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# Identification and development of new polymorphic microsatellite markers using genome assembly for *Ganoderma boninense*, causal agent of oil palm basal stem rot disease

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Abstract Ganoderma boninense is a telluric lignicolous basidiomycete and the causal agent of basal stem rot, one of the most devastating diseases of the oil palm (Elaeis guineensis). While the fight against G. boninense is of major concern in Southeast Asia, little information is available regarding the genetic diversity and evolutionary history of the fungus. In this context, the development of an informative molecular marker set to characterize the diversity of G. boninense is a key step towards understanding the biology of this pathogen. A G. boninense draft genome sequence of 63 Mbp, assembled using 454 and Illumina sequencing technology, was used to identify and develop a set of microsatellite markers (simple sequence repeats, SSRs). A total of 2487 SSRs were identified, for which 145 SSR primer pairs were designed. These SSRs are characterized by di- to hexanucleotide motifs with 5 to 34 repetitions. Ninety-seven SSR loci were successfully

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amplified on an initial small set of *G. boninense* isolates from Indonesia. A collection of 107 isolates from several regions in Southeast Asia were screened to characterize each locus for allele number, polymorphism information criterion and the presence or absence of null alleles at each locus. These results allowed us to propose an effective set of 17 SSRs for studying genetic diversity within *G. boninense*.

Keywords Ganodermaboninense · Oilpalm · Basalstem rot · Microsatellite · Genome sequence

# Introduction

The availability of new molecular biology tools, along with the exploration of genomic information, has made it possible to develop a large number of genetic markers, including microsatellites. Microsatellites, also known as simple sequence repeats (SSRs), are DNA regions comprising short tandem repeat units from one to six base pairs in length. Polymorphic microsatellites have become one of the most popular and commonly used molecular markers for studying genetic diversity, genealogy and molecular ecology (Jarne and Lagoda 1996). In particular, microsatellites remain the most powerful and widely used tool for studying population genetics and the geographical spread of plant pathogens (Schoebel et al. 2013a, b; Presti and Wasko 2014). This success results from their codominance, high polymorphism, high allele number and relatively high abundance in most genomes, and the ease of typing by polymerase chain reaction (PCR).

Until recently, developing microsatellite markers for a previously unsequenced species involved the construction of an enriched DNA library by cloning sequences in plasmids and sequencing them using the Sanger method (Zane et al. 2002), which was expensive and time-consuming. With the development of next-generation sequencing (NGS) technologies, large amounts of genomic sequence data can be made available at a low cost (\$0.05 for 1 Mbp in April 2014 against \$5292 for 1 Mbp in September 2001; NIH estimate: http:// www.genome.gov/sequencingcosts), providing a faster and cheaper solution for finding microsatellites without a priori knowledge of the target species. In recent years, NGS platforms such as Illumina GAII and Roche 454 have enabled the availability of larger sets of microsatellite markers by exploring complete genome sequences, even for non-model species (Abdelkrim et al. 2009; Zalapa et al. 2012; Cai et al. 2013; Schoebel et al. 2013a, b). Until now, most studies have searched raw NGS data for microsatellite patterns for the design of primers (Gardner et al. 2011). The use of NGS raw data, i.e. raw reads, has a number of limitations. The length of raw reads from the 454 (500 bp) and Illumina (250 bp) systems is often not sufficient to design primers for both sides of the SSR, which is a critical step in developing microsatellite markers (Castoe et al. 2010). Proper primer design for SSR genotyping, therefore, should satisfy several conditions, including a minimum PCR product size of 100 bp, in order to avoid confounding artefacts caused by primer peak or the same melting temperature (Tm) for both forward and reverse primers. Castoe et al. (2010) were able to design primers for only 49.6 % of the SSRs they identified in their data set of 454 raw reads, and Cai et al. (2013) were able to design primers for only 32.7 % to 40.1 % of the SSRs they identified in their Illumina raw read data set. Moreover, sequence redundancy in raw reads poses a major challenge in the design of locus-specific primers, as determining whether reads with similar sequences correspond to a single locus or to duplicate regions is not a trivial task (Castoe et al. 2010). New approaches based on exploring genome assemblies seem to be a promising way of circumventing both problems caused by the use of raw reads (Cai et al. 2013). Read assembly consists in aligning and merging reads into much longer DNA/RNA fragments in order to reconstruct the original molecule. This method thus provides much longer sequences-about 1 to 10 kbp for a raw assembly against a maximum of 600 bp for 454 raw reads or 300 bp for Illumina raw reads (Zalapa et al. 2012; Wei et al. 2014)—and reduces data complexity considerably.

Despite the advent of NGS, developing fungal microsatellite markers is still challenging (Dutech et al. 2007; Schoebel et al. 2013a, b). While the small size of fungal genomes, most ranging from 10 to 60 Mbp (Gregory et al. 2007), makes it easier to explore the genome to find microsatellite loci, it appears that microsatellites are less abundant in fungi than in any other organism (Dutech et al. 2007). Fungal microsatellite motifs are also known to have low numbers of repetitions (most fewer than eight), with a high proportion of mononucleotide motifs, making it difficult to find highly polymorphic SSRs (Dutech et al. 2007). Finally, the cross-species transferability of SSRs in fungi seems to be lower than in other organisms (Barbara et al. 2007; Dutech et al. 2007).

