.4%) [5–9], the only other sequenced species in the Ceratobasidiaceae family. Among the assembled contigs, Contigs 1191 and 8134 were identified to be part of the mtDNA and carry the ITS and 28S ribosomal RNA gene sequence of *C. theobromae*. The ITS sequence acquired was identical to that submitted by Samuels et al. [11] for *C. theobromae*. For quantitative assessment of genome completeness, BUSCO [16] analysis was conducted that indicated *C. theobromae* contains 99.1% of examined loci (85.3% complete genes and 13.3% fragmented genes). The *ab initio* gene prediction generated 9264 protein-coding genes in the *C. theobromae* genome with an average sequence length of 2,168 bp (Table 2). In comparison to the five strains of *R. solani*, the assembled CT2 genome has between 1225 to 4156 less genes.

Functional annotation of the 9264 predicted genes showed that 5364 (58%) could be assigned GO terms and 3055 (33%) could be mapped to the KEGG pathway database (Table 2; Additional file 1: Sheet 1). KEGG pathway analysis of the whole *C. theobromae* genome and two publicly available *R. solani* genomes (AG1-IA: AFRT00000000 and AG3: JATN00000000) showed very similar results, with nearly complete major metabolic pathways for both the species (Additional file 1: Sheet 3). This is another indication that, though the predicted *C. theobromae* genome is smaller with fewer genes compared to *R. solani*, its gene complement is almost complete. Bidirectional BLASTp analysis conducted with *Rhizoctonia* spp. has identified 800 species-specific genes for *C. theobromae* (Additional file 1: Sheet 2). Though the major part of these species-specific genes encodes hypothetical proteins, the most relevant difference was the 11 *C. theobromae* specific putative effector proteins (Additional file 1: Sheet 2).

* C. theobromae* being fastidious in nature, Koch’s postulates are not possible; therefore, we adopted the concept of sequence-based identification of the microbial pathogen [19] and relied on the transcriptome sequence

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**Table 2 Genome assembly and annotation statistics of Ceratobasidium theobromae**

| Contig category | Number of Contigs | Sequence information | % Sequence information |
|----------------|------------------|----------------------|------------------------|
| Total Contigs   | 105,861          | 44,646,720           | 100.0                  |
| Contig <200 bp  | 96,369           | 7,610,465            | 17.0                   |
| Bin 4 (bacterial)| 447              | 2,326,605            | 5.2                    |
| Other bacterial hit | 1371          | 822,507              | 1.8                    |
| Bin 1, 2 and 3 (R. solani) | 613   | 29,152,735           | 65.3                   |
| Other Rhizoctonia spp. hit | 2,078 | 2,248,445         | 5.0                    |
| Undefined       | 4,983            | 2,497,925            | 5.6                    |
| **Total**       | **6878**         | **33,899,105**       | **100.0**              |

**Fig. 3** Detection of bacterial contaminant in the initial genome assembly of Ceratobasidium theobromae. a The percentage of smaller contigs (< 200 bp), contigs sorted by MetaBAT2 and bacterial/Rhizoctonia blast hits over the contigs. b, c illustrate the different distribution of GC contents in the sequences considered as bacterial and putative *C. theobromae*
of symptomatic and asymptomatic cacao tissue. RNA-Seq analysis of four VSD stem segments verified to carry the *C. theobromae* ITS sequence identified 3551 transcriptionally active genes (with \( \geq 10 \) raw reads) from the *C. theobromae* transcriptome, compared to just four *C. theobromae* genes showing any read detection in the two asymptomatic samples (Additional file 1: Sheet 1). The relatively low level detection of *C. theobromae* transcripts in the VSD symptomatic tissue may be due to the low titer of the fungal RNA in the samples. This is consistent with earlier reports that the fungal load of *C. theobromae* in the symptomatic tissues is very low and the same PCR-based detection techniques used here often fail in VSD symptomatic cacao tissues [11].

