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Prevalence and diversity of TAL effector-like proteins in fungal endosymbiotic *Mycetohabitans* spp.

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Abstract

Endofungal Mycetohabitans (formerly Burkholderia) spp. rely on a type III secretion system to deliver mostly unidentified effector proteins when colonizing their host fungus, Rhizopus microsporus. The one known secreted effector family from Mycetohabitans consists of homologues of transcription activator-like (TAL) effectors, which are used by plant pathogenic Xanthomonas and Ralstonia spp. to activate host genes that promote disease. These 'Burkholderia TAL-like (Btl)' proteins bind corresponding specific DNA sequences in a predictable manner, but their genomic target(s) and impact on transcription in the fungus are unknown. Recent phenotyping of Btl mutants of two Mycetohabitans strains revealed that the single Btl in one Mycetohabitans endofungorum strain enhances fungal membrane stress tolerance, while others in a Mycetohabitans rhizoxinica strain promote bacterial colonization of the fungus. The phenotypic diversity underscores the need to assess the sequence diversity and, given that sequence diversity translates to DNA targeting specificity, the functional diversity of Btl proteins. Using a dual approach to maximize capture of Btl protein sequences for our analysis, we sequenced and assembled nine Mycetohabitans spp. genomes using long-read PacBio technology and also mined available short-read Illumina fungal-bacterial metagenomes. We show that btl genes are present across diverse Mycetohabitans strains from Mucoromycota fungal hosts yet vary in sequences and predicted DNA binding specificity. Phylogenetic analysis revealed distinct clades of Btl proteins and suggested that Mycetohabitans might contain more species than previously recognized. Within our data set, Btl proteins were more conserved across M. rhizoxinica strains than across M. endofungorum, but there was also evidence of greater overall strain diversity within the latter clade. Overall, the results suggest that Btl proteins contribute to bacterial-fungal symbioses in myriad ways.

Impact Statement

Many Mucoromycota fungi harbour endosymbiotic bacteria, including *Rhizopus* spp. that are food fermenters and pathogens of plants and immunocompromised people. *Rhizopus microsporus* has endofungal *Mycetohabitans* (formerly *Burkholderia*) spp. that deploy proteins related to DNA-binding 'transcription activator-like' (Btl) effectors of plant pathogens, which enter plant nuclei and activate disease susceptibility genes. By sequencing isolated bacteria and mining fungal holobiont sequences, we found Btl proteins in diverse *Mycetohabitans* strains, varying in predicted DNA binding specificity, and thus in potential host

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Keywords: effectors; endofungal bacteria; long-read sequencing; meta-assembled genomes; Mycetohabitans; Rhizopus.

Abbreviations: ANI, average nucleotide identity; Btl, *Burkholderia* TAL-like; COGs, Clusters of Orthologous Genes; HMM, hidden Markov model; JGI, Joint Genome Institute; MAG, metagenome-assembled genome; Mef, *Mycetohabitans endofungorum*; Mrh, *Mycetohabitans rhizoxinica*; NLS, nuclear localization signal; RVD, repeat variable diresidue; SCG, single-copy core gene; TAL, transcription activator-like; T3SS, type III secretion system. NCBI accession numbers are available in Table 1 and the Data Availability summary as individual data types were deposited in different ways.

Data statement: All supporting data, code and protocols have been provided within the article or through supplementary data files. Three supplementary figures and two supplementary tables are available with the online version of this article.



targets. Btl proteins were more conserved within *Mycetohabitans rhizoxinica*, suggesting distinctions among the two named species. The results in the context of phenotypic differences observed in other studies suggest that Btl proteins contribute to symbiosis in diverse ways, providing insight into effector evolution and arguing for functional characterization of additional Btl proteins to understand establishment and maintenance of these important fungal–bacterial interactions.

DATA SUMMARY

All genomic data are available through the corresponding NCBI accession numbers provided in Table 1. For each of the nine assembled genomes, *btl* genes are annotated within the genome accession. For the metagenome-assembled genomes, *btl* genes and fragments have been deposited separately as a third-party annotation (Accessions: BK063803–BK063866). Fasta files for the Btl protein amino acid sequences and scripts from R analysis are available on GitHub (cartercharlotte/BtlDiversity).

INTRODUCTION

Effector proteins are synthesized and secreted by bacterial pathogens and mutualists to manipulate their hosts. Investigation of effector protein families reveals mechanisms for disease, resistance and mutualism, while assessing their evolutionary patterns provides insight into host jumps and commonly targeted host pathways and processes. While some effectors are found across species, many are orphans or belong to smaller groups. Transcription activator-like (TAL) effectors, first discovered in the pepper and tomato pathogen *Xanthomonas euvesicatoria*, have been found in several plant pathogenic *Xanthomonas* (TALEs) and *Ralstonia* (RipTALs) spp. [1]. TAL effectors act in the host as transcription factors to directly upregulate specific host genes. Flanking a central, repetitive DNA recognition domain in TAL effectors are termini containing a signal for the type III secretion system (T3SS), multiple nuclear localization signals (NLSs) and an activation domain [2]. Each repeat in the central domain is composed of 33–35 amino acids that are highly conserved except for two that vary, termed the repeat variable diresidue (RVD) [3]. The target DNA sequence for a TAL effector can be predicted based on the RVDs present, as each RVD dictates the nucleotide-binding specificity of its repeat in a one-to-one manner [4, 5]. Using *tal* gene knockouts, host RNA-sequencing (RNA-seq) and computational prediction of binding sites, in a number of plant species TAL effectors have been found to activate susceptibility genes that benefit the pathogen, including sugar transporters known as SWEETs, transcription factor genes and a putative sulphate transporter [2]. Notably, not all plant pathogenic *Xanthomonas* and *Ralstonia* spp. or strains within a species have TAL effectors, while some species have more than 20 encoded in their genome [5].

Outside of plant pathogens, TAL effector-like protein sequences have been identified in the marine metagenome, referred to as marine organism TALE-like, i.e. 'MorTL', proteins [6], and in endofungal *Burkholderiales*, referred to as *Burkholderia* TAL-like, i.e. 'Bat' [7] or 'Btl' [8], proteins. Both MorTL and Btl proteins recognize DNA in the same modular fashion as TAL effectors, and they have attracted some interest alongside TAL effectors as building blocks for DNA-targeting reagents [6, 9]. Complete MorTL protein sequences have not yet been obtained, but some Btl protein sequences have. Compared to TALEs and RipTALs, the N- and C-termini of Btl proteins are much shorter, with no clear activation domain, though there is an N-terminal T3S signal and a single, C-terminal NLS [7, 8]. Also, the repeats of the Btl DNA recognition domain are altogether less conserved and contain some RVDs not yet observed in other TAL effectors [8]. While the taxonomic source of MorTL proteins is unknown, the identification of Btl proteins in endofungal *Burkholderiales*, now classified as *Mycetohabitans* spp. [10], has opened the door to a better understanding of TAL effector evolution and of bacterial–fungal symbiosis.

