

# Single-Molecule Real-Time Sequencing Combined with Optical Mapping Yields Completely Finished Fungal Genome

Luigi Faino,<sup>a</sup> Michael F. Seidl,<sup>a</sup> Erwin Datema,<sup>b</sup> Grardy C. M. van den Berg,<sup>a</sup> Antoine Janssen,<sup>b</sup> Alexander H. J. Wittenberg,<sup>b</sup> 🕩 Bart P. H. J. Thomma<sup>a</sup>

Laboratory of Phytopathology, Wageningen University, Wageningen, The Netherlands<sup>a</sup>; KeyGene N.V., Wageningen, The Netherlands<sup>b</sup>

L.F. and M.F.S. contributed equally to this work

ABSTRACT Next-generation sequencing (NGS) technologies have increased the scalability, speed, and resolution of genomic sequencing and, thus, have revolutionized genomic studies. However, eukaryotic genome sequencing initiatives typically yield considerably fragmented genome assemblies. Here, we assessed various state-of-the-art sequencing and assembly strategies in order to produce a contiguous and complete eukaryotic genome assembly, focusing on the filamentous fungus Verticillium dahliae. Compared with Illumina-based assemblies of the V. dahliae genome, hybrid assemblies that also include PacBiogenerated long reads establish superior contiguity. Intriguingly, provided that sufficient sequence depth is reached, assemblies solely based on PacBio reads outperform hybrid assemblies and even result in fully assembled chromosomes. Furthermore, the addition of optical map data allowed us to produce a gapless and complete V. dahliae genome assembly of the expected eight chromosomes from telomere to telomere. Consequently, we can now study genomic regions that were previously not assembled or poorly assembled, including regions that are populated by repetitive sequences, such as transposons, allowing us to fully appreciate an organism's biological complexity. Our data show that a combination of PacBio-generated long reads and optical mapping can be used to generate complete and gapless assemblies of fungal genomes.

IMPORTANCE Studying whole-genome sequences has become an important aspect of biological research. The advent of nextgeneration sequencing (NGS) technologies has nowadays brought genomic science within reach of most research laboratories, including those that study nonmodel organisms. However, most genome sequencing initiatives typically yield (highly) fragmented genome assemblies. Nevertheless, considerable relevant information related to genome structure and evolution is likely hidden in those nonassembled regions. Here, we investigated a diverse set of strategies to obtain gapless genome assemblies, using the genome of a typical ascomycete fungus as the template. Eventually, we were able to show that a combination of PacBiogenerated long reads and optical mapping yields a gapless telomere-to-telomere genome assembly, allowing in-depth genome analyses to facilitate functional studies into an organism's biology.

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ver the last years, the emergence and rapid evolution of whole-genome sequencing technologies have profoundly affected genomic research (1). In the past, whole-genome sequencing projects involved cost-intensive and laborious Sanger sequencing that typically delivered highly fragmented genome assemblies. The high demand for resequencing projects, as well as for de novo genome assemblies, has incited the emergence of novel technologies, caught under the umbrella term next-generation sequencing (NGS), which routinely produce gigabases of data in a short amount of time. NGS technologies can be divided in those that generate large amounts of short DNA sequence reads (<500 bp) that are typically characterized by their high quality (1) sequence error per 1 kb) and technologies that produce long reads (>1 kb), although often with relatively low quality (1 to 2 sequence errors per 100 bases) (2). The added value of short reads for *de novo* genome assemblies is typically hampered if repetitive

sequences in a genome are longer than the reads themselves, because these reads will be collapsed into a single element, leading to continuity breaks of the assembly at each genomic location that contains the repetitive element (3). Paired reads, consisting of two reads generated from a single DNA fragment and separated by a known distance, can help to increase the continuity of the assembly, provided that the distance between the pair of reads is longer than the repeat itself (3). However, the sequence spanned by these paired reads will often remain unknown, and therefore, de novoproduced short-read assemblies will mainly comprise nonrepetitive regions (4, 5). Different sequencing strategies, such as singlemolecule real-time (SMRT) and nanopore sequencing, can be used to characterize repeat-rich genomic regions (6-9). Both sequencing strategies generate long reads (up to ~50 kb) that can read through entire repeats and, thus, facilitate more contiguous genome assemblies (6, 10). SMRT sequencing is a well established

Source of	Value for indicated assembly and data sources										
data, metric	VerdaJR2v1.5	VerdaJR2v1.5	VerdaJR2v1.5	SPAdes 3.0	SPAdes 3.0	A5	A5	VDAG_JR2v4.0			
PE library	Х	Х	Х	Х	Х	Х	Х				
MP library	Х	Х	Х	Х	Х	Х	Х				
PacBio P4-C2		Х		Х		Х		Х			
PacBio P5-C3			Х		Х		Х	Х			
Optical map	Х	Х	Х					Х			
Contig metrics											
No. of contigs ( $\geq 0$ bp)	4,514	515	533	2,335	2,463	1,013	1,195	8			
No. of contigs ( $\geq$ 1,000 bp)	3,262	324	338	1,579	1,570	415	419	8			
Longest contig (bp)	99,830	2,178,335	2,251,806	227,026	543,223	2,308,962	2,304,878	9,275,483			
Total length (bp)	33,523,879	35,178,480	35,520,228	34,886,730	35,110,786	36,248,419	36,213,197	36,150,287			
$N_{50}$ (bp)	17,466	662,062	649,303	46,943	50,038	598,861	512,741	4,168,633			
No. of Ns/100 kb <sup><math>a</math></sup>	0	0	0	0	0	0	0	0			
Scaffold metrics											
No. of scaffolds ( $\geq 0$ bp)	9	9	9	1,334	1,510	606	599	8			
No. of scaffolds ( $\geq 1,000$ bp)	9	9	9	659	702	298	285	8			
Longest scaffold	9,141,183	9,180,926	9,215,033	1,263,620	1,066,798	2,912,494	2,937,429	9,275,483			
Total length	37,537,096	38,353,192	38,703,526	34,780,691	34,969,668	36,425,691	36,548,884	36,150,287			
$N_{50}$ (bp)	4,064,734	4,091,407	4,087,047	350,075	306,662	781,486	808,031	4,168,633			
No. of Ns/100 kb	10,691.34	8,277.57	8,224.83	109.82	100.64	652.57	1,082.47	0			

