What is in *Umbilicaria pustulata*? A metagenomic approach to reconstruct the holo-genome of a lichen

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Keywords: Metagenome assembly, SPAdes, Sequencing error, Symbiosis, Chlorophyta, Gene loss, organellar ploidy levels, Microbiome
Abstract

Lichens are valuable models in symbiosis research and promising sources of biosynthetic genes for biotechnological applications. Most lichenized fungi grow slowly, resist aposymbiotic cultivation, and are poor candidates for experimentation. Obtaining contiguous, high quality genomes for such symbiotic communities is technically challenging. Here we present the first assembly of a lichen holo-genome from metagenomic whole genome shotgun data comprising both PacBio long reads and Illumina short reads. The nuclear genomes of the two primary components of the lichen symbiosis – the fungus *Umbilicaria pustulata* (33 Mbp) and the green alga *Trebouxia* sp. (53 Mbp) – were assembled at contiguities comparable to single-species assemblies. The analysis of the read coverage pattern revealed a relative abundance of fungal to algal nuclei of approximately 20:1. Gap-free, circular sequences for all organellar genomes were obtained. The bacterial community is dominated by *Acidobacteriaceae*, and encompasses strains closely related to bacteria isolated from other lichens. Gene set analyses showed no evidence of horizontal gene transfer from algae or bacteria into the fungal genome. Our data suggest a lineage-specific loss of a putative gibberellin-20-oxidase in the fungus, a gene fusion in the fungal mitochondrion, and a relocation of an algal chloroplast gene to the algal nucleus. Major technical obstacles during reconstruction of the holo-genome were coverage differences among individual genomes surpassing three orders of magnitude. Moreover, we show that GC-rich inverted repeats paired with non-random sequencing error in PacBio data can result in missing gene predictions. This likely poses a general problem for genome assemblies based on long reads.
Introduction

The lichen symbiosis comprises a lichen-forming fungus (mycobiont), and a photosynthetic partner (photobiont), which is typically a green alga or a cyanobacterium. A bacterial microbiome and additional third-party fungi can also be part of the lichen consortium (Grube, et al. 2015; Spribille, et al. 2016). The bacterial microbiome in particular may contribute to auxin and vitamin production, nitrogen fixation, and stress protection (Erlacher, et al. 2015; Grube, et al. 2015; Sigurbjornsdottir, et al. 2016). Lichenized fungi are well known for synthesizing diverse, bioactive natural products (reviewed in (Muggia and Grube 2018)), which has recently stimulated research into biosynthetic pathways and gene clusters of these fungi (Abdel-Hameed, et al. 2016; Armaleo, et al. 2011; Bertrand and Sorensen 2018; Calchera, et al. 2019; Wang, et al. 2018). The estimated 17,500-20,000 species of lichens (Kirk, et al. 2008) are distributed across nearly all ecosystems (Ahmadjian 1993). Some lichens thrive as pioneering organisms in ecological niches that are otherwise adverse to eukaryotic life (Hauck, et al. 2009; Kranner, et al. 2008). The capability to inhabit such a diverse set of habitats is tightly connected with the lichen symbiosis itself. The nutritionally self-sustaining system harbors internal autotrophic photobionts, which provide carbohydrates to all other members of the association. Furthermore, some mycobiont species switch between different sets of environmentally adapted photobionts, and can thus occupy broad ecological niches (Dal Grande et al. 2018, Rolshausen et al. 2018).