First identified in 2005, *Mycetohabitans* (formerly *Burkholderia*) *rhizoxinica* (Mrh) is a facultative endosymbiont of *Rhizopus microsporus*, a Mucoromycota fungus that causes rice seedling blight and is an opportunistic human pathogen [11]. Mrh and the related species *Mycetohabitans endofungorum* (Mef) have been studied for their production of secondary metabolites [12, 13], their control of *R. microsporus* reproduction, both sexually and asexually [13, 14], and the mechanisms of their endofungal lifestyle. The bacteria rely on a T3SS to colonize the fungus, but the effectors they secrete through that pathway are unknown, except for Btl proteins [15]. Mechanistic studies have focused on the Mrh type strain B1 [16] and on Mef strain B13 [17], based on discovery order and tractability in a laboratory setting, leaving the understanding of overall diversity within *Mycetohabitans* limited [8, 18]. The prevalence of *Mycetohabitans* spp. across *Rhizopus* spp. has been surveyed in a few studies [18, 19], suggesting <20% of fungal isolates have endosymbionts. Accessibility of documented symbiotic isolates, lack of symbiosis documentation in culture collection accession metadata and the small scale of screening conducted so far have restricted the ability to conduct large-scale analyses of *Mycetohabitans*.

In earlier work, we functionally characterized the only Btl protein, Btl19-13, from Mef strain B13 [8]. It is not required for symbiosis, but it alters the host transcriptome and increases host tolerance to detergent stress, hinting at an influence on cell membrane composition. A Btl protein from a different strain, Btl18-14, with one fewer repeat than Btl19-13 and some differences in the sequence of RVDs, was unable to rescue the *btl19-13* knockout. Adaptation to allelic variation at the target in the two

Table 1. Fungal accessions and their bacterial symbionts examined in this study with the number of btl genes or gene fragments detected in each

Fungal accession	Fungal species	Bacterial strain IDs	Bacterial species	Substrate	Location	No. of btl hits	GenBank accession(s)
Our sequencin	g assemblies – long-rea	ad PacBio					
ATCC 52813	R. microsporus	HKI 0402/ B4/B13	M. endofungorum	Soil	Ukraine	2	CP132744-CP132745
ATCC 52814	R. microsporus	HKI 0403/ B7/B14	M. endofungorum	Soil	Georgia	3	CP132741-CP132743
ATCC 52812	R. microsporus var. Chinensis	B12	M. rhizoxinica	NA	USA	4	CP062175-CP062177
NRRL 5546*	R. microsporus	B46	Mycetohabitans sp.	Soil	Brazil	2	JADBGL000000000
NRRL 5547*	R. microsporus	B47	M. rhizoxinica	Soil	Phillipines	4	CP062173-CP062174
NRRL 5549	R. microsporus	B49	M. rhizoxinica	Rabbit dung	Wisconsin, USA	3	CP062171-CP062172
NRRL 5560	R. microsporus	B60	M. endofungorum	Corn	USA	5	CP062168-CP062170
CBS 112285	R. microsporus	HKI 0456/B5	M. endofungorum	Ground nuts	Mozambique	1	CP062178-CP062179
CBS 111563	R. microsporus	HKI 0455/B3	M. endofungorum	Rice wine tablet	Vietnam	1	CP062180-CP062181
NCBI public as	ssemblies and genes - s	short-read Illumina	ı				
ATCC 62417	R. microsporus	HKI 0454/B1	M. rhizoxinica	Rice	Japan	3	GCF_000198775.1
CBS 308.87	R. microsporus	HKI 0404/B8	Mycetohabitans sp.	Human infection	Australia	1	MN840541.1
CBS 261.28	R. microsporus	HKI 0513/B6	M. rhizoxinica	NA	USA	3	MN840537.1, MN840543 MN840546.1
ATCC 20577	<i>R.</i> sp. strain F-1360	HKI 0512/B2	M. rhizoxinica	Soil	Japan	3	MN891944.1, MN840542 MN840547.1
ZygoLife proje	ct assemblies – short-r	ead Illumina					
IMI 266680	R. oryzae		M. endofungorum	Soil	Sri Lanka	3	SRR12278013
NRRL 1514	R. oryzae		M. rhizoxinica	NA	NA	2	SRR9650540
NRRL 2582	R. oryzae		M. rhizoxinica	Hospital mattress	Ohio, USA	7	SRR9029062
NRRL 2934	Rhizopus sp.		M. rhizoxinica	Pack rat dung	California, USA	1	SRR9650546
NRRL 3373	Rhizopus sp.		Undetermined	NA	NA	2	SRR9650549
NRRL 5548	R. microsporus		M. rhizoxinica	Hospital	Illinois, USA	2	SRR9029145
NRRL 5550	R. microsporus		M. rhizoxinica	NA	New York, USA	3	SRR9029124
NRRL 5551	R. microsporus		M. endofungorum	NA	Philippines	7	SRR9712541
NRRL 5552	R. microsporus		M. rhizoxinica	Sediment	Ohio, USA	3	SRR9712532
NRRL 5553	R. microsporus		M. endofungorum	Soil	South Africa	2	SRR9029401
NRRL 5554	R. microsporus		M. endofungorum	Soil	South Africa	4	SRR9029403
NRRL 5558	R. microsporus		M. rhizoxinica	Corn	USA	2	SRR9720186
NRRL 62023	R. oryzae		M. rhizoxinica	Corn	Nebraska, USA	3	SRR9720300
NRRL 66675	R. americanus		M. endofungorum	Rabbit dung	Pakistan	3	SRR8485126
NRRL A- 11791	Rhizopus sp.		M. rhizoxinica	NA	NA	8	SRR9720220
NRRL A- 11376	R. oryzae		M. endofungorum	NA	NA	4	SRR9720377

Continued

Table 1. Continued

NRRL A- 21579	R. oryzae	M. endofungorum	NA	NA	1	SRR12354412
NRRL A- 26124	R. microsporus var. oligosporus	Undetermined	NA	NA	1	SRR9720297
BC1034	Apophysomyces sp.	Mycetohabitans sp.	Millipede	Oklahoma, USA	2	SRR7686242
BC1015	Apophysomyces sp.	Mycetohabitans sp.	Millipede	Oklahoma, USA	1	SRR7686240

^{*}NRRL 5546 and NRRL 5547 were sequenced with PacBio in this study and are also in the ZygoLife database.

respective host isolates, or altogether different functions, are both seen with *Xanthomonas* TALEs [1]. In that same study [8], we discovered by Southern blot that Btl genes or gene fragments are present, in varying number and genomic context, in each of 14 examined strains of *Mycetohabitans* spp. from a global collection of *Rhizopus* accessions; strains were chosen for their availability from culture collections and whether they were previously reported to have endosymbionts, though some novel interactions were identified [19–21]. Recently, another Btl protein was characterized, Btl21-1 (also referred to as Bat1 and MTAL1) from Mrh strain B1 [7, 22]. The predicted DNA binding specificity of Btl21-1 is distinct from that of Btl19-13. A *btl21-1* mutant strain was less successful at colonization due to fungal septa development that prevented the bacteria from moving within hyphae [22], and other Btl proteins from B1 appear to promote successful re-establishment of the symbiosis needed for sporulation [23]. The Southern blot, the negative results of the Btl19-13 and Btl18-14 rescue experiment, and the sequence and functional differences between Btl19-13 and Btl21-1 support the hypothesis that Btl proteins play varied roles in symbioses between different *Mycetohabitans* and *Rhizopus* strains.