TABLE 1 Statistics for the various genome assemblies of Verticillium dahliae strain JR2

<sup>a</sup> Ns, unknown nucleotides.

technology that produces long reads (on average, >15 kb and up to ~50 kb) that can be used to improve previously generated genome assemblies using a limited amount of data (~10 to 20× genome coverage) (11) or for *de novo* genome assemblies ( $\geq$ 50× genome coverage) (12). While prokaryote genomes can be completely assembled solely based on long reads (12), the assembly of complete chromosomes of eukaryotic genomes is less straightforward.

Optical mapping is a technique for constructing ordered highresolution restriction maps from single DNA molecules in a genomewide fashion. As the technique can be used to align *in silico*-generated restriction maps of genome assemblies, optical mapping can be used to direct the placement of individually assembled sequence contigs onto chromosomes (13–15). Nevertheless, optical mapping is not routinely used in genome sequencing initiatives.

Genomic studies have revealed the importance of noncoding regions, structural rearrangements, and repetitive elements for the lifestyle of many organisms (16–18). As these regions are notoriously difficult to assemble in genome sequencing projects, several approaches have been developed to reconstruct such regions (8, 19–21). In this study, we set out to find an optimal strategy to obtain gapless eukaryotic genome assemblies. To this end, we focused on the genome of a filamentous fungal plant pathogen, *Verticillium dahliae*, with a predicted genome size of 36 Mb, as a model. Previous analyses have revealed extensive genomic rearrangements between individual strains of this species and, furthermore, identified distinct genomic regions that are enriched in repetitive elements (18, 22–24). These findings necessitate the improvement of the current fragmented genome assemblies for this species and, thus, provides an excellent target for our study.

### RESULTS

Long reads increase genome assembly quality. We previously generated a genome assembly of *V. dahliae* strain JR2 based on

paired-end (PE) and mate-pair (MP) reads generated upon Illumina sequencing of short (500-bp) and long (5-kb) insert size libraries (24). In this so-called VerdaJR2v1.5 assembly, optical mapping was used to connect scaffolds, leading to about 4,500 contigs with a contig  $N_{50}$  of ~17 kb (Table 1) (22). In order to reduce the number and length of gaps in this assembly, here we used long reads generated with PacBio sequencing for gap filling and scaffolding (see Table S1 in the supplemental material). To this end, two data sets were generated using a conventional (P4-C2 [P4 polymerase and C2 chemistry]) and an improved (P5-C3) version of PacBio chemistry (see Table S1). Sequencing of four SMRT cells with P5-C3 chemistry resulted in a total yield of 702 Mb (19× predicted average genome coverage), while the P4-C2 chemistry produced 1.7 Gb ( $46 \times$  average coverage) (see Table S1). However, the reads generated with P5-C3 chemistry had an average length of 8.3 kb, while the average read length with P4-C2 chemistry was 6.8 kb (see Table S1). Thus, while the P4-C2 chemistry generated more sequence output, the P5-C3 chemistry generated longer reads. Gap filling and scaffolding with PBJelly2 (version 14.9.9) (11), using long reads derived from both PacBio chemistries independently, significantly improved the genome assembly, resulting in a little over 300 contigs that exceeded 1 kb, while the longest contig reached >2.1 Mb (Table 1). The overall genome assembly improvement after gap filling is furthermore evidenced by the  $N_{50}$  value, which increased from 17.5 kb to approximately 650 kb (Table 1). However, although the contiguity of the genome assembly drastically increased upon the use of long reads for gap filling independent of the applied PacBio chemistry, the final assembly still contained a large amount of unknownnucleotide sequences (Table 1).

In order to assess whether a single-step assembly that combines short- and long-read data, rather than gap filling with long reads of a previously generated assembly based on short reads, would increase assembly quality and contiguity, we assembled the ge-

	Value for data set:											
Metric	SMRT.4	SMRT.6	SMRT.8	SMRT.10	SMRT.12	SMRT.14	SMRT.18					
No. of SMRT cells used	4	6	8	10	12	14	18					
Coverage ( <i>n</i> -fold)	$46.4 \times$	72.1×	96.1×	$120 \times$	143.7×	167.1×	$248 \times$					
Contig metrics with HGAP <sup>a</sup>												
No. of contigs $\geq 0$ bp	246	45	49	41	41	34	35					
No. of contigs $\geq$ 1,000 bp	246	45	49	41	41	34	35					
Largest contig (bp)	957,075	8,522,516	9,231,296	5,501,910	5,496,487	5,496,279	9,231,737					
Total length (bp)	36,019,813	36,390,158	36,496,508	36,298,366	36,439,073	36,407,468	36,472,797					
N <sub>50</sub> (bp)	271,892	3,085,282	4,168,662	2,910,158	3,361,205	3,361,230	3,399,208					
No. of Ns/100 kb <sup><math>b</math></sup>	0.00	0.00	0.00	0.00	0.00	0.00	0.00					
Contig metrics with MHAP <sup>a</sup>												
No. of contigs $\geq 0$ bp	95	132	77	55	50	47	48					
No. of contigs $\geq 1,000$ bp	95	132	77	55	50	47	48					
Largest contig (bp)	3,355,274	1,305,931	3,215,544	3,814,805	5,484,470	4,267,138	5,486,069					
Total length (bp)	36,785,530	35,897,226	36,545,821	36,523,003	36,589,360	36,382,335	36,635,502					
N <sub>50</sub> (bp)	1,816,396	567,445	1,167,265	2,569,351	3,068,688	2,330,944	3,358,862					
No. of Ns/100 kb	0.00	0.00	0.00	0.00	0.00	0.00	0.00					

 TABLE 2 Verticillium dahliae strain JR2 assemblies based on different amounts of PacBio long reads

<sup>*a*</sup> Software used for genome assemblies.