Self-assembly of non-cacao RNA-Seq reads from the 4 VSD symptomatic tissue samples incorporated reads ranging from 88,978 to 746,046 (Additional file 2: Sheet 1–4), depending on the sample, representing less than 0.2% of the total reads of those samples. The number of de novo assembled transcripts from the four samples ranged from 425 to 2271 (Additional file 2: Sheet 1–4). When compared to sequences in Genbank, most of the de novo assembled transcripts showed homology to the related *R. solani* AG3. Almost every transcript showed higher homology to the *C. theobromae* genes predicted from the genome assembled in this study compared to *R. solani* AG3 genes (Additional file 2). A closer inspection of the alignments between the self-assembled transcripts and the predicted genes indicates that the self-assembled transcripts are much shorter and have gaps compared to predicted genes, but otherwise have near perfect sequence matches (result not shown). This further strengthens our assertion that we have sequenced the genome of the fungus causing VSD in cacao and that the predicted genes from the assembled genome are nearly complete.

After aligning the *C. theobromae* specific RNA-Seq reads (Table 3) from the 4 VSD symptomatic tissue samples against its genes assembled in this study, variant calling generated 4326 putative single nucleotide polymorphisms (SNPs) and small insertions and deletions (INDELs). After filtering the variants (SNPs and INDELs) as mentioned in the methods, a total of 2278 putative CDS (coding DNA sequence)-based variants were obtained in 1051 *C. theobromae* genes (Additional file 2: Sheet 5). Among the 4 libraries, the total number of putative variants (QUAL \( \geq 999 \), \( DP \geq 30 \), and GQ \( \geq 40 \)) ranged between 307 and 1673 (Table 3). The ratios between heterozygous and homozygous variants ranged between 0.021 and 0.063 (Table 3). Though the sampling size is small and these variants are not validated yet, the presence of very low heterozygosity suggests local genetic variability among *C. theobromae* isolates from the South Sulawesi Province of Indonesia may be limited compared to regional variability. Moreover, there was not a single homozygous and heterozygous variant combinations for the same allele in the data set (Additional file 2: Sheet 5) suggesting, at a minimum, the pathogen is homothallic. Although sexual reproduction is known to occur in *C. theobromae* [11], we failed to identify mating type genes within the *C. theobromae* genome.

Proteins secreted by plant pathogenic fungi have the potential to interact with and alter host cells and therefore, their identification and characterization is essential to understanding virulence and the mechanism of infection [20]. As predicted by SignalP version 5.1 [21], 998 proteins were considered potential components of the *C. theobromae* secretome (Additional file 1: Sheet 1), accounting for 10.7% of its proteome. Fungal effectors among the secretome were identified using the machine learning program EffectorP 2.0 [22]. The 138 effector proteins predicted (Additional file 1: Sheet 4) are considered of importance due to their potential to suppress the pathogen-associated molecular pattern triggered plant immunity and are potential candidate genes for future studies on virulence, symptom expression and host range. Although most of these effector proteins have homologs in the *R. solani* genomes, most were identified as hypothetical proteins. The predicted secreted proteins and effectors in five *R. solani* strains range between 391–1142 and 68–126 respectively [7].

To understand the potential *C. theobromae* genes involved in the modification and degradation of cell wall and other organic matter, the predicted genes were analyzed by BLASTp against the Carbohydrate-Active enzymes database (CAZymes) using a threshold value of E \( < 10^{-10} \) and >40% sequence identity. A total of 119 CAZymes families mapping to 664 predicted *C. theobromae* proteins were identified (Table 4). The web server based automated CAZyme annotation using dbCAN2 [23] also generated similar result and identified 663 *C. theobromae* proteins as potential CAZymes. Among the potential CAZymes, 355 are predicted to encode secreted genes.