Importantly, the distribution and diversity of Btl proteins across *Mycetohabitans* is as yet unknown, as is their overall relationship as a group to *Xanthomonas* and *Ralstonia* TAL effectors. Determining Btl distribution and diversity across species and strains from multiple hosts or niches is key to understanding the ecological roles and evolution of these proteins. The limited availability of known *Rhizopus* accessions with *Mycetohabitans* symbionts and genomic resources for those accessions has been a barrier to better understanding how Btl proteins and other symbiotic factors are used. So far, only three *Mycetohabitans* genomes have been sequenced deeply enough to fully assemble the *btl* genes [24, 25]. The repetitive nature of TAL effector-family genes makes them a challenge with low coverage or short-read sequencing. Using long-reads, we sequenced and assembled seven new *Mycetohabitans* genomes and re-sequenced two of the three previously sequenced genomes, and examined the encoded Btl protein sequences to determine their distribution, diversity and evolutionary relationships. To expand this analysis, we also mined deep, short-read holobiont contig assemblies for a large collection of fungal isolates from the ZygoLife project (Joint Genome Sequencing Project doi: 10.46936/10.25585/60001062) [18, 26–28]. Here we report the findings of this two-part approach, 46 Btl proteins and 48 *btl* gene fragments from Mrh and Mef genomes, revealing widespread distribution and substantial variation in sequence and predicted DNA binding specificity, as well as species-specific patterns of conservation.

METHODS

DNA extraction

Mycetohabitans spp. were extracted from their fungal hosts by crushing fungal tissue from nonsporulating cultures with a scalpel on an empty Petri dish, removing hyphal fragments by syringe filtration and plating on Lysogeny broth (LB) amended with 1% glycerol [14]. Plates were incubated at 28°C until colonies formed (approximately 1 week), which were then inoculated to liquid LB and grown with shaking at 28°C until turbid (2–5 days). Genomic DNA was prepared from 3 ml of turbid liquid culture with the MasterPure Gram Positive DNA Purification Kit (Lucigen). DNA was quantified by using a Nanodrop device (Thermo Scientific) and integrity was assessed by agarose gel electrophoresis.

Genome sequencing and assembly

Libraries were prepared and sequencing was carried out using PacBio long-read technology (Pacific Biosciences), at the Mount Sinai Icahn School of Medicine Genomic Core Facility. Strains B13 and B14 were sequenced on an RSII machine using one SMRT cell each as described [29] and were assembled with HGAP and polished with Quiver in the resequencing pipeline in SMRT Analysis v2.2.0. The seven additional strains were multiplexed and run in the same SMRT cell on a Sequel I machine. Initial assembly was done by the Core Facility using SMRTLink v.8(8.0.080529) and the full set of reads. This resulted in closed assemblies with circular contigs for bacterial isolates from accessions CBS112285, CBS111563 and NRRL5549, but several linear contigs each for the others. For the incomplete assemblies, reads were downsampled to 50% using the SMRT Link command line tool bamsieve (using --percentage 50 and --blacklist options) to create two mutually exclusive, downsampled datasets, and each 50% was separately

assembled using SMRTLink v.9(9.0.0.92188). This method yielded closed assemblies for genomes of the bacterial isolates from NRRL 5547, NRRL 5560 and ATCC 52812. All closed genomes were run once through the SMRTLink v.9 resequencing pipeline with the complete read set and resulting coverage graphs were inspected for anomalies. Output circular contigs were rotated to the putative replicative origin in SnapGene (www.snapgene.com): chromosomes were rotated to a low GC area just upstream of *dnaA*, the larger megaplasmids to a low GC area immediately upstream of *repA* and smaller megaplasmids to just upstream of a predicted *repO*. Final assemblies were analysed for completeness with BUSCO [30]. Clusters of Orthologous Genes (COGs) analysis of the chromosomes and megaplasmids for the complete genomes was done with GenoVi [31] and visualized in R.

Genomic analyses

The reference sequence alignment-based phylogeny builder (REALPHY) pipeline [32] was used to infer the phylogenetic relationships of the ten *Mycetohabitans* genomes using *Burkholderia pseudomallei* strain K96243 as an outgroup (NCBI: BX571965-BX571966). Anvi'o v.8 [33] was used to carry out the generation and visualization of a pangenome following the workflow outlined previously [34]. Using GenBank files as input, anvi-script-process-genbank was used to generate FASTA as well as NCBI_PGAP-generated gene calls and annotations [35]. Specifically, anvi-gen-contigs-database was run on the FASTA files for each genome with specifying --external-gene-calls as NCBI_PGAP. Functions and gene calls were included in contigs databases using anvi-import-functions. Further, hidden Markov model (HMM) and single-copy core gene (SCG) taxonomy were run on the contigs databases for further annotation [36, 37]. The pangenome was assembled with the command anvi-pan-genome, utilizing a genomes storage database generated by the command anvi-gen-genomes-storage with the flag for --external-genomes [33].

Additional contig-level genomes from NCBI (B8, GCF_021991875.1; B6, GCF_021991795.1; B2, GCF_021733735.1) and from raw fungal data (see below) were retrieved and run through REALPHY. Average nucleotide identity (ANI) analysis on all genomes from was done using the enveomics collection ANI Matrix tool [38]. Phylogenetic trees were visualized and prepared for publication using FigTree (http://tree.bio.ed.ac.uk/software/figtree/).

Identification of novel Mycetohabitans hosts using raw fungal genome sequence data

Raw fungal genomic sequencing data associated with individual strains were acquired from the National Center for Biotechnology's Short Read Archive (NCBI SRA) for Mucoromycota and Zoopagomycota (i.e. Zygomycota *s.l.*) fungi. These data were processed into FASTQ files with the SRA toolkit (https://github.com/ncbi/sra-tools). Contigs were assembled using SPAdes v.3.15.2 [39] using the pipelines developed for the project (https://github.com/zygolife/DDD). Briefly, data were filtered, assembled with SPAdes v.3.15.2 with --meta option and also run in plasmid mode (--meta and -plasmid options) to detect circularized plasmids from metagenomes. These produced assemblies were binned with Autometa v.1.0.2 [40] to produce metagenome-assembled bacterial genomes (MAGs). The putative taxonomy of MAGs were assigned by GTDB-Tk [41] and MAGs were extracted and included in ANI and phylogenetic analysis described above.

Identification and annotation of Btl genes

In the complete genomes, *btl* gene sequences were identified by querying with the DNA sequence of *btl19-13* using BLAST. The hits were inspected and ORFs annotated manually. For the MAGs, a TBLASTN search of non-binned (fungal and bacterial) contigs was performed using the amino acid sequences of Btl19-13 and Btl21-1 as queries, and all contigs with hits to either protein were extracted. A manually curated list of Btl proteins from all available genomes was used to create a reference database in Geneious Prime (v.2022.1.1, Biomatters), and the 'Annotation and Prediction' function was used to annotate the contigs (settings: 50% similarity, 'best match' option). Annotation of the contigs with Prokka (v.1.14.5) [42] and extraction of features containing 'effector' from the annotations, and additional TBLASTN analyses of the contigs using *Xanthomonas* (PthXo1, WP_041182630.1) and *Ralstonia* (RipTALIV-2, LN874063.1) TAL effector sequences as queries were used to validate the final annotations. The promoter region (~300 bp upstream) of each *btl* gene was inspected for any match to the *hrp*_{II} box consensus sequence, TTCG-N16-TTCG [15, 43, 44]. Type III secretion signals were predicted using EffectiveDB [45].

Phylogenetic analyses of Btl protein sequences

Btl protein sequences from the nine PacBio bacterial assemblies and the reference *M. rhizoxinica* genomes were analysed using the QueTAL webtools DisTAL (with RVDs removed option) and FuncTAL [46].