In this study, we developed an SSR marker set from a draft genome assembly of Ganoderma boninense, a telluric white rot basidiomycete that is the causal agent of oil palm basal stem rot (BSR). G. boninense is a non-model plant pathogen fungus whose biology is poorly known (Ho and Nawawi 1986; Pilotti et al. 2002), but it belongs to the same genus as Ganoderma lucidum, a very popular medicinal fungus that has already been sequenced (Chen et al. 2012). G. boninense can affect up to 50 % of palm trees in a plantation, ranging from lower productivity to tree death (Corley and Tinker 2003), and is of major economic importance in Southeast Asia (Turner 1981). As palm oil represents the largest share of the world's vegetable oil market, with 54 million tons produced in 2012 (United States Department of Agriculture: http://apps.fas.usda.gov/), 80 % of which is used by the food industry, BSR may have a serious effect on global food security. Most studies of G. boninense have been carried out to select resistant oil palm progenies (Idris et al. 2004; Durand-Gasselin et al. 2005; Breton et al. 2006) or to develop management practices to limit the spread of the disease (Susanto et al. 2005). To fight G. boninense more effectively, we must improve our understanding of its biology. Our knowledge of the physiology (Pilotti 2005; Rees et al. 2007) and genetics (Miller et al. 1999; Pilotti et al. 2003) of G. boninense is insufficient to explain its current range across an area as large as Southeast Asia. In recent years, population genetics has proved to be an important tool for understanding the biology and spread of plant pathogens (Jarne and Lagoda 1996; Travadon et al. 2012a, b). Genetic diversity patterns can provide information about the respective contributions of clonal and sexual reproduction (Travadon et al. 2012b) and the existence of gene flow barriers (Ali et al. 2014). Neutral molecular markers can be used to reconstruct the main dispersion paths of cosmopolitan or invasive plant pathogens (Montarry et al. 2010), and can even lead to the identification of the original hotspot of an emerging disease (Dutech et al. 2012).

The current work contributes to our knowledge of the genetic diversity of *G. boninense*. In this study, we used a "Seq-Assembly-SSR" approach (Cai et al. 2013) in our preliminary genome assembly, which was based on two sequencing methods (454 and Illumina), in order to develop a suitable set of SSR markers that was then used to produce an initial description of the genetic diversity of a collection of *G. boninense* isolates from southern Asia. As the microsatellite compatibility between closely related fungal species is often described as anecdotal (Dutech et al. 2007), we also tested the transferability of those markers to a closely related species, *G. resinaceum*. The development of this microsatellite marker set would seem to be essential for gaining a better understanding of the

diversity of *G. boninense*, and may be a first step towards improving crop and disease management practices.

# Materials and methods

### **Fungal material**

As G. boninense is the main species found on oil palms (Ho and Nawawi 1985), we based our sampling on the morphological traits of basidiocarps as described by Ho and Nawawi (1986) and Flood et al. (2000). The basidiocarps may be stalked or sessile, flat or bracketshaped. The dorsal surface is glossy, blackish-brown in color, with concentric markings. The edge is white when fresh, and the undersurface is also white. Five G. boninense specimens were collected from infected palms in industrial plantations owned by PT Socfin Indonesia in North Sumatra (Indonesia). Mycelium was purified and maintained in a standard potato dextrose agar (PDA) medium (39 g  $L^{-1}$  PDA from Sigma-Aldrich<sup>®</sup>,  $0.5 \text{ g L}^{-1}$  chloramphenicol, pH 7.0) in the dark at 34 °C. We assumed that these isolates are constituted of dikaryotic mycelium. One isolate, NJ3, which was also used for genome sequencing, and four others were used for the first SSR validation step. To assess the transspecific amplification pattern for the SSRs, we obtained two isolates of G. resinaceum, a species closely related to G. boninense, from the CBS-KNAW Fungal Biodiversity Centre's culture collection (http://www.cbs.knaw.nl/index. php/collection).

One hundred and seven fruiting bodies were collected from peninsular Malaysia (40 fruiting bodies), Borneo (20 fruiting bodies) and North Sumatra (47 fruiting bodies). Geographical locations and sample distribution are summarized in Table 1. Each sample was isolated from a single oil palm tree and identified as *G. boninense* using morphological criteria. Only dikaryotic tissues from the fruiting bodies were kept and freeze-dried before being packed with silica gel.