Table 3 List of filtered transcriptome sequence variants in RNA-Seq libraries from VSD-infected stems

| RNA-Seq library | No. of RNA reads in each library | No. of variants (QUAL \( \geq 999 \), \( DP \geq 30 \), and GQ \( \geq 40 \)) |
|-----------------|---------------------------------|---------------------------------------------------------------|
|                 |                                 | Homozygous alternate                                           |
|                 |                                 | SNPs | INDELs | SNPs | INDELs |
| INS_19          | 108,224                         | 258  | 39     | 10   | 0      |
| INS_23          | 283,192                         | 620  | 135    | 26   | 0      |
| INS_31          | 888,258                         | 1357 | 280    | 34   | 2      |
| INS_57          | 109,242                         | 279  | 37     | 16   | 4      |
proteins based on the presence of a signal peptide (Table 4; Additional file 1: Sheet 1). Glycoside hydrolases (GH) formed the largest group followed by glycosyltransferases (GT) and auxiliary activities (AA) (Table 4). Among the secreted CAZymes, again the largest group is GH followed by AA and polysaccharide lyases (PL). Amongst these large families, PL1, 3, GH28, GH43 and CE8 are often involved in pectin degradation, a component of the plant primary cell wall and middle lamella [24]. Other enzymes (AA9, GH5, GH3, and GH16 for example) often target cellulose and hemicellulose [25]. Together, the genes encoding secreted CAZymes targeting pectin, cellulose, hemicellulose and xylan during infection are expected to support the infection and colonization of cacao by *C. theobromae* (Additional file 1: Sheet 1). In addition to secreted effectors and plant cell degrading enzymes, fungi have a broader arsenal of secreted proteins/enzymes at their disposal when causing disease. Among the non-CAZyme and non-effector secreted protein coding genes, proteases/peptidases are the largest group with known putative functions, followed by lipases and Cytochrome P450 genes (Additional file 1: Sheet 1). However unidentified hypothetical proteins remain the largest group and require more detailed study.

### Conclusions

The assembled *C. theobromae* genome and its analysis provide insight into the genetic makeup of this important cacao pathogen. *C. theobromae* has the tools within its relatively compact 33.89 Mbp genome, like other plant pathogens, to cause plant disease. The *C. theobromae* genome presented supports a typical pathogenesis model, where the fungus secretes effector proteins involved in plant defense suppression along with enzymes required for degradation of cell walls and other cell components. Why the pathogen should be fastidious is unclear and whether its association with *Actinomyces* sp. has anything to do with the virulence or fastidious nature, requires more study. Although, a preliminary variant study suggests limited regional variation, future exploitation of the assembled genome should provide the basis for a better understanding of the genetic variability of the pathogen and support the development of potential disease resistance sources. The effector proteins identified here-in offer potential targets for molecular manipulation and resistance development. Due to the high number of genes that encode hypothetical proteins in this genome and limited detection of *C. theobromae* transcripts in planta, more research is needed to fully understand the disease mechanism of this fungus.

### Materials and methods

#### Isolation and DNA extraction

VSD symptomatic cacao petioles were collected from Soppeng District of South Sulawesi Province, Indonesia. Petioles were surface sterilized by submerging in 6% (v/v) sodium hypochlorite for three minutes followed by three rinses in sterile water. Segments were cut transversely exposing the xylem tissue and placed on water agar. Slow growing mycelium was transferred to liquid CCM [10] then transferred to fresh liquid CCM a second time. After a total of 3 months in liquid CCM at 25 °C, mycelia were harvested in 1.5 ml microtubes and collected by centrifugation (14,000 rpm for 15 min). The mycelial pellet was freeze dried and shipped to USDA-APHIS-PPQ.

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**Table 4 Number of CAZymes family genes of *Ceratobasidium theobromae***

| CAZymes familya | Total genes | Secretedb | Non-secretedb |
|-----------------|-------------|-----------|--------------|
| AA3             | 17          | 5         | 12           |
| AA1             | 15          | 13        | 2            |
| AA5             | 12          | 8         | 4            |
| AA9             | 22          | 22        | 0            |
| Other AA (5)    | 14          | 8         | 6            |
| CBM1            | 15          | 7         | 8            |
| CBM13           | 24          | 15        | 9            |
| CBM5            | 11          | 7         | 4            |
| Other CBM (8)   | 22          | 12        | 10           |
| CE4             | 18          | 14        | 4            |
| CE16            | 13          | 5         | 8            |
| CE8             | 11          | 10        | 1            |
| Other CE (6)    | 17          | 11        | 6            |
| GH0             | 13          | 7         | 6            |
| GH43            | 15          | 7         | 8            |
| GH13            | 13          | 6         | 7            |
| GH3             | 12          | 6         | 6            |
| GH28            | 19          | 17        | 2            |
| GH5             | 24          | 13        | 11           |
| GH16            | 30          | 16        | 14           |
| GH18            | 19          | 7         | 12           |
| GH7             | 12          | 10        | 2            |
| Other GH (44)   | 134         | 71        | 63           |
| GT2             | 15          | 0         | 15           |
| GT4             | 15          | 0         | 15           |
| Other GTs (28)  | 70          | 5         | 65           |
| PL1             | 26          | 23        | 3            |
| PL3_2           | 19          | 17        | 2            |
| Other PLs (5)   | 16          | 11        | 5            |