<sup>b</sup> Ns, unknown nucleotides.

nome of *V. dahliae* strain JR2 based on the previously generated PE and MP reads (see Table S1 in the supplemental material) (24) in combination with either of the two PacBio data sets (see Table S1) using SPAdes (version 3.0.0) (25). The assembly based on the P4-C2 data set contained 2,335 contigs with an  $N_{50}$  of ~47 kb and 659 scaffolds of >1 kb (Table 1). Interestingly, similar assembly statistics were obtained with the P5-C3 data set, showing that the differences in yield and read length obtained with the two chemistries did not affect the assembly process (Table 1).

Single-step assembly software packages were typically developed to assemble short reads only and are therefore not optimized to utilize long reads for gap filling. An alternative methodology is a two-step de novo assembly approach in which short reads are assembled, after which gap filling is performed with long reads (20, 26). Thus, we assembled the short reads using the A5 pipeline (version 2014.01.13) (27), followed by gap filling and scaffolding using PBJelly2 (version 14.9.9) (11) with a relatively low (between  $10 \times$  and  $50 \times$ ) average genome coverage of PacBio reads (Table 1; see also Table S1 in the supplemental material). This two-step approach generated approximately 400 contigs of >1 kb with a contig  $N_{50}$  of about 2.3 Mb, thus clearly outperforming the singlestep assemblies (Table 1). The superior quality of the two-step genome assembly is even more evident after scaffolding (Table 1). The gap-filled VerdaJR2v1.5 genome has contig metrics comparable to those of the two-step assembly that did not include optical map data (Table 1), demonstrating that inclusion of optical mapping in the assembly strategy does not improve contiguity. However, when comparing scaffold metrics, the addition of the optical map data resulted in superior genome assembly quality through the placement of more contigs into scaffolds. Nevertheless, the final assembly still contains a considerable amount of gaps (Table 1).

*De novo* genome assembly based on long reads only. In order to assess whether we could assemble the *V. dahliae* JR2 genome based on long reads only, additional PacBio sequencing was performed using the P4-C2 chemistry. In total, 14 SMRT cells were used, resulting in 6 Gb of sequence (167× average genome coverage). This data set was randomly sampled and used to form subsets representing sequencing results from 4 (46×; SMRT.4), 6 (72×; SMRT.6), 8 (96×; SMRT.8), 10 (120×; SMRT.10), or 12 (144×; SMRT.12) SMRT cells, and all data sets were assembled using HGAP (version 2.0), as well as MHAP (version 1.5b1) software (see Table S2 in the supplemental material) (12, 28). The poorest assembly was obtained with HGAP (version 2.0) based on four SMRT cells, resulting in 246 contigs with an  $N_{50}$  of less than 0.3 Mb and a largest contig of less than 1 Mb (Table 2). However, all assemblies based on six or more SMRT cells generated comparable assembly outputs, with a total assembly size of ~36.5 Mb composed of up to 49 contigs, an  $N_{50}$  that exceeded 2.9 Mb, and a largest contig exceeding 5.5 Mb in all cases (Table 2). The fewest contigs, namely, 34, were produced based on 14 SMRT cells.

In order to determine its quality, the genome assembly based on 14 SMRT cells was aligned to the previously generated optical map (24), revealing that only a single contig was wrongly assembled (see Fig. S1 in the supplemental material). Nevertheless, all assemblies based on PacBio sequencing outperformed the hybrid assemblies as long as the sequencing depth exceeded  $72 \times$  (Tables 1 and 2).

Assembly of a gapless genome of *V. dahliae* strain JR2. In an attempt to generate a gapless genome assembly of *V. dahliae* strain JR2, we generated a new assembly using HGAP software (version 2.0) (12) solely based on PacBio reads derived from two types of chemistries, comprising 2.7 million reads equivalent to 8.9 Gb (~248× coverage; 18 SMRT cells) (see Table S3 in the supplemental material). The assembly comprised 35 contigs, with the longest contig being ~9 Mb and an  $N_{50}$  of 3.4 Mb (Table 2). We subsequently aligned the contigs to the previously generated optical map (22), revealing five contigs that represented complete chromosomes (chromosomes 1, 3, and 6 to 8). The remaining three chromosomes could be assembled by ordering 12 contigs based on the optical map, followed by gap filling using PBJelly (version 14.9.9) (11). The gap-filled sequences were polished for errors that

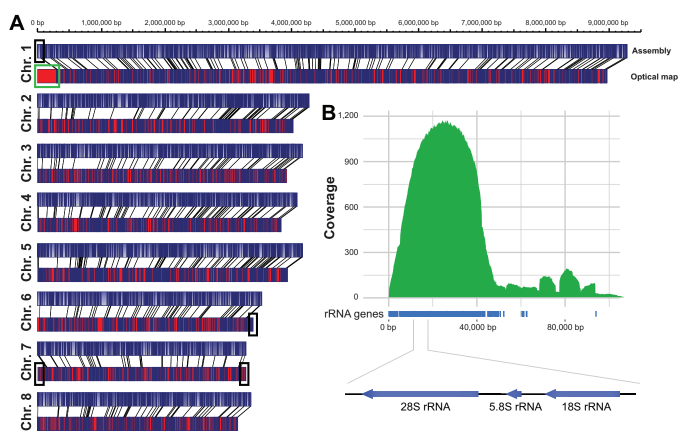


FIG 1 A gapless genome assembly of *Verticillium dahliae* strain JR2. (A) Alignment of the gapless genome assembly of *V. dahliae* strain JR2 with the optical map displays nearly perfect agreement. Represented in blue with blue lines is the genome assembly, while the optical map is represented in red with blue lines. Each blue line represents an NheI restriction site. Black lines represent alignments between the assembly and the optical map. Indicated in black and green boxes are length discrepancies between the assembly and the optical map due to the collapse of repetitive elements in the assembly. (B) Data for rRNA gene cluster located on the distal end of chromosome 1 (see green box in panel A). Local high read coverage (>1,000×) compared with the genomewide average of 15× coverage indicates the collapse of this region during the genome assembly. A single repeat unit of the *V. dahliae* rRNA gene is displayed, and its location in the assembly is marked.