# **DNA** extraction

Dikaryotic tissues from the fruiting bodies and 2-week-old pure cultures were crushed to powder, and the DNA was extracted using the MATAB [mixed alkyltrimethylammonium bromide] protocol, adapted from the CTAB [cetyltrimethylammonium bromide] protocol (Doyle 1987), with some modifications to improve the DNA extraction yield (Tris HCl 1 M pH 8, NaCl 5 M, EDTA 500 mM, MATAB, PEG 6000 and sodium sulfite). A first step with cell wall digestion by Glucanex (lysing enzymes from *Trichoderma harzianum*, Sigma-Aldrich<sup>®</sup>) buffer (0.5 g Glucanex<sup>®</sup>, 70 ml NaCl 1 M, 30 ml H<sub>2</sub>O, buffer at pH 6) produced a higher DNA yield and removed most of the polysaccharides. The MATAB protocol was also modified for 2-ml tubes to facilitate the extraction of a large set of samples. The DNA concentration (ng  $\mu l^{-1}$ ) of each sample was determined by fluorometry (Invitrogen Qubit<sup>®</sup> 2.0 Fluorometer; Thermo Fisher Scientific) to avoid measurement bias due to the high polysaccharide content.

## Genome sequencing and assembly

The draft genome sequence was obtained using total genomic DNA extracted from the NJ3 isolate. A combination of 454 GS FLX reads (single reads with theoretical length of 400 bp) and Illumina HiSeq 2000 reads (paired reads with insert size of 3 Kb and theoretical length of 50 bp) were used for contig assembly. The reads were assembled using CLC Genomics Workbench 4.6.1 (CLC bio/QIAGEN), with standard parameters, and the N25, N50 and N75 genome assembly statistics were calculated. The N values are standard quality measurement values for genomic data (N25, N50, N75 are calculated by summing the lengths of the largest contigs until N % of the total contig length is reached, and the minimum contig length in this set is the number usually used to report the N value of a de novo assembly). This whole genome shotgun project has been deposited at DDBJ/EMBL/GenBank under accession number LFMK00000000. The version described in this paper is version LFMK01000000.

#### SSR development and primer design

The draft genome of G. boninense (NJ3 isolate) was screened with msatcommander 1.0.8 (Faircloth 2008) to identify SSRs. The parameters were set to search for di- to hexanucleotide perfect SSR motifs with a minimum of five repeats. In order to increase the success of the primer design and to allow for greater flexibility in the choice of fragment length, only candidate SSRs with flanking sequences of at least 400 bp were kept. Primers were designed using Primer-BLAST (http:// www.ncbi.nlm.nih.gov/tools/primer-blast/), with the following requirements: optimum primer size of 20 bp, product size between 150 and 500 bp, primer melting temperatures (Tm) from 57 °C to 63 °C, with optimum at 60 °C, maximum 5bp complementarity between primer pairs, and primer location at least 2 bp away from targeted microsatellite locus. Each primer pair (forward and reverse) was tested with in silico PCR, using the NJ3 genome sequence and a local blastn procedure (Altschul et al. 1990) to select primer pairs that matched on a single genome location. Selected primer pairs were built with an M13 (5'CACGACGTTGTAAAACGA3') (Boutin-Ganache et al. 2001) tail added on the 5' side of the forward primer. This M13 sequence is complementary to the M13 fluoro-tagged primer.

Country	Region	Plantation	Plantation GPS coordinates	Samples
Indonesia	Sumatra	Aek loba	2°37'46.00"N 99°37'42.00"E	AL1, AL2, AL3, AL4, AL5/ AL7, AL8
		Mata pau	3°32'52.67"N 99° 5'11.17"E	311, 312, 318, 319, 325, 334, 341, 347, 357, 366
		PSBB	3°19'5.32"N 99° 2'37.19"E	200, 210, 218, 226, 261, 271, 282, 290, 299, 306
		Sei liput	4°20'32.80"N 98° 3'52.50"E	SL01, SL03, SL05, SL08, SL10, SL12, SL17, SL20, SL22, SL24
		Tanah gambus	3°14'26.70"N 99°26'0.17"E	374, 382, 389, 396, 398, 406, 419, 412, 420, 427
Malaysia	Borneo	Sabah	5°10'52.41"N 119°12'38.20"E	1, 2, 3, 4, 5, 6, 7, 10, 11, 12, 22, 23, 25, 26, 27, 28, 30, 31, 32, 33
	Peninsular Malaysia	Johor bahru	1°44'34.27"N 103°38'52.07"E	70, 71, 72, 73, 74, 75, 76, 77, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91
		Negeri sembilan	2°57'16.08"N 102°29'20.58"E	113, 114, 115, 116, 117, 118, 119, 120, 121, 122, 123, 124, 125, 126, 127, 128, 129, 130, 131, 133

 Table 1
 Geolocation of the 107 Ganoderma boninense samples from the genotyping population

The first in vitro validation of the designed primer pairs was performed on the five Indonesian *G. boninense* isolates maintained as pure cultures (the 5-Indonesian-isolate set) to identify primer pairs exhibiting a mono-locus amplification pattern and polymorphism. Only primer pairs producing mono-locus amplification and polymorphic products were used to genotype the set of 107 *G. boninense* fruiting bodies and two *G. resinaceum* isolates. All primer characteristics are given in Table 2.