*a Number within parentheses indicates the number of CAZymes families

*b As determined by SignalPE version 5.1 and BLASTp search against Carbohydrate-Active enzymes database at the threshold value of E < 10<sup>−10</sup> and > 40% similarity
facility in Beltsville, USA and transferred to the USDA-ARS Sustainable Perennial Crops Laboratory in Beltsville after inspection. DNA was extracted from the mycelia as described by Ali et al. [26].

Isolation of RNA from symptomatic plant material
VSD symptomatic and asymptomatic cacao stems were collected from the Luwu District of South Sulawesi Province, Indonesia and cut into 10 cm segments. Samples were freeze dried and shipped as mentioned above. Freeze dried cacao stems were ground in a mortar and pestle in liquid nitrogen and RNA was extracted as described by Bailey et al. [27].

Molecular identification of C. theobromae
For species confirmation of the mycelial DNA samples, PCR amplification of the ITS region of the template DNA of 5 C. theobromae samples, all originating from the same initial water agar culture, were performed using the ITS4 and ITS5 primers described by White et al. [28]. PCR amplification, product purification and sequencing was performed as previously described by Ali et al. [26]. Similarly, to confirm the presence and absence of C. theobromae in the symptomatic and asymptomatic cacao stems, cDNA was synthesized from the extracted RNA as described by Bailey et al. [27] and diluted cDNA (1:20) was subjected to nested PCR assays, first using fungal specific ITS 4 and 5 primers [28] followed by C. theobromae specific primers Than_ITS1 and 2 [11]. PCR products were sequenced as mentioned above.

Molecular phylogenetic analysis of C. theobromae isolates and different Rhizoctonia spp. representing 30 anastomosis groups [29] was carried out based on DNA sequence data of ITS 1 and 2 regions. Sequences were aligned using the ClustalW2 tool [30] under default settings. A phylogenetic tree was reconstructed using the Maximum Likelihood method based on the Poisson correction model [31] in MEGA6 [32].

Genome sequencing, assembly and binning
Ceratobasidium theobromae isolate CT2 genomic DNA was sequenced using Illumina paired-end short-read technology (library preparation and sequencing done by Beijing Genome Institute, Shenzhen, China). For the initial assembly, 360,198,964 short reads (100 bp) were trimmed using BBMap version 37.58 [17] and were assembled using SPAdes Genome Assembler version 3.11.0 [33] in read error correction and assembling mood. K-mers was set at K21, K33, K55. Genome bins were recovered based on tetranucleotide frequencies and read coverage using MetaBAT2 [15] with default parameters.

Ab initio gene prediction
The ab initio gene prediction was performed from the assembly results using AUGUSTUS version 2.7 [34] trained with R. solani AG3 gene models (GenBank: JAT000000000). The predicted proteins were compared against NCBI non-redundant (NR) protein databases by BLASTp to identify biological functions [35]. Open reading frames were also annotated using Blast2GO (http://www.blast2go.com/b2ghome) [36] and the KEGG–database of metabolic pathways [37].

Transcriptome sequencing
To validate the expression of the C. theobromae predicted genes, RNA-Seq was performed on RNA from 4 independent field collected VSD symptomatic cacao stems (validated as carrying C. theobromae above). Two asymptomatic samples were also included. RNA-Seq library preparation and paired-end sequencing was performed using the Illumina HiSeqX platform by Beijing Genome Institute, Shenzhen, China. RNA reads from RNA-Seq libraries ranging from 72 to 104 million reads in fastq format were trimmed up using BBdUk version 37.58 [17], using adapters.fa with parameters ktrim = r, k = 23, mink = 11, hist = 1, tpe, tbo. Trimmed reads were aligned using HISAT2 2.1.0 [38] to the CDS of the C. theobromae genome assembled in this study.