Extracted *btl* genes and fragments were translated to amino acid sequences and parsed with an R script called Btl RVD Finder (Github: cartercharlotte/BtlDiversity) to identify key motifs and repeats based on regions conserved among the initial Btl protein set. Btl proteins were named using the number of repeats, starting with the conserved YD repeat as 1, and the bacterial strain number (or abbreviated host strain number when the bacteria are unnamed), following Carter *et al.* [8]. Proteins lacking a methionine start, a C-terminal NLS, or both the conserved N-terminal pseudorepeat and the YD first repeat were labelled as incomplete. The sequence context of fragments was ascertained manually. Fragments were named in the same way as intact proteins except substituting a unique letter for the (unknown) number of repeats (e.g. BtlA-12 from host strain '12' abbreviated from ATCC 52812).

All amino acid sequences were aligned with MUSCLE and used for a maximum likelihood phylogeny with 1000 bootstraps in MEGAX [47]. The resulting output was converted to a Newick file and visualized using RStudio with the script Btl Phylogeny (Github: cartercharlotte/BtlDiversity). Additional information on species name and data origin was added manually to the table of Btl sequences and features (generated by Btl RVD Finder) and used to annotate the tree. Separately, to eliminate the confounding lack of node support resulting from the presence of the incomplete proteins, a subset of complete Btl protein sequences was similarly processed. Sequence logos were generated [48] from repeat sequences extracted from the final list of all intact Btl proteins.

All TAL effector family comparison

Representative proteins from all TAL-like families described to date were collected and added to a subset of Btl proteins. From *Ralstonia solanacearum*, four RipTALs were chosen to represent the four designated RipTAL classes [49]: RipTALIV (CTQ57112.1), RipTALII (CTQ57109.1), RipTALIII (CTQ57110.1) and Brg11(Q8XYE3.1). Several *Xanthomonas* TALEs were chosen to represent the known diversity within that family: *Xanthomonas euvesicatoria* AvrBs3 (P14727.2) and AvrBs4 (CAA48680.1), *Xanthomonas oryzae* pv. oryzae PthXo1 (WP_041182630.1) and AvrXa10 (Q56830.1), *Xanthomonas oryzae* pv. oryzicola Tal5 (AJQ86298.1) and Tal8 (AJQ88017.1), *Xanthomonas citri* PthA (ACZ62653.1), and *Xanthomonas translucens* pv. undulosa Tal1 (WP_108084612.1), Tal2 (WP_108084616.1) and Tal4a (WP_108084830.1). To represent the family of TAL-like proteins found in the marine metagenome, MOrTL1 (ECG96326.1) and MOrTL2 (EBN1909) [6] were included. Also included were three TAL-like protein fragments identified in a *Mycoplasma* genome [50]. The full set of 25 protein sequences was aligned with Muscle and a maximum likelihood phylogeny was reconstructed using all sites with 1000 bootstraps in MEGAX [47]. The same protein set was analysed with DisTAL [46], though not all proteins were recognized by that software as it was built for *Xanthomonas* TALEs.

RESULTS

De novo sequencing of seven and re-sequencing of two Mycetohabitans genomes using long-read technology

Previously, we identified seven *btl* genes in the genomes of strains B1, B13 and B14, yielding a very limited data set, but by Southern blot probed with *btl19-13* we found hybridizing bands in each of 11 other strains, with dissimilar banding pattens suggesting high sequence diversity [8]. To examine this diversity, we attempted to purify high-quality DNA for long-read whole genome sequencing on the PacBio Sequel I platform. We succeeded for seven strains, including that of the type strain of Mef, B5, which was previously only available as a contig-level assembly that was missing *btl* genes [10]. Of the seven, we obtained complete assemblies for B5 and five others, with BUSCO genome completeness scores [30] ranging from 99.1 to 99.4%, and a contig-level assembly for one, B46, with a BUSCO score of 98.2% (Table 2). B46 was underrepresented in our total read set, probably because it was underrepresented in the multiplexed library.

In our previous study of *btl19-13*, we used complete genome assemblies of B13 and B14 that we generated by sequencing on the PacBio RSII platform, and an assembly for B1 that was publicly available (GenBank accession PRJEA51915). During that study, the Joint Genome Institute (JGI) deposited assemblies for B13 and B14 (GenBank accessions PRJNA303198 and PRJNA303197); those yielded the same *btl* gene content as our assemblies, so we cited them in our paper [8]. We have since deposited our B13 and B14 assemblies (Table 2). While we were completing our analyses for the present study, eight Illumina-based *Mycetohabitans* genome assemblies were made publicly available on NCBI, including some for strains that we sequenced; however, those assemblies are contig level, not closed [51].

Our six complete assemblies reveal chromosomes ranging from 2592175 to 2776699 bp, and one or two additional replicons each. In all six assemblies there is a megaplasmid of 716–857 kb as there is in B1, B13 and B14. These megaplasmids are similar to pBRH01 of the reference strain *M. rhizoxinica* B1, with the megaplasmids of B12, B47 and B49 being very similar (99% identity over 91–94% of pBRH01, using megablast), and those of B3, B5 and B60 less similar (87–93% identity over 69–72% of the sequence). Strains B12 and B60 both have additional large plasmids that have some similarity to the plasmid of reference strain B1, pBRH02, with that of B12 being more similar (99% identity over 75% of the sequence) than that of B60 (94% identity over 50% of the sequence). A basic analysis of the COGs in the megaplasmid and plasmid across strains showed enrichment of unclassified genes, defence mechanisms, mobilome and repair compared to the chromosome (Fig. S1, available in the online version of this article). The plasmid has very few genes related to transport and metabolism. The larger megaplasmid is conserved across strains, whereas the smaller plasmid, while it is not restricted to one species, is not present in all strains.

Genomic relationships and species of the sequenced strains

To determine the diversity of strains and the species represented by the nine genome assemblies we generated, we performed whole genome phylogenetic analysis [32] including in the analysis the Mrh type strain B1 [25] (Fig. 1a). Among our newly sequenced genomes is the type strain for Mef, B5. Except for B46, all strains have at least 94.6% ANI to one or the other of these type strains, meeting the threshold conventionally used as a bacterial species boundary of 94–96% depending on genus [52, 53]. The strains indeed separated clearly into two clades with B46 as an outlier, though B1 shared higher ANI with strains designated as Mrh than B5 did with strains designated as Mef. From this limited number of strains, there was no strong association of species with

Table 2. Mycetohabitans spp. genome assemblies generated in this study*

Strain	Genome size (bp)	Coverage	Contigs	Chromosome (bp)	Plasmid 1 (bp) Plasmid 2 (bp)		Protein coding genes	tRNAs	%GC	NCBI accessions	Fungal host accessions
Mycetohabitans endofungorum B3	3541851	463×	2	2686710	855141	N/A	2771	47	9.09	CP062180-CP062181	CBS 111563
Mycetohabitans endofungorum B5	3355449	402×	7	2592175	763274	N/A	2676	47	61.2	CP062178-CP062179	CBS 112285
Mycetohabitans endofungorum B60	3610257	434×	ю	2736534	716486	157237	2869	48	8.09	CP062168-CP062170	NRRL 5560
Mycetohabitans rhizoxinica B47	3471072	770×	7	2646852	824220	N/A	2805	47	61.1	CP062173-CP062174	NRRL 5547
Mycetohabitans rhizoxinica B12	3802698	734×	ю	2776699	857052	168947	2984	47	60.7	CP062175-CP062177	ATCC 52812; CBS 262.28
Mycetohabitans rhizoxinica B49	3533946	925×	7	2710688	823258	N/A	2815	47	6.09	CP062171-CP062172	ATCC 46352; NRRL 5549
Mycetohabitans sp. B46	3293065	139×	21	ND	ND	ND	2668	51	61.0	JADBGL000000000	NRRL 5546
Mycetohabitans endofungorum B13	3550038	136×	2	2760408	789630	N/A	2747	47	9.09	CP132744-CP132745	ATCC 52813
Mycetohabitans endofungorum B14	3643928	172×	æ	2674176	768340	201412	2932	47	60.6	CP132741-CP132743	ATCC 52814

*ND, Not determined; N/A, no second plasmid present.