### PCR amplification and genotyping

Each microsatellite was amplified in simplex 10-µl PCR reactions containing 5 ng DNA, 100 nM of each of fluorotagged M13 (FAM, NED, VIC, PET), 100 nM of reverse primer, 80 nM of forward primer, 0.2 mM dNTPs, 0.6 mM MgCl<sub>2</sub>, 1X reaction buffer and 0.2 U QIAGEN *Taq* DNA polymerase, 5 % DMSO and BSA at 0.8 µg µl<sup>-1</sup>. After amplification, 2 µl of simplex PCR product for each dye (two markers per dye) were pooled in a plate, and 10 µl of a mixture of GeneScan 600 LIZ size standard and formamide (in a ratio of 1:125) was added to each well for genotyping.

For SSR validation using the 5-Indonesian-isolate set, genotyping was performed on a LI-COR 4300 automated DNA analyzer with IRDye sizing standards of 50–350 bp and 50–700 bp (LI-COR, Inc.). For SSR genotyping of the 107 *G. boninense* samples, PCR products were analyzed using a 3500xl genetic analyzer (Applied Biosystems; Thermo Fisher Scientific). All alleles were scored using GeneMapper software (version 4.1; Applied Biosystems).

# SSR characteristics and diversity analysis

Allele number and genetic diversity are key marker characteristics, as they enable the quantification of marker information content. For each locus, the allele number, observed  $(H_o)$  and expected heterozygosity  $(H_e)$  were calculated using GENETIX 4.05.2 (Belkhir et al. 2004) for all samples together and for each geographical area: peninsular Malaysia, Borneo and Sumatra. Polymorphism information content (PIC) was calculated using the R package PopGenKit, based on Botstein et al. (1980). In addition, the number of observed genotypes was calculated for each locus for both the three geographical areas and the whole area.

Linkage disequilibrium (LD) is used to test for and quantify the statistical correlation between loci or alleles, and in particular, is able to detect whether markers convey redundant information due to their physical linkage. LD analysis between allele pairs was conducted at two levels, first on the whole set of samples and second for each geographical area, using FSTAT 2.9.3. Significance was assessed using a permutation test accounting for Bonferroni corrections with 40800 permutations [http:// www2.unil.ch/popgen/softwares/fstat.htm, developed by Goudet (2002)].

A potential drawback to using SSRs is the presence of null alleles, which for various reasons (Dakin and Avise 2004) fail to amplify to the level of detection. Such null alleles generate false homozygote genotypes and may cause bias in population genetic estimates (Pemberton et al. 1995; Chapuis and Estoup 2007). For example, in genetic structure analysis, null alleles may reduce accuracy in assigning samples to populations and may overestimate F<sub>st</sub> (Carlsson 2008). It is necessary, therefore, to identify loci with such null alleles and to exclude them from further analysis. Null alleles can be identified using indirect methods that account for excess homozygotes at null allele loci compared to other loci. As recommended by Dabrowski et al. (2014), we combined the results of two programs based on indirect methods, ML-NullFreq (Kalinowski and Taper 2006) and Micro-Checker (Van Oosterhout et al. 2004), to determine loci with null alleles. These two programs are complementary in the sense that Micro-Checker has a high detection rate associated with a high false-positive rate, while 
 Table 2
 Characteristics of the 17 polymorphic microsatellite loci developed for Ganoderma boninense