Mining of putative variants
RNA-Seq reads in fastq format from each VSD symptomatic library (INS_19, INS_23, INS_31 and INS_57) were aligned to the CDS of the C. theobromae genome assembled in this study using a memory-efficient short-read aligner Bowtie-0.12.7 [39] and the variant calling and quality control was performed as previously described by Ali et al. [40]. In short, the files were outputted into SAM and converted to BAM format using SAMtools [41]. Variant calling was performed using SAMtools mpileup and bcftools with default parameters. The resulting VCF file was filtered manually using Excel functions. All variants (SNPs and INDELs) with QUAL < 999 and DP < 30 were removed and further filtered based on genotype quality (GQ) and retained if there was at least one library supporting it (GQ ≥ 40) (Additional file 2: Sheet 5).

De novo transcriptome assembly
To further validate the assembled genome of C. theobromae, the causal agent of VSD, we conducted independent de novo transcriptome assemblies of the non-cacao reads from the 4-field collected VSD symptomatic cacao stems as validated above. Trimmed reads for each sample were initially aligned using HISAT2 2.1.0 [38] to the CDS of the cacao genome [42], and the unaligned reads
were assembled using SPAdes Genome Assembler version 3.11.0 [33] with K-mers set at K23. The ab initio gene prediction was performed from the assemblies using AUGUSTUS version 2.7 [34] trained with *R. solani* AG3 gene models. The predicted transcriptomes were compared against *C. theobromae* genome assembled in this study and closely related *R. solani* genome (GenBank: JATN00000000) by BLASTn and against NCBI non-redundant (NR) protein databases by BLASTx analysis.

### Supplementary information

**Supplementary information** accompanies this paper at https://doi.org/10.1186/s40694-019-0077-6.

- **Additional file 1.** *C. theobromae* genome and predicted genes.
- **Additional file 2.** *C. theobromae* de novo transcriptome assemblies and variants.

### Abbreviations

AA: auxiliary activities; AG: anastomosis group; BLAST: basic local alignment search tool; bp: base pair; BUSCO: benchmarking universal single-copy orthologs; CAZYme: carbohydrate-active enzyme; CCM: curtiptom culture medium; CDS: coding DNA sequence; CE: carbohydrate esterase; DP: raw read depth; GC: guanine–cytosine; GH: glycoside hydrolases; GO: gene ontology; GG: genotype quality; GT: glycosyltransferases; INDEL: small insertions and deletions; ITS: internal transcribed spacer; Kbp: kilobase pair; KEGG: Kyoto encyclopedia of genes and genomes; Mbp: megabase pair; NCBI: National Center for Biotechnology Information; PCR: polymerase chain reaction; PL: polysaccharide lyases; RNA-Seq: ribonucleic acid sequencing; SNP: single nucleotide polymorphism; VSD: vascular-streak dieback.

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### Authors’ contributions

Conceived and designed the research: SSA, AA, APF, PM and BAB. Sample collection: AA, APF, AWS, AR, TYK and MJ. Performed the research: SSA, APF and AA. Bioinformatics and data analysis: SSA and JS. Contributed reagents/materials/analysis tools: BAB, PM, DG and LWM. Wrote the paper: SSA and BAB. Provided intellectual and editorial comments: SSA, BAB, PM, LWM and DG. All authors read and approved the final manuscript.

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### Availability of data and materials

The complete nucleotide sequence assemblies and the Whole Genome Assembly of the *C. theobromae* isolate CT2 has been deposited at GenBank under the Accession SSOP000000000, under BioProject PRJNA524910. The combined transcriptome assembly from multiple tissues have been uploaded as additional files.

### Ethics approval and consent to participate

Not applicable.

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**Consent for publication**

All the authors have read the manuscript and gave consent for publication.

**Competing interests**

The authors declare that they have no competing interests.

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