could have been introduced by PBJelly2 (11) using Quiver (12). Thus, 17 of the 35 contigs obtained in the assembly spanned 98.1% of the predicted genome size in eight DNA molecules of contiguous sequence (Table 1), displaying perfect alignment to the optical map except for one edge of chromosome 1, one edge of chromosome 6, and both edges of chromosome 7 (Fig. 1A). However, mapping of the PacBio reads onto the assembly using BLASR (29) revealed a particularly high read coverage at the edges of these chromosomes, indicative of the collapse of repetitive elements. To investigate this hypothesis, de novo annotation of repetitive elements was performed (Table 3), indeed revealing the location of repetitive elements at these regions with high read coverage. More particularly, the high coverage at the edge of chromosome 1 could be attributed to of the collapse of the 300-kb ribosomal DNA repeat region to 50 kb by the assembly software (Fig. 1B). Thus, the length discrepancies between the assembly and the optical map are most likely the result of repetitive element collapse during the assembly process, and we conclude that we obtained a complete and gapless genome assembly of V. dahliae strain JR2.

To further assess the quality of the genome assembly, the correct assembly of the telomeres that comprise the chromosomal ends was assessed. To this end, we investigated the presence of the typical telomeric fungal repeat (TTAGGG) at each of the ends of the assembled chromosomes. Simple repeat analysis identified the typical fungal telomeric repeat in multiple copies (between 7 and 19) on both edges of each of all eight DNA molecules. In conclusion, we assembled the complete genome of *V. dahliae* strain JR2 in eight chromosomes from telomere to telomere without gaps.

Quality assessment of genome assemblies. The quality of genome assemblies is correlated with the quality of the sequencing reads used to generate the assembly. While Sanger and Illumina sequencing produce high-quality reads (2), PacBio sequencing generates long reads of relatively low-quality (~1 to 2 errors per 100 bp) that can only be used for genome assemblies after error correction. To assess how sequencing errors affect genome assemblies generated by PacBio long reads, we used high-quality short reads (PE and MP Illumina data) derived from V. dahliae strain JR2 (30) and mapped them independently onto the genome assemblies, after which sequence variants were identified as a proxy for sequence errors. The smallest amount of errors was observed in the SMRT.18 assembly generated by HGAP (version 2.0) (12) (Table 4). Interestingly, with the exception of the SMRT.4 assembly, all assemblies generated by HGAP (version 2.0) (12) carried fewer sequencing errors than the hybrid assemblies and assemblies generated by MHAP (version 1.5b1) (28) (Table 4). Our data therefore indicate that sequence errors that are intrinsic to long-

	Value for <i>V. dahliae</i> strain:										
	JR2			VdLs17							
Type of element <sup>a</sup>	No. in genome	Coverage (bp) <sup>b</sup>	Coverage (%) <sup>c</sup>	No. in genome	Coverage (bp) <sup>b</sup>	Coverage (%)					
TEs											
SINEs	15	665	0	16	811	0					
LINEs	324	124,209	0.34	311	167,003	0.46					
LTR elements	1,071	2,428,443	6.72	1,006	2,430,766	6.76					
DNA elements	269	114,336	0.32	272	150,768	0.42					
Unclassified	1,557	1,286,043	3.56	1,351	1,098,298	3.05					
Summary of TEs		3,953,696	10.94		3,847,646	10.7					
Other repeats											
Small RNA	125	22,942	0.06	114	18,050	0.05					
Satellites	74	7,336	0.02	71	7,003	0.02					
Simple repeats	10,210	423,998	1.17	10,208	424,918	1.18					
Low complexity	832	40,802	0.11	835	40,340	0.11					
Total amt of repeats		4,446,122	12.3		4,336,001	12.05					

TABLE 3 Summary of tra	insposable elements and other type	es of repetitive elements identi	ified in V. dahliae strains JR2 and VdLs17

<sup>a</sup> TEs, transposable elements; SINEs, short interspersed elements; LINEs, long interspersed elements; LTR, long terminal repeat.

<sup>b</sup> Total bases matching the element.

<sup>c</sup> % of genome covered by the element.

read sequencing do not affect genome assemblies generated by HGAP (version 2.0) (12) as long as sufficient read depth is used (minimum SMRT.6, corresponding to  $72 \times$  average genome coverage). In contrast, genome assemblies based on a low coverage of

long reads, such as the hybrid assemblies and the assembly solely based on a low coverage of PacBio long reads (SMRT.4, corresponding to  $46 \times$  average genome coverage), as well as genome assemblies that are generated using MHAP (version 1.5b1) (28),