Locus	GenBank accession number	Repeat motif	Primer sequences (5'-3')	F Start/R End (bp)	Tm (°C)	Size (bp)	
3d	KT124389	(CA) <sub>8</sub>	F:AGCCCGCAGAGCCAATGTCC	7382	59.41	197–225	
			R:GGCGTGCGACTGCTGGTTCA	7584	60.25		
6d	KT124390	(AG) <sub>8</sub>	F:ACCTGAGGCGCTGAAAAGCCA	9509	58.97	319–331	
			R:TGACCAGCCCCACCGTCCTC	9817	60.18		
7a	KT124391	(CAG) <sub>6</sub>	F:TCGGGTAGGCTCGCAGGTGG	4916	60.32	338-356	
			R:GGGCCGCACAGGTCGAGAAA	5245	59.35		
11a	KT124392	(CCGA) <sub>5</sub>	F:GCGTGATGAAGTCGGGCGGG	2636	60.45	248-284	
			R:GCAGCCGGGATAGTGCGTGT	2883	59.45		
11d	KT124393	(AGG) <sub>8</sub>	F:TCTGCGGTGGGGGTTGTGCT	8190	60.68	274–289	
			R:GTCATCGCATCGGGGGGTCGG	8456	59.63		
17b	KT124394	(AGC) <sub>19</sub>	F:CGGGAAGTGGTGAACGGTGGT	7604	58.98	234–243	
			R:GGGTGGCTTGACAGCGGCAT	7820	59.97		
18a	KT124395	(CAC) <sub>6</sub>	F:GCGTTGTCGGATGGACGGGG	60050	60.11	263–293	
			R:TGCGGAGGTGGATGTCGGGT	60301	59.90		
19a	KT124396	(AG) <sub>6</sub>	F:GGCCGAGCCGAAAACGTCACA	62519	60.24	208-216	
			R:GGGCGGAGCGAGTCAGTTGT	62717	59.08		
20d	KT124397	(GCA) <sub>9</sub>	F:CGCCATGCCCACCACCAGAG	6268	60.04	283-325	
			R:GACCCGGCTGCCCGAATGAG	6545	59.84		
22a	KT124398	(CAG) <sub>6</sub>	F:TCGGAAGGCAGACGGACGGA	67367	59.62	205-217	
			R:CTAGGCGTGCGTGGCAGAGC	67565	60.45		
24d	KT124399	(CAG)11	F:GCACAGGCACAAGCGCAAGG	10128	59.71	204–267	
			R:CGACGACCGCCCCAAAGGAT	10359	59.14		
33a	KT124400	(CCA) <sub>9</sub>	F:AGCTCCCCTCCCAGCTCCAAC	43018	59.90	171-186	
			R:GAATGCGGCGGGGAAACGGA	43188	60.04		
37d	KT124401	(CGG) <sub>8</sub>	F:TCCACGCCTGAGGGCCAGAA	7092	59.82	311-350	
			R:TCGGAGTCGAAGCCACCCTTCA	7420	59.60		
42d	KT124402	(GT) <sub>9</sub>	F:ACAAGGCTCAAGGCAGCGCA	17415	59.55	212-224	
			R:GCACACCCCAGCAACAGGAGG	17621	59.91		
47d	KT124403	(GA) <sub>10</sub>	F:GGCGACGAGGGCACGAGAGA	392	60.66	273–297	
			R:CCGCACTTTCGCCAACCACC	655	59.09		
51d	KT124404	(CGG) <sub>9</sub>	F:CGAACGCCATCCGAGCAACAGA	8504	59.80	281-317	
			R:TTGGCTCAAAGCCCCGCAAA	8782	57.64		
52d	KT124405	(GA) <sub>8</sub>	F:CCTTGTGGTGGTAGGCGGCA	3911	58.99	313-327	
			R:ATCTCCGTGGCACCGCATCCT	4212	60.31		

ML-NullFreq is less powerful but detects fewer false positives. Null allele detection was carried out independently for the three geographical areas, and we considered that loci with at least five negative results over six (three populations tested with two software programs) could be considered as having no null alleles and kept for further analysis.

A descriptive analysis of the genetic structure of the samples was carried out using the ade4 R package (Dray and Dufour 2007). To measure possible genetic differentiation between geographical areas,  $F_{st}$  was calculated between samples from the three geographical areas and a principal component analysis (PCA) performed to study possible sample differentiation. The calculation of  $F_{st}$  was based on Nei's genetic

distance (Nei 1973), with heterozygosities weighted by group size and significance assessed using a standard permutation test based on Markov chain (1000 permutations). Both PCA and  $F_{st}$  calculations were carried out twice, first with the SSR set with no null alleles and then with the full SSR set.

# Results

#### Genome sequencing and assembly

The 454 pyrosequencing of the NJ3 isolate generated 2,599, 370 reads, with an average read length of 378 bp. Illumina

sequencing generated 276,922,790 reads, with an average read length of 51 bp. After assembly using the CLC Genomics Workbench 4.6.1 software, 21,425 contigs were obtained, with lengths varying from 501 to 89,905 bp and a mean length of 2864 bp. This assembly generated a genome with total length of 63,034,190 bp. The N25, N50 and N75 isolates were 10,965, 5913 and 2563 bp in length, respectively.

# Characterization of microsatellite markers in *G. boninense*

Based on the draft genome assembly, msatcommander software identified a total of 2487 candidate SSR markers in the 21,425 contigs. The number of loci detected for each type of microsatellite loci and the number of repeats decreased from dinucleotide to hexanucleotide motifs, with the exception of the pentanucleotide motif, which had the lowest number of detected loci (Table 3).

The 1687 SSR markers with flanking sequences of at least 400 bp were selected to facilitate primer design. This selection was fairly equitable for most of the SSR types, with a mean of 32.5 % of loci discarded for di-, tri-, tetra- and hexanucleotide SSR markers and 61 % of the candidate pentanucleotide SSR markers. To save time, 145 SSR markers were randomly selected for the design of primer pairs to further assess marker properties. All primer pairs were designed to produce an expected PCR product size ranging from 150 to 400 bp. After in silico PCR was performed, only 110 primer pairs showed the expected mono-locus amplification pattern. Upon completion of in vitro PCR using these 110 primer pairs, 17 primer pairs (15.5 %) showed no amplification, while 93 (84.5 %) showed reproducible mono-locus amplification. Among these 93 primer pairs, 46 (49.5%) exhibited polymorphic amplification products from the five Indonesian isolates. The SSR loci amplified by these 46 primer pairs were then used as markers for genotyping the collection of 107 G. boninense isolates and the two G. resinaceum isolates. After this step was completed, only 17 SSR markers showed proper amplification, i.e. migration patterns characterized by a main peak of over 200 RFU (relative fluorescent units), no more than two amplified peaks,