There is an increasing interest in genomic resources on lichens, because lichens are valuable models in symbiosis research (Grube, et al. 2015; Grube and Spribille 2012; Wang, et al. 2014), and promising sources of biosynthetic genes for biotechnological applications (see above). Most lichenized fungi grow slowly, resist aposymbiotic
cultivation, and are generally poor candidates for experimentation. Therefore, researchers increasingly use genomic data as sources of novel information on the lichen symbiosis (e.g. (Armaleo, et al. 2019)). Genome sequences of about 19 lichenized fungi and of two algal photobionts have been published to date (Table 1). Most genome sequences stem from lichens whose symbionts were grown in axenic culture. The few studies using metagenomic data to reconstruct the fungal genomes reported highly fragmented assemblies comprising more than 900 scaffolds (Allen, et al. 2018; Liu, et al. 2019; McDonald, et al. 2013; Meiser, et al. 2017). Some assemblies range in an expected total length (McDonald, et al. 2013; Meiser, et al. 2017), and achieve comparable BUSCO (Simao, et al. 2015) scores to assemblies derived from single-species cultures (Meiser, et al. 2017). However, the only two publicly available genome sequences of lichenized fungi that were assembled from metagenomics data, *Cetradonia linearis* and *Alectoria sarmentosa*, (Allen, et al. 2018; Liu, et al. 2019) have more than 20% BUSCO genes missing (Table 1). They are thus far from complete. Moreover, discontinuous assemblies are of limited use for functional genomics analyses, which rely on a comprehensive and accurate annotation of genes and even more so of gene clusters (Denton, et al. 2014; Dunne and Kelly 2017). Attempts to assemble the entire holo-genome of a lichen have not been reported, thus far. Also, a genome assembly strategy based on long read sequencing technology, e.g. PacBio, as well as hybrid approaches, has not yet been applied to lichens. Obtaining the complete set of genome sequences from organisms forming obligate symbioses is challenging. Large-scale cultivation of the individual partners is often not feasible, or aposymbiotic cultivation of the symbionts is entirely impossible. This precludes efforts to obtain pure, single-species DNAs. The alternative approach, reconstructing high-quality genomes from multi-species, metagenomic samples, can be
methodologically demanding (Greshake, et al. 2016). For example, genomic representation can be skewed towards one partner in the association (e.g. the host species), resulting in uneven coverage of individual genomes (Greshake et al 2016). Further methodological challenges include the risk of creating chimeric contigs, i.e. assemblies of reads from multiple genomes, or selecting the appropriate assembly software (Greshake, et al. 2016; Meiser, et al. 2017). Moreover, inaccurate post-assembly taxonomic assignment (binning) can lead to chimeric draft genome sequences, which comprise contigs from multiple species (Sangwan, et al. 2016). Thus, it is highly desirable to assess and develop methods for obtaining metagenome-assembled genomes of eukaryotes, and eventually achieve similar assembly qualities and reporting standards as in prokaryotes (Bowers, et al. 2017).

Here we report the reconstruction of the holo-genome for the lichen *Umbilicaria pustulata* entirely from metagenomic DNA. Details on the biology and distribution of *U. pustulata* have been published elsewhere (e.g. (Dal Grande, et al. 2017; Hestmark 1992)). We inferred the genome sequences of the lichenized fungus *Umbilicaria pustulata*, its green algal symbiont *Trebouxia* sp., and its bacterial microbiome. We combined Illumina short reads from different whole genome shotgun library layouts with PacBio long reads, and integrated results from complementary assembly strategies.

Specifically, we addressed the following questions: What is the quality of fungal and algal organellar and nuclear genomes based on hybrid short and long read assemblies obtained from a metagenomic lichen sample? What are the relative genome copy numbers and the relative taxon abundances of the microorganisms involved in the lichen symbiosis? What is the composition of the bacterial microbiome of a lichen individual? Is there evidence for horizontal gene transfer from algae or bacteria into the
fungal genome? What are the methodological pitfalls associated with reconstructing the holo-genome of symbiotic communities from metagenomic reads, and with their integration into comparative genomics studies focusing on gene loss?

**Materials and Methods**

*Sample collection and DNA extraction*

Thalli of *U. pustulata* were collected near Olbia (Sardinia, Italy) and Orscholz (Saarland, Germany) between May 2013 and December 2014. DNA was extracted using the CTAB method (Cubero and Crespo 2002) and subsequently purified with the PowerClean DNA Clean-Up Kit (MO BIO, Carlsbad, CA, USA).