 TABLE 4
 Statistics for the various genome assemblies of Verticillium dahliae strain JR2 generated using Quast software and using VDAG\_JR2v4.0

 as the reference genome

	Data	source									
	Library		PacBio chemistry				Scaffold metric (no. of instances)				
Assembly, software used <sup>a</sup> PE					Optical map	Avg PacBio coverage <sup>b</sup>		Misassemblies	Genes that are:		
	PE	MP	P4-C2	P5-C3			SNPs <sup>c</sup>		Complete	Partial	Missing
VerdaJR2v1.5	Х	Х			Х		1,146	493	10,855	570	5
VerdaJR2v1.5	Х	Х	Х			46	988	544	11,271	158	1
VerdaJR2v1.5	Х	Х		Х		19	1,018	511	11,266	160	4
SPAdes 3.0	Х	Х				46	636	251	10,982	447	1
SPAdes 3.0	Х		Х			19	775	223	10,959	463	8
A5	Х	Х				46	661	369	11,349	78	3
A5	Х		Х			19	768	365	11,344	81	5
HGAP											
SMRT.4			Х			46	1,089	21	11,149	167	114
SMRT.6			Х			72	283	16	11,429	1	0
SMRT.8			Х			96	175	12	11,429	1	0
SMRT.10			Х			120	160	18	11,424	1	5
SMRT.12			Х			143	146	21	11,429	1	0
SMRT.14			Х			167	75	5	11,430	0	0
SMRT.18			Х	Х		248	41	13	11,430	0	0
VDAG_JR2v4.0			Х	Х	Х	248	113	0	11,430	0	0
MHAP											
SMRT.4			Х			46	10,270	15	11,410	16	4
SMRT.6			Х			72	15,683	12	11,256	81	93
SMRT.8			Х			96	11,256	15	11,397	25	8
SMRT.10			Х			120	8,521	12	11,411	13	6
SMRT.12			Х			143	7,579	13	11,425	4	1
SMRT.14			Х			167	6,535	14	11,405	12	13

<sup>a</sup> HGAP 2.0 (12) and MHAP 1.5b1 (28) were used to generate assemblies.

<sup>*b*</sup> Average genome coverage of the PacBio data set used for the assembly.

<sup>c</sup> SNPs, single-nucleotide polymorphisms.

which lacks a polishing step after the assembly, are characterized by much higher error rates (Table 4). In conclusion, an error-free assembly of a genome like that of *V. dahliae* can be obtained upon HGAP (version 2.0) (12) assembly of PacBio long reads with  $\sim$ 72× average genome coverage, guaranteeing sufficient coverage for error correction and high sequence contiguity.

To further assess the quality of the different genome assemblies, we performed Quast analyses (31) that use a reference genome assembly and annotation to identify potential misassemblies. Here, we used the final V. dahliae JR2 genome (VDAG\_JR2v4.0) assembly and annotation as a gold standard to investigate the quality of the other genome assemblies, as this assembly has been verified using the optical map. All genome assemblies generated based on long reads are characterized by a high level of contiguity, resulting in a high number of predicted genes (Tables 1, 2, and 4). However, for assemblies generated by MHAP (version 1.5b1) (28), a less-complete gene annotation was inferred compared with assemblies generated by HGAP (version 2.0) (12) (Table 4). Notably, all genome assemblies displayed misassemblies compared with the final genome assembly (Table 4). Therefore, the completeness and, thus, quality of the genome assembly are directly correlated with the amount of predicted genes in the assembly. Incomplete genome assemblies may therefore lead to a considerable underestimation of predicted genes and, thus, directly affect further biological studies.

Assembly of a gapless genome of V. dahliae strain VdLs17 using long reads and optical mapping. Different strains of V. dahliae are characterized by extensive structural rearrangements and chromosomal size polymorphisms (22). We previously showed that, despite a high degree of sequence identity, the genome of V. dahliae strain VdLs17 is structurally rearranged compared with that of V. dahliae strain JR2 (22). Thus, in order to evaluate the robustness of the approach described here, we made an attempt to assembled the genome of V. dahliae strain VdLs17 (23) based on PacBio reads and optical mapping. To this end, we generated about 300,000 reads equivalent to 1.6 Gb (~44× coverage) using P5-C3 chemistry (4 SMRT cells) (see Table S3 in the supplemental material). The assembly based on HGAP (version 2.0) (12) resulted in 119 contigs, with the longest contig being ~2.5 Mb and an  $N_{50}$  of 711 kb (Table 5). We subsequently aligned the contigs to the previously generated optical map of V. dahliae strain VdLs17 (23), revealing that about 98% of the genome was covered by the assembly. Surprisingly, this alignment revealed contig edges with considerable overlap that were not merged by the assembly software. Therefore, these overlapping edges were manually merged, gap filled with PBJelly (version 14.9.9) (11), and polished using Quiver (11, 12). This manual assembly produced eight gapless DNA molecules that matched the optical map almost perfectly (Table 5; see also Fig. S2 in the supplemental material). Similar to the genome assembly of V. dahliae strain JR2, an assembly collapse was identified at the edge of chromosome 1 where the ribosomal DNA cluster resides (see Fig. S2). Furthermore, two additional collapsed regions were identified on chromosome 4 (see Fig. S2). Similar to the V. dahliae strain JR2 assembly, telomeric repeats were found at both edges of each of the eight VdLs17 chromosomes. Thus, we also inferred a gapless genome assembly of V. dahliae strain VdLs17 in eight telomere-totelomere chromosomes.

To determine the improvement of the gapless assembly generated here over the previously generated genome assembly based

 TABLE 5
 Statistics of Verticillium dahliae strain VdLs17 genome assemblies

	Value for assembly using:						
Metric	Sanger sequencing + optical mapping <sup>a</sup>	PacBio + optical mapping					
Contig metrics							
No. of contigs:							
≥0 bp	1,562	119	8				
≥1,000 bp	1,525	118	8				
Largest contig (bp)	216,594	2,545,020	6,210,300				
Total length (bp)	32,902,348	36,288,516	35,973,870				
$N_{50}({\rm bp})$	43,309	711,766	5,894,008				
No. of Ns/100 kb <sup><math>b</math></sup>	0	0	0				
Scaffold metrics							
No. of contigs:							
≥0 bp	9	119	8				
≥1,000 bp	9	118	8				
Largest contig (bp)	6,048,892	2,545,020	6,210,300				
Total length (bp)	36,874,636	36,288,516	35,973,870				
$N_{50}$ (bp)	4,180,501	711,766	5,894,008				
No. of Ns/100 kb	10,770.33	0	0				

<sup>a</sup> Genome assembly described in reference 23.