Table 3 Repartition and characteristics of detected microsatellite loci

Motif	Total number of	Number of repeats				
	detected markers	Minimum	Maximum			
Dinucleotide	1630	5	26			
Trinucleotide	731	5	24			
Tetranucleotide	50	5	16			
Pentanucleotide	18	5	7			
Hexanucleotide	58	5	14			

and a maximum of 10 % missing data. As the primer pairs were designed with a wide range of expected PCR product lengths, for each dye, two markers (differing by at least 16 nucleotides) were able to be multiplexed for genotyping (i.e. 8 markers per plate), increasing genotyping speed for large-scale analysis. All properties of the 17 properly amplified polymorphic SSR markers are summarized in Table 2. The set comprises six dinucleotides, ten trinucleotides and one tetra-nucleotide, with the number of repeats ranging from 5 to 11 and with PCR products 171 to 356 bp in length. All primers have a theoretical melting temperature between 57.6 °C and 60.7 °C and a difference between forward and reverse primers of no more than 2.2 °C (Table 2).

#### SSR characteristics and diversity analysis

We identified 136 alleles for the 17-SSR set in genotyping the 107 G. boninense isolates. The allele number per locus varied from 3 (locus 17b) to 14 (locus 24d), with an average value of 7 (Table 4). The number of identified genotypes across all geographic areas varied from 5 (locus 17b) to 38 (locus 20d), with an average of 17 (Table 4). No PCR amplification failure was observed for five of the loci (locus 7a, 22a, 33a, 42d, 51d), whereas up to 8.4 % failure was found for locus 18a. There was no significant difference between mean observed  $(H_o)$  and expected heterozygosity  $(H_e)$  for any of the geographical areas. PIC values for the 17 SSR markers ranged from 0.298 (locus 17b) to 0.813 (locus 20d). LD was significant (p value<0.01) between alleles for four pairs of loci in the full set of samples: 18a and 47d, 19a and 20d, 3d and 20d, and 11d and 47d (ranging from 0.001 to 0.009). All of these markers belonged to different contigs. Among the various geographical areas, LD was significant only for the pair 19a and 20d in the peninsular Malaysia samples (0.002) and the pair 3d and 20d in the Sumatra samples (0.001).

Based on our decision rules, nine loci (7a, 22a, 33a, 6d, 11d, 20d, 37d, 47d, 51d) showed no convincing indication of null alleles (see supplementary information Table 1), whereas eight loci (11a, 18a, 19a, 17b, 3d, 24d, 42d, 52d) showed evidence for the presence of null alleles.

#### Genetic structure

The main result shown by PCA (first axis, eigenvalue 4.36) was the clear separation between samples from Borneo and those from Sumatra and peninsular Malaysia. Inclusion or exclusion of loci with null alleles did not affect this result (first axis eigenvalue 4.8 %; see Fig. 1a and b). Samples collected from peninsular Malaysia and Sumatra were interspersed within a homogenous cluster, indicating that they belonged to the same genetic group. The PCA results are in agreement with pairwise  $F_{st}$  values calculated between samples from the

Table 4 Genetic diversity characteristics of the Ganoderma boninense 17-SSR set<sup>a</sup>

	Ove	Overall			Indonesia, $n=47$				Malaysian Borneo, n=20				Peninsular Malaysia, n=40					
Locus	N	$N_G$	PIC	$N_p$	$N_A$	$N_{G/P}$	$H_o$	$H_e$	$N_p$	$N_A$	$N_{G/P}$	$H_o$	$H_e$	$N_p$	$N_A$	$N_{G/P}$	$H_o$	$H_e$
07a	7	13	0.408	47	5	8	0.2979	0.4136	20	2	3	0.3500	0.4090	40	6	10	0.3500	0.4541
11a	10	22	0.686	47	5	11	0.5106	0.7156	13	8	9	0.4615	0.8400	39	8	15	0.7179	0.7696
18a	11	27	0.711	47	10	18	0.3191	0.6156	17	6	6	0.0588	0.7968	34	9	16	0.3235	0.7401
19a	5	10	0.545	47	4	8	0.3191	0.6083	20	3	4	0.4500	0.4218	39	5	9	0.5641	0.6810
22a	5	10	0.54	47	5	8	0.5957	0.5749	20	3	4	0.3500	0.3679	40	4	7	0.5750	0.5991
33a	6	13	0.575	47	5	8	0.6170	0.5758	20	5	8	0.8000	0.7590	40	5	9	0.5000	0.6434
17b	3	5	0.298	45	2	3	0.1111	0.4057	20	2	3	0.1500	0.4500	40	3	4	0.1000	0.2291
03d	10	20	0.771	47	7	16	0.5319	0.8065	19	6	11	0.5263	0.7980	40	7	13	0.5250	0.7889
06d	6	14	0.658	47	6	11	0.6596	0.7008	19	4	6	0.4737	0.5277	36	4	8	0.5278	0.6553
11d	6	16	0.568	47	5	10	0.4681	0.5706	20	4	8	0.8000	0.7577	39	6	9	0.4872	0.5311
20d	13	38	0.813	47	13	26	0.8298	0.8296	19	7	12	0.8421	0.8307	40	11	22	0.8000	0.8468
24d	14	29	0.79	46	9	17	0.4130	0.8067	20	8	12	0.4000	0.8346	40	11	18	0.4500	0.7801
37d	9	14	0.505	47	6	9	0.3830	0.4310	20	6	7	0.5500	0.5205	38	5	9	0.5263	0.6281
42d	5	8	0.411	47	4	6	0.1064	0.4482	20	3	3	0.3000	0.3038	40	4	7	0.1750	0.4402
47d	11	21	0.584	47	8	13	0.4894	0.6719	18	6	9	0.6111	0.7206	40	6	8	0.3750	0.5589
51d	8	22	0.717	47	8	15	0.7234	0.7081	20	6	7	0.6500	0.5590	40	7	16	0.7000	0.7924
52d	7	15	0.694	47	6	10	0.3404	0.6495	18	6	9	0.2222	0.8111	39	6	11	0.3077	0.7263
Mean	8	17	0.604	46	6	12	0.4386	0.5986	19	5	7	0.4325	0.5959	39	6	11	0.4360	0.5980