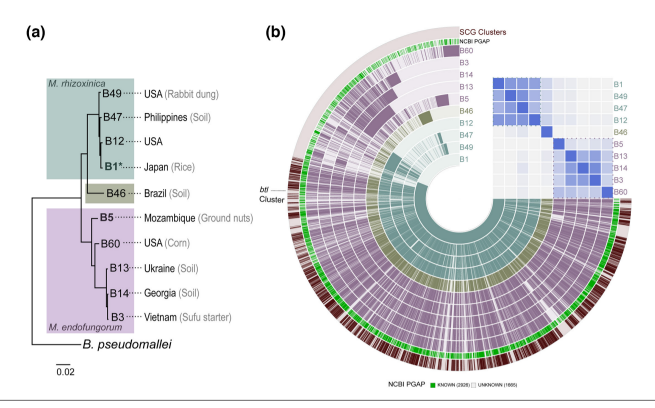


Fig. 1. Genomic relationships, species and origins of the sequenced strains. (a) Output from the reference sequence alignment-based phylogeny builder (REALPHY) for the nine strains sequenced in this study and the *M. rhizoxinica* type strain B1 (denoted by *). B5 is the type strain for *M. endofungorum*. The geographical and substrate origin of the host isolate for each strain is listed when known. *Burkholderia pseudomallei* strain K96243 was used as an outgroup. Scale bar indicates substitutions per site. (b) Pangenome of the ten strains with indicated average nucleotide identity (ANI) heatmap. The blue, green and purple layers represent *M. rhizoxinica, Mycetohabitans* sp. B46 and *M. endofungorum* genomes respectively, where gene clusters are arranged based on synteny with B1. The outermost dark red layer highlights single-copy core gene (SCG) clusters found in all ten genomes. In the heatmap, blue dotted lines indicate clusters of strains with 94.6% or greater ANI to one another.

location or substrate from which the host was isolated: Mrh strains appear in fungal isolates from Asia and North America, and Mef strains in isolates from Asia, North America, Africa and Europe, and both species appear in fungi isolated from soil or plant material, which represent the majority.

We generated a pangenome for the ten *Mycetohabitans* genomes using anvi'o [33]. We used B1 synteny as reference for displaying alignments and included ANI analysis (Fig. 1b). The pangenome contained 29051 genes within 4591 gene clusters. Single-copy core genes (SCGs) were identified in a total of 1731 gene clusters within the pangenome; all 22 SCGs were found to have 100% annotation based on the Genome Taxonomy Database (GTDB) [36]. We identified 24 putative *btl* genes within the pangenome. These were distributed across all strains except B5 and included a maximum of four paralogues, found in B1, B12 and B14. These genes were grouped in the same gene cluster, which had a combined (geometric and functional) homogeneity index of 0.73, indicating a high degree of similarity across strains in the residues as well as in the distribution of gaps across the multiple sequence alignment.

Btl protein diversity across the ten strains

As the goal of our sequencing was specifically to interrogate btl genes, we extracted the identified btl gene sequences from each of our seven new genome assemblies to compare to the seven known btl sequences from the other three genomes [8]. Btl gene sequences were found on the chromosome, the smaller plasmid, the large one or a combination (Table S1). Not all encode full-length proteins. Though not identified in our pangenome analysis, a single sequence with homology to btl genes could be found in the (complete) assembly for Mef type strain B5 (also called HKI 0456; fungal accession CBS 112285) but it only encodes two repeats, and they are out of frame with each other. The contig-level genome [10] assembled from short-read Illumina data for this strain (NZ_PRDW00000000.1) also contains this btl gene and none other, but there is a separate deposited sequence (MN891945) attributed to this strain encoding a 20-repeat Btl protein that is not present in our B5 assembly. Similarly, strain B46 from the fungal accession NRRL 5546 contains only a small (479 bp) btl gene fragment. The remaining five strains have 11 intact btl genes and six btl gene fragments between them (Table S1). Most of the gene fragments are short, containing one to two repeats at most and many stop codons or frameshifts, and they are often interrupted by transposable elements.

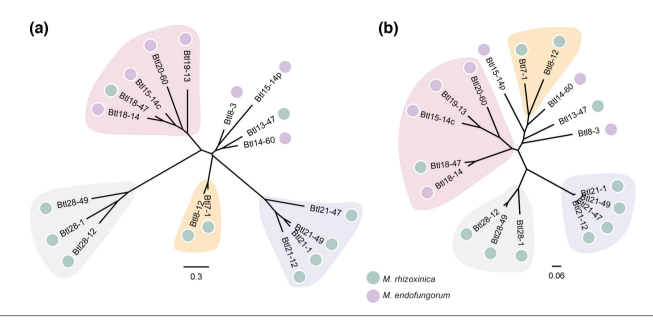


Fig. 2. Btl proteins do not exclusively cluster by species. Unrooted trees depicting (a) DisTAL and (b) FuncTAL phylogenetic analyses based on the amino acid sequence with RVDs removed and RVDs only, respectively, for the putative Btl proteins encoded in ten *Mycetohabitans* genomes. More information on these proteins can be found in Table S1. Clades present (with the same members) in both trees are highlighted by background shapes of the same colour. Scale bars correspond to DisTAL and FuncTAL scores.

Previously identified full-length btl genes have a sequence within ~200 bp upstream of the start codon that matches the hrp_{II} box binding element, TTCG-N16-TTCG, for the T3SS regulator HrpB found in R. solanacearum and Burkholderia pseudomallei [8, 43, 44]. The intact btl genes in our newly sequenced strains likewise harbour the element at this location (Table S2). Despite this evidence for co-regulation with the T3SS, most of the Btl proteins do not have computationally predicted N-terminal signals for T3S (Table S2). Btl19-13 is among these, yet it has been shown to be T3 secreted [8]. Thus, the prediction is lacking, and based on N-terminal sequence similarity to Btl19-13, it seems likely that all the Btl proteins in fact transit the T3 pathway. A possible exception is the eight-repeat Btl protein encoded by a chromosomal gene in strain B60, from fungal accession NRRL 5560 (Table S2). This gene lacks the hrp_{II} element due to an interruption of the putative promoter region and 5' end of the btl ORF by a transposase ORF, which may eliminate expression and is likely to be in the process of pseudogenization.

To understand the relationships among the different intact Btl protein sequences, we used the QueTAL webtools DisTAL and FuncTAL for phylogenetic analysis [46]. DisTAL removes the RVDs, i.e. the determinants of binding specificity, and aligns based on the remaining repeat 'backbone' sequences, while FuncTAL compares only the RVD sequences of the proteins. This allows for differentiation between the potentially rapid evolution of the DNA targeting function versus the presumably slower evolution of the backbone. The Btl proteins group similarly in these two analyses (Fig. 2). There are differences in the relationships of the clades to each other, and certain proteins, like Btl8-12 and Btl7-1, appear in different relative locations in the two trees. However, the similar clustering overall suggests that the RVDs and the rest of the repeats are evolving in concert. While certain clades comprise only proteins from one *Mycetohabitans* species, such as the set of four 21-repeat Btl proteins present in four Mrh strains, one clade contains sequences from both species. In this small sample set, the degree of conservation varies depending both on the protein and the species.