<sup>b</sup> Ns, unknown nucleotides.

on Sanger sequencing and optical mapping (23), we compared both genome assemblies (Table 5). Surprisingly, a whole-genome alignment displayed a high number of inversions between the two assemblies (see Fig. S3 in the supplemental material). To resolve this observation, the previously generated genome assembly (23) was aligned to the optical map that had been used to generate this assembly. Unanticipated, a large number of assembly mistakes were revealed (see Fig. S4). Closer inspection of the assembly mistakes showed that, although the location of the placement of the scaffolds on the chromosomes was correct, the orientation of many of the scaffolds on the chromosomes was not (see Fig. S4). Thus, using long reads and optical mapping, we were able to identify and correct assembly mistakes in the previously generated and published VdLs17 genome assembly.

Characterization of transposable elements in V. dahliae benefits from gapless genome. The biggest challenge in any genome assembly is the correct assembly of repetitive elements. Usually, relatively long repetitive elements, such as transposable elements (TEs), are poorly assembled in genome assemblies generated based on short reads, leading to an underestimation of the amount of TEs in the genome. However, TEs are important drivers of genome evolution (16) and, therefore, are relevant to many biological processes. For example, in the genomes of many plantpathogenic fungi, TEs are found to accumulate at particular genomic regions, leading to genome plasticity that allows novel virulence factors called effectors to evolve (16). Moreover, TEs have been implicated in a phenomenon called "repeat-driven expansion," where the expansion of plastic genomic regions carrying highly variable genes with roles in pathogen virulence is mediate by TEs (32). Similar to those in the genomes of other fungal pathogens, TEs in the genome of V. dahliae are concentrated at distinct genomic locations and have been associated with genes that are important for virulence (23). Previous analyses have estimated that about 4% of the genomes of V. dahliae strains JR2 and VdLs17

is composed of repetitive elements (22, 23, 33). However, these estimations were likely compromised by the high level of fragmentation of these previous genome assemblies (23, 33). Using the gapless genome assemblies of V. dahliae strains JR2 and VdLs17, we now reinvestigated the amounts and types of TEs that can be found in these genomes. De novo TE prediction was performed on the V. dahliae strain JR2 genome, and about 20 TE families could be classified (see Table S4 in the supplemental material). Out of all of the TE families annotated, we identified 14 retrotransposon families and a few DNA transposon families. The retrotransposon families comprised one LINE (long interspersed element) retrotransposon family and 13 long-terminal-repeat (LTR) retrotransposon families that were further classified based on the presence of predicted open reading frames (ORFs) in the DNA sequence (34, 35). Thus, we identified seven retrotransposon families (VdL-TRE1 to VdLTRE4, VdLTRE6, VdLTRE7, and VdLTRE12) displaying ORFs within the DNA sequence and four retrotransposon families (VdLTRE8 to VdLTRE11) lacking ORFs (see Table S4 in the supplemental material). Retrotransposons displaying ORFs in the DNA sequence are classified as autonomous elements, while retrotransposons lacking ORFs are considered nonautonomous elements. Interestingly, when we compared the TE repertoire of V. dahliae strain JR2 with that of strain VdLs17 that was previously assembled using Sanger sequencing (33), three families of autonomous LTRs (VdLTRE6, VdLTRE7, and VdLTRE12) and all nonautonomous LTR families (VdLTRE8 to VdLTRE11) found in JR2 were lacking in VdLs17, while one family (VdLTRE5) was only identified in VdLs17 and lacking in JR2 (33). However, when using the gapless VdLs17 genome assembly, all the TE families that were missing from previous analyses of the VdLs17 genome (33) were recovered, showing that the two strains have highly identical TE catalogues, which also agrees with their high levels (>99%) of overall genome nucleotide identity (22).

Finally, we reassessed the genomewide abundance of repetitive elements in the finished genome assemblies of *V. dahliae* strains JR2 and VdLs17, using the repertoire of nonredundant sequences generated by the combination of TE families identified in both genomes. LTR retrotransposons, in particular VdLTRE9 (see Table S4 in the supplemental material), are the most abundant TEs in both genomes (Table 3). Strikingly, in total, the repetitive elements in the *V. dahliae* genomes amount to 12% (Table 3), which is 3 times higher than all previous estimates for these genomes (22, 23, 33).

# DISCUSSION

Advances in NGS technologies have allowed biologists to explore the genome sequences of a multitude of organisms across the tree of life to gain insight into their biology (20). However, only in a few cases has the massive amount of sequencing data that are typically obtained with these technologies been adequately transformed in to high-quality genome assemblies. Rather, highly fragmented assemblies that are biased toward genic regions have typically been obtained, while repetitive elements have remained significantly underrepresented. Considering the importance of repetitive elements for eukaryotic genome functioning and evolution, their correct and complete assembly is imperative for full understanding of the biology of an organism (36, 37). This is particularly relevant for filamentous pathogens, as transposable elements have been found to play crucial roles in the evolution of effector catalogues through repeat-driven expansion (16, 22, 26, 38). Fungal repetitive elements typically range in length between a few hundred base pairs and several kilobases (33, 39). Several sequencing and assembly strategies have recently been developed to assemble repeat-rich genomic sequences based on short reads (40, 41). However, these approaches are laborious and still challenging, and thus, their routine application may not be obvious. A way to improve assembly contiguity is by exploiting long reads that can span entire genomic repeats (20). Long-read technologies, such as SMRT and nanopore sequencing, can generate reads that span entire repetitive elements (6–10, 12, 42). Here, we show that the fragmented genome assemblies of the vascular wilt fungus *V. dahliae* that were previously obtained based on Sanger (23) and Illumina (22) sequencing could be significantly improved with the aid of long-read sequencing.