*Quantitative PCR*

Quantitative PCRs (qPCR) targeted the fungal and algal single copy genes, *mcm7* (Forward - gaatgcaaggaacacatc, Reverse - ttgtactgtctatccgctgg) and *g467* (COP-II coat subunit; Forward - cctcaagtcctatctcg, Reverse - gcactgaaggaagac), respectively. DNA concentrations extracted from four thalli were measured with the Qubit dsDNA High Sensitivity Kit (Life Technologies) according to the manufacturer’s instructions. For qPCR measurements, we used the *GoTaq qPCR Master Mix* (Promega) at a total volume of 10 µl. PCR (95°C for 2 min; 40 cycles of 95°C for 15 sec, 55°C for 30 sec and 60°C for 1 min) was carried out in an *ABI 7500 Fast Real Time PCR system cycler* (Applied Biosystems). Four lichen thalli were measured in three technical replicates. To determine the total copy numbers, we used a standard curve approach with serial ten-fold dilutions of plasmids engineered to contain single copy PCR templates (pGEM®-T Easy Vector, Promega).
**Whole genome shotgun sequencing**

We generated a whole-genome paired-end library with the Illumina TruSeq DNA Sample Prep v2 (Illumina, San Diego, CA, USA), selecting for a mean fragment length of 450 bp with the SPRIselect reagent kit (Beckman Coulter, Krefeld, Germany). A mate pair library with an insert size of 5 kb was created with the Nextera Mate Pair Sample Prep Kit (Illumina, San Diego, CA, USA). The paired-end and mate pair libraries were sequenced on an Illumina MiSeq machine. Long-read sequencing was performed on the PacBio RS II system (Pacific Biosystems of California, Menlo Park, CA, USA), using 16 SMRT cells in total.

**Read preprocessing**

Low quality 3’-ends and adapter sequences were removed from the Illumina paired-end reads with Trimmomatic v0.32 (Bolger, et al. 2014) (*ILLUMINACLIP: IlluminaAdapter.fasta:2:30:10*). Mate pairs were processed with nextclip v0.8 (Leggett, et al. 2014) to remove adapters and to bin them according to read orientation. PacBio sequence reads were error corrected with two alternative strategies. For an intrinsic error correction, we used canu v1.20 (Koren, et al. 2017). Since an intrinsic error correction requires a high long-read coverage, which might not be achieved for the less abundant genomes in the lichen holo-genome, we additionally corrected the PacBio reads using Illumina data as extrinsic information. We merged the Illumina paired-end reads with FLASH v1.2.8 (Magoc and Salzberg 2011), using standard parameters. The processed Illumina read- and mate-pair data were then assembled with MIRA v4.0, using the *genome,denovo,accurate* flags (Chevreux, et al. 1999). The resulting contigs were then used for correcting sequencing errors in the PacBio reads with ECTools ([https://github.com/jgurtowski/ectools](https://github.com/jgurtowski/ectools), last accessed Feb. 27 2020).
requiring a minimum alignment length of 200 bp with a WIGGLE_PCT of 0.05 and a CONTAINED_PCT_ID of 0.8 for the read mappings. Only PacBio reads with lengths after correction of above 1000 bp were retained.

De novo metagenome and metatranscriptome assembly

We employed a multi-layered strategy to target different parts of the lichen holocell genome (see supplementary text, Supplementary Material online for a detailed description of the assembly strategies and supplementary figure S1, Supplementary Material online for the workflow). In brief, we first generated an assembly of the \textit{U. pustulata} metagenome with FALCON v0.2.1 (Chin, et al. 2016) using the uncorrected PacBio reads. The resulting contigs were scaffolded with SSPACE-Long v.1.1 (Boetzer and Pirovano 2014). In parallel, we assembled the error-corrected PacBio reads with the Celera assembler wgs v8.3rc2 (Berlin, et al. 2015). Finally, we made a hybrid assembly with SPAdes v3.5.0 (Bankevich, et al. 2012), that made use of all Illumina reads, the ECTools error-corrected PacBio reads, and the uncorrected PacBio reads to support scaffolding. Subsequent to taxonomic assignment with MEGAN v.5.10 (Huson, et al. 2016) (see below), we binned all algal and bacterial contigs, respectively. They were then merged into single assemblies using \textit{minimus2} (Treangen, et al. 2011) followed by a scaffolding step with SSPACE-Long with the help of the PacBio reads.

For the genome of the fungus \textit{U. pustulata}, the SPAdes contigs of at least 3 Kbp in length were used to further scaffold the FALCON assembly with SSPACE-Long. The final assemblies were polished with Pilon v1.15 (Walker, et al. 2014) using the Illumina short reads.