Btl proteins encoded in fungal hologenomic sequences

To explore Btl protein diversity more fully, we probed fungal genome sequencing data from the ZygoLife Project which aimed to understand taxonomic diversity and evolutionary history of the Mucoromycota and Zoopagomycota fungi [18, 27, 28, 54–57]. An aspect of this project involved low- to medium-coverage genome sequencing of NRRL collection isolates which we utilized to search for *Mycetohabitans* genomic evidence and *btl* genes contained within. To identify endosymbiont hosts and probe endosymbiont diversity, the raw genome sequence data were assembled into metagenome assemblies and binned into bacterial MAGs. Bacterial MAGs within the data set were associated with a species designation using the GTDB Tool Kit [41], but we searched for *btl* sequences in all contigs, not just bacterial assemblies, and indeed identified *btl* gene fragments in two accessions, NRRL 3373 and NRRL A-26124, that did not have sufficient bacterial reads for identification of the symbiont genome. All accessions with both *btl* sequences and bacterial MAGs corresponded to *Rhizopus* spp. or *Apophysomyces* spp. accessions and the bacteria

were putatively identified as *Mycetohabitans* spp. (Table 1). New *btl* genes were named using the last two numbers of the fungal accession ID, or, if not unique, the last three.

In addition to the ZygoLife hologenomes, we also searched GenBank for matches to Btl proteins and found deposited fragments from three *Mycetohabitans* strains, HKI 0404/B8, HKI 0513/B6 and HKI 0512/B2, which are symbionts of fungal isolates CBS 308.87, CBS 261.28 and ATCC 20577, respectively [23]. Subsequently, the contig-level genomes of these strains were also deposited (see genome phylogenetics below). Combined, the total number of *btl* genes or fragments identified increased to 94 (Table 1).

Hologenomic sequences vs. long-read bacterial genome assemblies

The Mrh strain associated with accession NRRL 5547, hereafter referred to as strain B47, provides a useful case study for assessing how accurately repetitive genes of interest could be extracted from ZygoLife short-read MAGs versus our long-read sequenced genomes. We identified three intact *btl* genes in the complete B47 genome. Five different contigs from the B47 MAG contain *btl* homologous sequences: one contains the full-length *btl21-47*, and the four others contain the 5' or 3' end of *btl18-47* or *btl13-47*, with these fragments falling at the end or beginning of the contig. Highlighting a limitation of short-read sequencing for assembling repetitive sequences like *btl* genes, for each *btl* gene in the B47 MAG, a few repeats within the repeat region captured are missing. However, the translated sequences of the gene fragments match perfectly with the intact Btl18-47 and Btl13-47 encoded in the long-read assemblies, minus the missing central repeats. B46 (from NRRL 5546) was also present in both data sets, but we did not retrieve either of its *btl* fragments in our search of the ZygoLife MAGs.

Btl protein repeat variation

The function of TAL-family proteins is determined by their DNA-binding specificity and requires nuclear localization, so the repeat types present and variation in repeats that confer specificity, as well as variation in any nuclear localization signals, is of interest for inferring functional variation of Btl proteins within and across strains [7–9]. Comparison across all of the sequences revealed four mostly conserved types of repeats in Btl proteins (Fig. S2): (1) a 1 as shorter pseudo-repeat harbouring the RVD HS, HY, NY or QY; (2) a first, sequence-divergent repeat that has the RVD CD, HD or YD; (3) the central, standard repeats; and (4) an again sequence divergent final repeat usually harbouring the RVD N* (where the asterisk represents a missing residue). This finding agrees with the initial observations of two N- and one C-terminal cryptic repeats in a set of just three Btl proteins analysed by de Lange *et al.* [7], who found that the 'first repeat' does not confer specificity. Comparison across the proteins also revealed variation in the nuclear localization signal following the repeats, either RIRK or QIRK. The RIRK motif is necessary for nuclear localization and changes to these residues or neighbouring ones may impact localization [8].

Btl protein diversity in M. rhizoxinica relative to M. endofungorum strains

Using the *btl* gene sequences identified in our assemblies, the ZygoLife MAGs and NCBI, we re-assessed Btl protein diversity, as well as distribution. We conducted a maximum likelihood phylogenetic analysis and excluded incomplete *btl* gene sequences, which may represent assembly gaps or actual pseudogenes, leaving a total of 46 sequences (Fig. 3; analysis with all Btl proteins in Fig. S3). Six clades were evident based on tree topology as well as differentiation by motifs, and these were named I–VI. In addition to the whole protein analysis, we extracted and compared just the RVDs of the intact Btl proteins. For automated extraction, we created our own tool (see Methods), which better handles Btl protein repeats than tools designed for *Xanthomonas* TAL effectors. We grouped Btl proteins by clades from Fig. 3 and aligned the RVD sequences to determine if potential DNA targets are conserved within each clade (Fig. 4). Where there were multiple options for the alignment of an RVD due to a mismatch, deletion or insertion, we used the backbone repeat sequence to guide the alignment; however, aligned RVDs are not necessarily part of identical repeats.

RVD sequences, repeat number and key motifs are highly conserved within Clades I, IV and V, with proteins often exhibiting only RVD polymorphism and at only one or two repeats. In contrast, Clade II, containing the protein we characterized previously, Btl19-13 [8], does not have high conservation of number of repeats per protein, ranging from eight to 20, and has relatively high protein to protein polymorphism in the RVDs. Clade III is similarly variable, and these proteins are on average shorter, containing 9–15 repeats. Clade V consists of sequences from the ZygoLife MAGs only and includes the only proteins with a 'CD' RVD in the first repeat. The proteins in Clade VI have only 7–11 repeats, the shorter ones apparently having lost middle repeats. All members have variants of all key motifs that are unique to this group. Clade VI includes Btl7-1 (also BAT3), the single Btl protein from B1 that does not appear to bind DNA [6], potentially due to these variations in sequence, or the small number of repeats, compared to other clades, highlighting the functional diversity brought about by sequence variation.

To examine distribution, we carried out whole genome-based phylogenetic analysis on the larger set of strains and mapped the Btl protein content (by clade) onto the resulting tree (Fig. 5a). We further examined strain relationships by ANI (Fig. 5b). Strains that did not have bacterial assemblies (NRRL 3373 and NRRL A-26124) or were insufficiently complete (BC1015 and BC1034) were excluded from these analyses. Within this dataset, Mrh strains all have *btl* genes and have more per strain on average than Mef strains. Furthermore, Mrh strains always have a Btl protein from Clade I or VI, and rarely Clades II, III and V. Both the

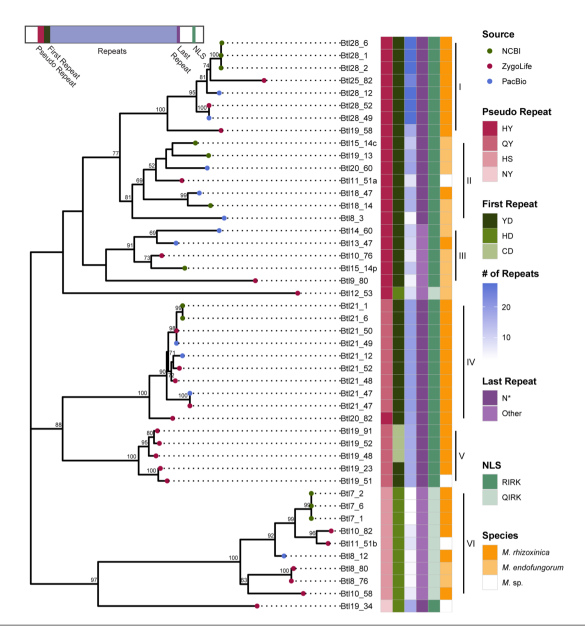


Fig. 3. Maximum likelihood phylogenetic analysis of 46 intact Btl protein sequences. Bootstrap values (as a percentage of 1000) are shown at the nodes for those with >50% support. As shown in the key at right, coloured circles at branch ends indicate the source of the data for each sequence, and coloured boxes detail *Mycetohabitans* species, number of repeats, and presence or absence of key protein motifs, including the nuclear localization signal (NLS), shown using the single letter amino acid code. An asterisk indicates a missing amino acid where one would be expected in a typical conserved repeat, i.e. 'N*. A diagram of a Btl protein with coloured, corresponding motif locations is in the top left.