Our assembly results show that an average of  $\sim 20 \times$  genome coverage of long reads is sufficient for gap filling of the original V. dahliae strain JR2 genome assembly (Table 1), which was generated using a combination of short reads and optical mapping (22). Interestingly, the sequence contiguity of the gap-filled VerdaJR2v1.5 assembly was similar to that of the assembly based solely on long reads when using  $\sim 40 \times$  genome coverage (Table 2). This suggests that short reads do not contribute to the assembly quality if the depth of long reads is sufficient. Indeed, recent data on bacterial genomes show that genome assemblies obtained solely using long reads always outperform hybrid assemblies that include short reads, provided that the long-read sequencing depth is sufficient (~40 to  $50 \times$ ) (12, 42). However, when we compared genome assemblies generated by HGAP (version 2.0) (12) and MHAP (version 1.5b1) (28), it appeared that assemblies generated by HGAP (version 2.0) (12) carried fewer sequence errors (Table 4). The smaller amount of sequence errors in the HGAP (version 2.0) (12) assemblies is likely due to an extra step performed during the assembly procedure to polish the sequences and, thus, reduce the sequencing errors, which is lacking in MHAP (version 1.5b1) (28). Interestingly, however, MHAP (version 1.5b1) (28) produces fewer contigs than HGAP (version 2.0) (12) when using a long-read sequencing depth of  $\sim 40 \times$  (Table 2). These results confirm previous data from the MHAP developers indicating that MHAP (version 1.5b1) (28) produces fewer contigs than other software when only a low sequencing depth of long reads is used (28). The MHAP assembly results provide an appealing perspective for the application of long reads to larger genomes, using less data to achieve acceptable assembly statistics. Interestingly, the genome assembly quality did not increase with sequencing depths  $>72 \times$  (Table 2).

Although single repetitive elements and small clusters of repeats may be assembled based on long reads, the assembly of larger repeat clusters like those found in centromeres and in rRNA clusters remains challenging. Previously, optical mapping has been used successfully by us and others to order contigs into chromosomes for various bacterial (43), fungal (22, 44), plant (45, 46), and animal genome assemblies (14, 47). Therefore, the employment of optical mapping to order contigs on chromosomes that were not fully assembled by HGAP appeared to be a critical step to obtain a complete and gapless genome assembly of the two *V. dahliae* strains. Moreover, using optical mapping, we were able to correct mistakes that were generated by the assembly software at a relatively low sequencing depth (Fig. S1 in the supplemental material), as well as a large number of assembly mistakes in the genome assembly of strain VdLs17 that is publically available through the Broad Institute repository (23). Comparison of the new, finished genome assemblies with the previously generated ones of *V. dahliae* strains VdLs17 and JR2 confirmed the assumption that the fragmentation of the original assemblies was mainly due to unassembled repetitive elements. Consequently, the amount of repetitive elements increased from 4% in the previous assemblies (22, 23) to 12% in the completed assemblies (Table 3). This finding will not only facilitate future studies of the role of particular repetitive elements in genome function and evolution but also allow studies addressing the *V. dahliae* genome structure, including, for instance, characterization of centromeric regions.

Nowadays, a prokaryotic Escherichia coli genome can be completely assembled based on sequencing of just a single SMRT cell (42). However, the complete assembly of eukaryotes is much more challenging, and only 10 gapless eukaryotic genome sequences have been reported thus far. Seven of these concern yeasts that have smaller genome sizes (<20 Mb) than filamentous fungi and a small amount of repetitive elements ( $\sim 4\%$ ), while the remaining three concern filamentous fungi, namely, the dimorphic basidiomycete Cryptococcus neoformans, which has a relatively small genome (19 Mb) (48), and the ascomycetes Myceliophthora thermophila and Thielavia terrestris, which, like V. dahliae, belong to the class of Sordariomycetes and have larger genomes (37 to 39 Mb) (49). Importantly, the three filamentous fungal genome assemblies were generated based on Sanger sequencing, and for the two largest genomes, specific target sequencing of repetitive elements was used (49). Such an approach is laborious and expensive and not suitable for high-throughput eukaryotic genome sequencing in a routine manner. Thus, despite their relatively small genome sizes, still no complete, gapless genome assemblies have been reported for fungal organisms based on NGS technologies. Even the assembly of the fungal wheat pathogen Zymoseptoria tritici (formerly known as *Mycosphaerella graminicola*) still suffers from the presence of a few minor gaps despite the fact that this genome is generally portrayed as finished (50, 51). Nevertheless, it can be anticipated that the number of finished fungal genomes will increase significantly in the near future due to the application of long-read sequencing, as well as further dissemination of the use of optical mapping or the integration of sequence-based highresolution genetic or physical maps. More recent technological advances have been able to automate optical mapping, as well as the subsequent imaging and data analysis. The resulting so-called whole-genome mapping can be executed in a high-throughput fashion and is therefore also suitable for more complex genome assembly projects (14). This, together with the ever-increasing read lengths due to improved sequencing chemistries, will bring gapless whole-genome assemblies within reach for complex eukaryotic genomes as well.

**Conclusions.** The complete genome assembly of two *V. dahliae* strains highlights the power of long-read DNA sequencing technology and establishes a standard for *de novo* genome assembly of haploid fungal genomes. We show that  $\sim 50 \times$  PacBio coverage is sufficient to achieve a high-quality genome assembly. However, an error-free assembly of a genome like that of *V. dahliae* can be obtained upon HGAP (version 2.0) (12) assembly of  $\sim 72 \times$  average genome coverage of PacBio long reads. Based on the combination of *de novo* assembly of long reads and optical mapping, we were able to assemble a gapless genome in a cost-effective manner. Technological advances in optical mapping, as well as in sequencing chemistries, will bring gapless whole-

genome assemblies within reach for complex eukaryotic genomes as well. Importantly, finished genome assemblies will disclose genomic information that is imperative to fully appreciate an organism's biological complexity.