<sup>a</sup> N number of alleles per locus;  $N_G$  number of identified genotypes per locus over all populations; PIC polymorphism information content based on Bolstein et al.'s method;  $N_p$  number of samples positively amplified per locus per population;  $N_A$  number of alleles per locus per population;  $N_{G/P}$  number of identified genotypes per locus per population;  $H_o$  observed heterozygosity;  $H_e$  expected heterozygosity

three geographical areas. For the nine markers without null alleles,  $F_{st}$  was 0.0792 (*p* value 0.0009) between Sumatra and Borneo samples and 0.0773 (*p* value 0.0009) between peninsular Malaysia and Borneo samples, but only 0.0111 (*p* value 0.1778) between Sumatra and peninsular Malaysia samples (Table 5). The results were similar for the 17-marker set (Table 5), with slightly smaller but still significant  $F_{st}$  values of 0.056 between the Borneo and Sumatra samples and 0.061 between the Borneo and peninsular Malaysia samples.

#### Cross-species transferability of SSR marker set

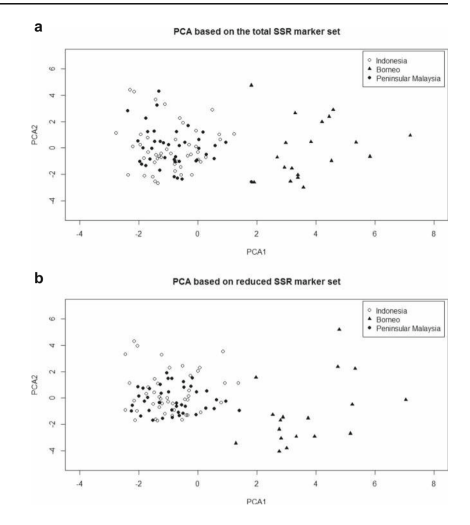
All 17 SSR markers in the *G. resinaceum* CBS19476 strain were successfully amplified, and 16 of the SSR markers in the *G. resinaceum* strain CBS22036 were able to be amplified, with locus 18a yielding no PCR products. For the complete 17-SSR marker set, only two new alleles were discovered on the *G. resinaceum* isolates. Locus 37d had a specific allele for strain CBS22036 only, with a length of 326 bp, which is within the range of detected allele sizes in *G. boninense* (311–350 bp). Locus 52d had a specific allele for both strains of *G. resinaceum*. This allele was 309 bp in length, outside the range of sizes found for *G. boninense* isolates (313–327 bp).

#### Discussion

In the present study, we first identified 2487 SSR loci in a G. boninense draft genome assembly. Then, focusing on the fastest way to obtain a workable minimum microsatellite set for genetic diversity analysis, we developed 17 SSR markers. These markers exhibited satisfactory amplification and polymorphism properties for large-scale population genotyping. To reduce the time and cost of genotyping, these SSR markers were designed to be used in multiplex. This is the first set of locus-specific SSR markers developed for G. boninense. Until now, only random amplified microsatellites (RAMS) (Zakaria et al. 2005), rapid amplification of polymorphic DNA (RAPD) (Abu-Seman et al. 1996; Zakaria et al. 2005) and internal transcribed spacer (ITS) (Moncalvo et al. 1995b; Smith and Sivasithamparam 2000; Utomo et al. 2005) markers have been developed for pathogenic strains of Ganoderma species found on oil palms.