phylogenetic analysis and the ANIs suggest that Mef is composed of two subspecies, though additional resolution is needed. In one, which contains none of the PacBio-sequenced strains, there are many *btl* gene fragments unassigned to a clade. In the other, which consists mostly of PacBio-sequenced strains, a Btl protein from Clade II is present in all strains, but *btl* gene content varies otherwise. In only B14 are two Btl proteins from the same clade observed. Two strains reside outside of both named *Mycetohabitans* spp. The assemblies for these and three strains in the Mef group have only incompletely assembled *btl* genes or pseudogenized *btl* genes or fragments (e.g. B46 and B5).

Btl proteins form a distinct but variable clade of TAL effectors

Next, we explored the evolutionary relationship of Btl proteins to TAL effectors and TAL effector-like proteins by generating a maximum likelihood tree based on the amino acid sequences. We chose (1) a representative set of full-length Btl proteins based on Fig. 3; (2) a set of TALEs from *Xanthomonas oryzae* pv. oryzae (Xoo), *Xanthomonas oryzae* pv. oryzicola (Xoc), *Xanthomonas*

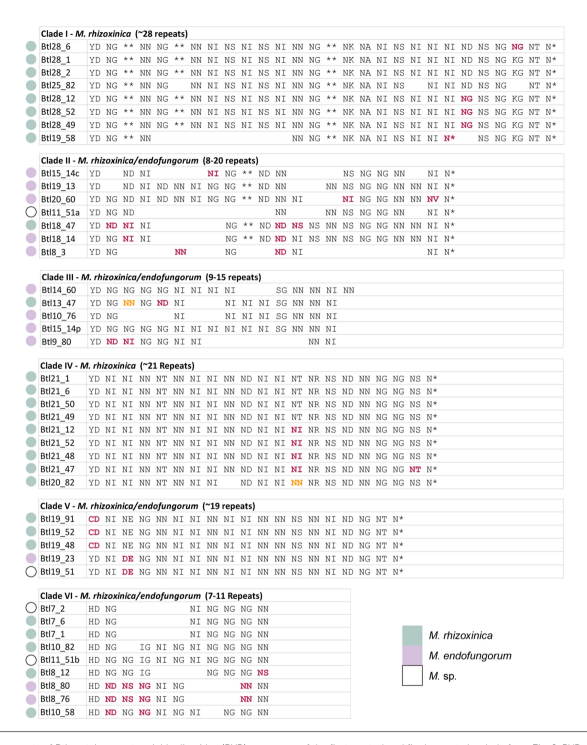


Fig. 4. Alignments of Btl protein repeat variable diresidue (RVD) sequences of the first, central and final repeats by clade from Fig. 3. RVDs matching the consensus are in black font. RVDs that vary from the consensus are in bold pink font, or bold orange to indicate a second variant at that position. An asterisk (*) represents a missing amino acid at the respective position of the RVD. Coloured circles indicate species, following the key at bottom right.

euvesicatoria (Xe), Xanthomonas citri (Xc) and Xanthomonas translucens pv. undulosa (Xtu); (3) RipTALs from Ralstonia solanacearum (Rs); (4) all available MorTL sequences [6]; and (5) Mycoplasma E-protein sequence fragments, which were suggested to represent distant members of the TAL effector protein family [50]. As shown in Fig. 6(a), Btl proteins cluster distinctly from TAL effectors and RipTALs, which form two closely related clades. One of the MorTL sequences groups more closely with Btl proteins than with the TALE and RipTAL groups. The other appears to have diverged earlier, from a common ancestor of Btls, TALEs, RipTALs and the first MorTL. The Mycoplasma sequences cluster on a separate branch. Finally, to further probe evolutionary

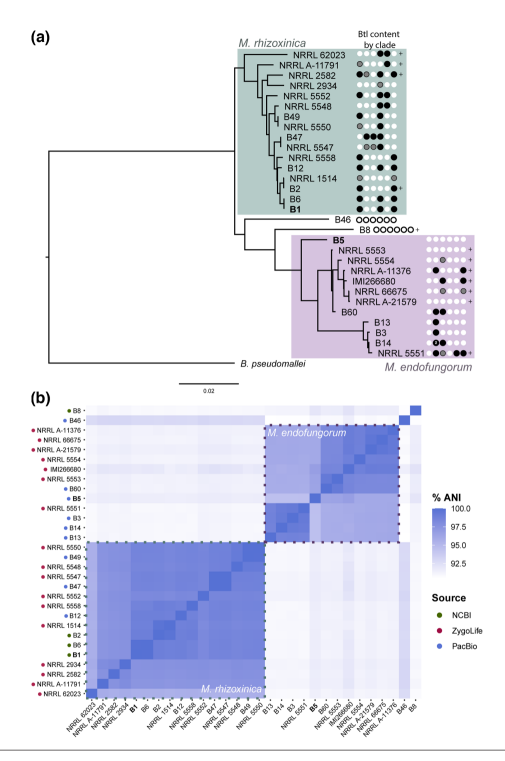


Fig. 5. Genomic relationships, species and *btl* gene content of *Mycetohabitans* strains examined in this study. (a) Output from the reference sequence alignment-based phylogeny builder (REALPHY) for the nine strains we sequenced, the metagenome bacterial assemblies and publicly available *Mycetohabitans* spp. genomes. Taxa are labelled by bacterial strain name (B#), or fungal accession number where a bacterial strain has not been named. Circles to the right of each taxon represent the *btl* gene content by clade (I–VI from left to right): white represents no gene, black represents one gene from that clade, or two in the case of strain B14, and grey represents an incompletely assembled gene that closely match that clade. Plus signs (+) indicate additional incompletely assembled fragments that were not assigned to a clade. *Burkholderia pseudomallei* strain K96243 was used as an outgroup. Scale bar indicates substitutions per site. (b) Average nucleotide identity analysis of the ten strains. Dotted lines indicate clusters of strains with 95% or greater ANI to one another, indicative of species. The source of a strain's sequence is indicated by coloured dots along the left axis. B1 is the type strain for *M. rhizoxinica* and B5 is the type strain for *M. endofungorum*.