#### MATERIALS AND METHODS

**DNA library preparation.** The PacBio libraries were constructed using approximately 10  $\mu$ g of genomic *V. dahliae* DNA that was mechanically sheared to a size of ~22 kb, using g-TUBES (Covaris, Inc., Woburn, MA) according the manufacturer's instructions. PacBio SMRTbell libraries were prepared by ligation of hairpin adaptors at both ends of the DNA fragment (52) using the PacBio DNA template preparation kit 2.0 (Pacific Biosciences of California, Inc., Menlo Park, CA) for SMRT sequencing on the PacBio RS II machine (Pacific Biosciences of California, Inc.) according to the manufacturer's instructions. Libraries were purified using Agencourt AMPure beads (Beckman Coulter, Inc., Brea, CA) to remove short inserts of <1.5 kb. Libraries were size selected using the BluePippin preparation system (Sage Science, Beverly, MA) with a minimum cutoff of 7 kb. The sheared DNA and final library were characterized for size distribution using an Agilent Bioanalyzer 2100 (Agilent Technology, Inc.).

DNA sequence data generation. PacBio sequencing data were generated at KeyGene N.V. (Wageningen, the Netherlands) using the PacBio RS II instrument. The DNA/polymerase binding kit 2.0 (Pacific Biosciences of California, Inc.) was used to anneal sequencing primers and DNA polymerase to the DNA fragments in order to make a complex for small-scale libraries, according to the manufacturer's recommendations. DNA template, polymerase, and primer complexes were diluted to a final concentration of 5 nM. The complexes were loaded onto the SMRT cells (Pacific Biosciences of California, Inc.). The sequencing kit 2.0 (Pacific Biosciences of California, Inc.) was used for sequencing using a 180-min sequence capture protocol along with stage start to maximize read length. Two different polymerases and chemistries were used to generate sequence reads for the V. dahliae strain JR2 genome. The P4 polymerase and C2 chemistry (P4-C2; Pacific Biosciences of California, Inc.) was applied to 14 SMRT cells, while the P5 polymerase and C3 chemistry (P5-C3; Pacific Biosciences of California, Inc.) was applied to four additional SMRT cells. The genome of V. dahliae strain VdLs17 was sequenced using four SMRT cells sequenced with P5-C3 chemistry.

**Sequence assembly.** Assemblies using SPAdes (version 3.0.0) (25) were performed using paired-end (PE) and mate-pair (MP) reads generated upon Illumina sequencing of short (500 bp) and long (5 kb) insert size libraries supplemented with long reads generated by either P4-C2 or P5-C3 PacBio chemistries. Gap filling of the previously generated VerdaJR2v1.5 genome assembly (22) was performed with PBJelly (version 14.9.9) (11), using long reads produced by either P4-C2 or P5-C3 PacBio chemistries. Our previously described hybrid assembly strategy (20) was used in combination with long reads produced with both PacBio chemistries.

Long-read assemblies were generated with HGAP (version 2.0) (12) and MHAP (version 1.5b1) (28) using default settings. Assembled sequences were aligned to the optical map with MapSolver (version 3.2) software (OpGen, Gaithersburg, MD) and manually ordered to compose chromosomes. Gap filling of the manually merged contigs was performed with PBJelly (version 14.9.9) (11) with default settings, except for the assembly stage, where the –maxTrim and –maxWiggle were set at 10,000 bp. Quiver (12) analysis was used to correct errors after gap filling.

**Genome quality assessment.** Telomeric repeats were identified by Tandem Repeats Finder (version 4.07b) (53). Repetitive elements were predicted using RepeatMasker (version 4.0.5) (54). An exhaustive and complete *Verticillium dahliae* TE database was generated by combining repetitive element sequences identified by LTRFinder (version 1.0.5) (55) and RepeatScout (version 1.0.5) (56), TEs previously identified in *V. dahliae* strain VdLs17 genome (33), and the RepBase database (57). Errors in the genome assembly generated based on long reads were identified by using Bowtie2 (58) to align previously generated high-quality short reads of *V. dahliae* strain JR2 (24) onto the genome assemblies that made use of the long reads and subsequent calling of sequence variants using FreeBayes (version 0.9.20) (59). Only sequence variants called with an accuracy higher than 99% and coverage of >10-fold were used in the analysis.

Accession numbers. Assembly data for the complete genomes can be found at the NCBI database under accession numbers GCA\_000400815.2 and GCA\_000952015.1 for *V. dahliae* strain JR2 and VdLs17, respectively.

## SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at http://mbio.asm.org/ lookup/suppl/doi:10.1128/mBio.00936-15/-/DCSupplemental.

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Figure S1, PDF file, 0.8 MB.
Figure S2, PDF file, 0.5 MB.
Figure S3, PDF file, 0.4 MB.
Figure S4, PDF file, 0.7 MB.
Table S1, PDF file, 0.1 MB.
Table S2, DOCX file, 0.02 MB.
Table S3, DOCX file, 0.02 MB.
Table S4, DOCX file, 0.02 MB.
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L.F. conceived of the study, participated in its design and coordination, performed analyses, and helped write the manuscript, M.F.S. participated in study design and coordination, performed analyses, and helped write the manuscript, E.D. generated assemblies for the different longread data sets, G.C.M.V.D.B. produced the biological material used in the study, A.J. participated in the design of the study, A.H.J.W. participated in the design and coordination of the study, and B.P.H.J.T. conceived of the study, participated in its design and coordination, and helped write the manuscript. All authors read and approved the final manuscript.

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