Our study confirms the advantages of using assembled reads (Seq-Assembly-SSR) rather than raw reads for the detection and development of SSR markers for non-model species, as previously shown by Cai et al. (2013). We were able to design primers for 100 % of the selected SSRs, compared to 49.6 % for studies based on 454 raw reads, and 30 to 40 % for studies based on Illumina raw reads. Among the 145 primer

Fig. 1 Global PCA results based on two sets of markers. a Global genetic diversity observed in PCA analyses based on total SSR marker set. b Global genetic diversity observed in PCA analyses based on the reduced 9-SSR marker set including only markers with no predicted null alleles



pairs tested in vitro, 93 (84.5 %) showed mono-locus amplification, which is comparable to results reported by Cai et al. (2013), who obtained 90.7 % locus-specific amplification from primer pairs designed using a genome assembly. The effectiveness of the Seq-Assembly-SSR approach in targeting locus-specific markers and primer design is the result of two improvements. First, assembled contigs provide longer

Table 5 F<sub>st</sub> calculation based on three populations

	Sumatra/Peninsular Malaysia	Sumatra/ Borneo	Peninsular Malaysia/ Borneo
9 Mark- ers	0.0111	0.0792**	0.0773**
17 Mark- ers	0.0133	0.0564**	0.061**

Pairwise  $F_{st}$  based on the method described by Nei (1973); 9 markers figures the set containing markers with a negative result to both allele null tests at 99 %; 17 markers figures the total set of microsatellite markers; \*\* indicates significance for *p* value<0.01 after 1000 permutation

sequences, defining primers with an average contig length of 2864 bp after assembly, compared to raw reads of 51 bp for the Illumina and 378 bp for the 454 technique. Second, sequence redundancy is reduced by the assembly process: while the amount of raw reads was around 4 Gb, the assembly process reduced the total amount of data to around 61 Mb. This reduction in redundancy improves the efficiency of prescreening for primers amplifying single-locus PCR products using in silico PCR (Cai et al. 2013). Furthermore, the selection of microsatellite loci with flanking sequences of at least 400 bp facilitates primer design and multiplexed genotyping, which reduces genotyping costs for large sets of samples.

The number of SSRs detected in the *G. boninense* genome draft assembly in this study falls within the range of numbers of SSRs detected in fungal genomes as reviewed by Karaoglu et al. (2005), which ranged from 124 (*Encephalitozoon cuniculi*) to 8376 (*Neurospora crassa*). However, the number detected here is almost twice that of SSRs identified in *G. lucidum* (1584) by Qian et al. (2013). As our *G. boninense* genome assembly is only a draft, no firm conclusions can be drawn. However, a comparison of the

estimated genome size of these two fungi (43 Mb for *G. lucidum* and 65 Mb estimated for the *G. boninense* draft assembly), as well as the much larger number of SSRs, suggests that there may have been a large duplication or deletion event at genome scale in the *Ganoderma* genus.

While most studies recommend a minimum of 8 to 11 SSR markers for performing sound statistical analysis of population genetics data (van Asch et al. 2010), our study developed 17 SSR markers with good amplification and polymorphism properties, which should allow comprehensive large-scale population studies in G. boninense. Fungus population studies have used on average 10 SSR markers and generally no more than 16 SSR markers (Schoebel et al. 2013a, b; Wang et al. 2014). Of the 17 SSR markers developed in this study, the mean PIC value was approximately 0.604, and 14 of the 17 markers presented PIC values greater than 0.5, indicating that they are highly informative (PIC > 0.5), according to Botstein et al. (1980). Compared to a mean PIC value of 0.33 obtained for SSRs described on another Polyporales by Zhang et al. (2015), our SSRs show high polymorphic information content. Among our 17 SSRs, 9 had no null alleles according to criteria used in this study. While the LD detected at the whole sample scale may be the result of sample structure, LD was significant for only two pairs of alleles within one of three populations, indicating that the information conveyed by each SSR marker was non-redundant. Visualization of putative relationships among investigated samples by PCA showed clear differentiation of a group of samples from Borneo against a mixed group of samples from Sumatra and peninsular Malaysia. PCA results were confirmed by  $F_{st}$  values: about 0.07 between Borneo and Sumatra or Borneo and peninsular Malaysia, while only about 0.01 between Sumatra and peninsular Malaysia. PCA and  $F_{st}$  results were similar regardless of whether the full 17SSR set or the set of 9 SSR with no null alleles was used. This indicates a potential high gene flow between the Sumatra and peninsular Malaysia populations, even though they are separated by the Malacca Strait, and suggests that the Borneo population is partially isolated.

Amplification of 16 of the SSRs for two *G. resinaceum* isolates, a species closely related to *G. boninense* (Moncalvo et al. 1995a, b; Binder et al. 2013), raises the question of species discrimination using this SSR set and species delimitation within the *Ganoderma* genus. SSRs are often considered to be poorly transferable between fungal species (Dutech et al. 2007; Schoebel et al. 2013a, b), with the exception of certain particular cases involving species complexes (Perez et al. 2012). The non-transferability of SSR loci between *G. lucidum* and *G. boninense*, however, were able to be verified by comparing the flanking sequences of the SSRs we detected for *G. boninense* to sequences in the *G lucidum* genome (Chen et al. 2012), although it was impossible to identify motifs of at least 20 bp that could be used to design transspecific SSR primers. Finally, the limited results for

*G. resinaceum* prevent us from drawing any firm conclusions. A larger number of individuals from *G. resinaceum* and the sequencing of fungi barcode markers such as ITS or tubulin genes (Moncalvo et al. 1995a, b) would be necessary to determine the species status of these two taxa.

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