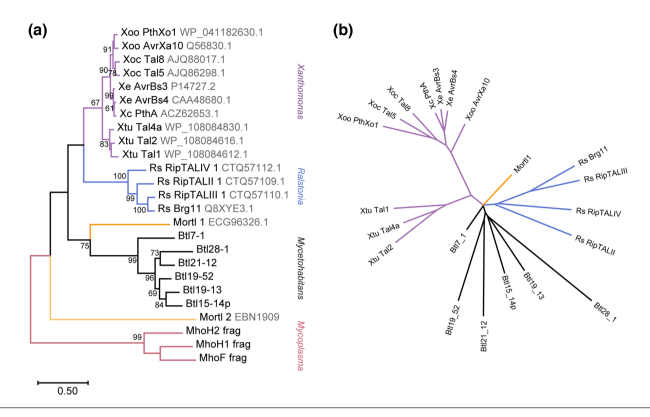


Fig. 6. Relationship of Btl proteins to TALEs, RipTALs and other TAL-like proteins. (a) Maximum likelihood tree based on amino acid sequences of the indicated proteins. Nodes with bootstrap values >50% (out of 1000 replicates) are labelled. Scale bar indicates substitutions per site. Where available, identifiers from UniProt or GenBank are given in grey font next to the protein names. Branches are coloured and clades are labelled based on the genus from which the sequence was taken, except for the marine metagenome sequences, which are in orange. (b) DisTAL analysis of the same group of protein sequences presented as an unrooted tree. Line colouring matches that in (a). Proteins too divergent from Xanthomonas TAL effectors to be parsed correctly by DisTAL are absent from the tree.

relationships, we analysed the sequences using DisTAL. The more distant of the two MorTL sequences and the *Mycoplasma* sequences were too far diverged from the *Xanthomonas* TALEs to be processed by the algorithm correctly and were therefore not in the resulting tree. However, an interesting pattern emerged (Fig. 6b): the individual Btl protein branch lengths are on average longer than those of the other proteins in the tree. This difference reflects a greater degeneracy of the repeat backbone sequences within each Btl protein relative to TALEs and RipTALs, hinting at the possibility that they have had more time to diverge or are not as subject as TALEs and RipTALEs to selection for rapid evolution via recombination.

DISCUSSION

In this study, we sequenced seven new genomes of *Rhizopus*-associated *Mycetohabitans* spp., six completely and one to a high-quality contig-level assembly. We are also making available our complete assemblies for the genomes of two Mef strains that were previously sequenced by the JGI. When assessing the diversity of the *Mycetohabitans* spp. genomes themselves, there were two clear clades aligning with the two named species, though the Mrh clade had higher ANI between members. All strains had one to five *btl* genes or fragments per genome, with most genomes having two or three intact *btl* genes, and some having only fragments. We expanded our analysis by identifying and retrieving *btl* sequences from fungal metagenome assemblies through the ZygoLife Project. Our ability to extract a large number of intact *btl* genes from the ZygoLife MAGs underscores the value of such large sequencing efforts. Though mining ZygoLife did not consistently capture all *btl* genes and fragments captured by long-read sequencing of the smaller set of strains, the metagenome analysis yielded both intact and clearly pseudogenized *btl* genes, as well as incompletely assembled *btl* gene fragments at the ends of contigs. This larger data set enabled analyses that yielded new insight into the conservation and diversification of Btl proteins across *Mycetohabitans* species. Additionally, we found that Mef strains split into two subclades with the type strain as an outlier, potentially representing subspecies or novel species; further investigation with more phenotypic data will be required to determine which, if either, is the case. The high Btl variability, evidence of pseudogeniziation and overall lack of conservation across closely related strains are indicative of these effectors undergoing rapid evolution, probably facilitated by their repetitive nature.

The number of intact btl genes retrieved from the MAGs was somewhat surprising, given the repetitive nature of these genes, the use of short-read sequencing technology in that project and the anticipated scarcity of bacterial DNA compared to fungal DNA within the library preparations. The short-read assemblies and low coverage also result in shorter contigs overall, in some cases yielding btl sequences on contigs with few or no other genes, and making it impossible to definitively attribute those sequences to specific bacterial species, or even to the symbiont rather than the fungal host itself. Indeed, anticipating poor yield and false negatives, we took an inclusive approach to mining: we did not filter metagenomes by symbiont sequence coverage, nor did we filter for *Rhizopus* spp. In total, we identified btl sequences in the metagenomes of 19 fungal accessions out of the more than 900 total accessions, of which 200 are Rhizopus spp. Interestingly, all accessions yielding btl sequences were Rhizopus spp., except two, both accessions of a different mucoralean genus, Apophysomyces, that has also been reported as an opportunistic human pathogen. All of the corresponding bacterial assemblies were identifiable as Mycetohabitans spp., except for two, which did not have sufficient bacterial contigs for species designation. Importantly, we did not experimentally verify absence of btl sequences from accessions for which the metagenome yielded no hits. Yet, we can deduce that some of these are in fact false negatives. Namely, three Rhizopus accessions with extracted bacterial genomes had no detected btl fragments, yet one of these is B46, which we know from our long-read sequencing assemblies has btl gene fragments, albeit short ones. Future, targeted long-read resequencing of some of the hologenomes or isolation and direct sequencing of the bacterial strains would be of interest to fully resolve the btl gene repertoires as well as the genomic context for these genes.

One pattern that emerged is a greater conservation of Btl protein content across Mrh than Mef isolates. Each Mrh strain in our analysis has a Btl protein from Clade I or IV, the most conserved Btl clades, with six of ten strains having one of each. Within these conserved clades the RVD sequences are often nearly identical. The observation by Richter *et al.* [22] that Btl21-1/MTAL1, a member of Clade IV, contributes to the establishment of symbiosis by Mrh strain B1 suggests that Mrh generally may rely on Btl proteins for successful interaction with the fungus. In contrast, the Mef genomes we examined overall have fewer *btl* genes and have no Btl proteins in these conserved clades, and one strain successfully colonizes its host even when its single, Clade II *btl* gene is knocked out [8]. All strains in one Mef subclade contain Btl proteins from Clade II that are still much more variable in repeat sequence than the Clade I and IV Btl proteins found in Mrh strains more distant from one another. Altogether, these observations suggest that in contrast to Mrh, Btl proteins play varied roles in Mef, non-essential to symbiosis. The apparent positive effect of Mef Btl19-13 on host membrane stress tolerance [8], however, hints non-exclusively at effects beneficial to the host that may maintain symbiosis over evolutionary time.

It is also possible that some Btl proteins in Mrh and Mef have the same or functionally equivalent host targets, but those targets are sequence-divergent across the Mef host isolates, and the Mef Btl proteins correspondingly adapted and diversified. As Btl proteins have many putative effector binding elements across their hosts genomes, experimental studies validating host targets will be critical for comparison of relevant sequences within host genomes. Or, the Btl protein content variability may be insignificant, simply an artefact of the still relatively small sample size. The Clade V Btl proteins only being from ZygoLife MAGs and some Btl proteins being outliers to the six clades indeed suggests that there are more Btl clades than were captured in this data set.

This study identified many novel Btl proteins, but only a few have been characterized; this is a major caveat to extrapolating Btl function among clade members. Ultimately, functional characterization of representative members of each of the clades will help clarify whether the pattern of Btl protein distribution and conservation we observed reflects a diversity of functions, adaptation to a diversity of host genotypes, or some combination, and will lead to a more nuanced understanding of the importance of Btl proteins in *Mycetohabitans–Rhizopus* symbioses. Additionally, though comparative genomics within *Mycetohabitans* spp. analogous to recent studies with endofungal *Mycoavidus* spp. [58] was outside the scope of this study given the limited number of high-quality assemblies, identifying and sequencing additional strains would enable a deeper investigation into the *Mycetohabitans* pangenome, shedding more light on the evolutionary trends within and between the species themselves that will inform hypotheses of effector evolution.

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Conflicts of interest

J.E.S. was a paid consultant for Zymergen, Sincarne and Michroma.

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