For the reconstruction of the organellar genomes, we used a baiting strategy. We aligned the canu-corrected PacBio reads against the organellar genomes of the
lecanoromycete fungus *Cladonia grayi* (JGI Clagr3 v2.0) and the green alga *Asterochloris glomerata* (JGI Astpho2 v2.0) (Armaleo, et al. 2019) with BLAT v35 (Kent 2002), using no cut-offs. The baited reads were assembled with canu v1.20, and the resulting organellar genomes were circularized with the help of the canu-corrected PacBio reads and circlator v.1.2.0 (Hunt, et al. 2015). Assembly polishing was performed as described above.

For the reconstruction of the metatranscriptome, we assembled the RNAseq data provided in (Dal Grande, et al. 2017) with Trinity release 2013-11-10 (Haas, et al. 2013), using the --jaccard-clip --normalize_reads parameters.

*Reconstruction of 16S rRNA gene trees*

16S rRNA genes were extracted from the bacterial fraction of the holo-genome assembly. This data was complemented with the 16S rRNA sequences from two new species recently found to be associated with lichens, *Lichenibacter ramalinae* gen. nov., sp. nov. (Pankratov, et al. 2019) and *Lichenihabitans psoromatis* gen. nov., sp. nov. (Noh, et al. 2019). Each gene served as a query for a BlastN search (Altschul, et al. 1997) against the 16S rRNA database of NCBI. The best 5 hits were extracted for each query, except for the sole 16S rRNA gene representing a member of the *Chitonophagaceae*, where we considered the best 10 hits. A non-redundant set of 16S rRNA sequences was generated, and we distinguished five taxonomic bins representing the *Rhizobiales*, *Acidobacteria*, *Chitonophagaceae*, *Actinobacteria*, and *Rhodospirillales*, respectively. Sequences in each bin were aligned with MUSCLE v.3.8.1551 (Edgar 2004) and maximum likelihood phylogenetic trees were computed with RAXML v.8.2.12 (Stamatakis 2014) using the GTRGAMMA model of sequence evolution. Branch support was assessed by performing 100 non-parametric bootstrap
replicates. Phylogenetic trees were visualized and edited with FigTree v.1.4.4 (http://tree.bio.ed.ac.uk/software/figtree/).

**Taxonomic assignment**

All reads and contigs were used individually as query for a DIAMOND v.0.6.12.47 search (Buchfink, et al. 2015). Contigs were searched against a custom database comprising 121 fungi, 20 plants, eight animals, 1471 bacteria and 560 viruses (supplementary table S1, Supplementary Material online), and reads were searched against the NCBI nr database. All sequences were subsequently taxonomically classified with MEGAN v5.10 (Huson, et al. 2016) requiring a minimum DIAMOND alignment score of 50. For MEGAN analyses including more than one read set, we normalized counts to the smallest read set in the analysis. Metagenomic compositions were visualized with Krona (Ondov, et al. 2011).

**Read mapping and coverage distribution analysis**

Reads from the three WGS libraries were mapped to the assembled scaffolds with bowtie2 (Langmead and Salzberg 2012). RNAseq reads of *U. pustulata* (Dal Grande, et al. 2017) were mapped with HISAT2 (Kim, et al. 2015), setting the maximal intron length to 3000 bp and keeping standard parameter values otherwise. To visualize the variation of the WGS read coverages and of the GC content across the different genomes, we split all scaffolds into partitions of 20 Kbp in length, and subsequently clustered the individual partitions by their tetra-nucleotide frequencies. For each partition, we then plotted the mean read coverage for each WGS library, and the mean GC content with Anvi’o (Eren, et al. 2015).
Nuclear and organellar genome annotation

Interspersed repeats were annotated with the RepeatModeler/RepeatMasker pipeline (Smit, et al. 2015). The fungal nuclear genome was annotated with funannotate (https://funannotate.readthedocs.io, last accessed Feb. 27 2020). As training data, we used the proteomes of Xanthoria parietina JGI v1.1 and Cladonia grayi JGI v2.0 (Armaleo, et al. 2019), together with U. pustulata transcripts. The transcripts were obtained in the following way. RNAseq data from U. pustulata (Dal Grande, et al. 2017) was de-novo assembled with Trinity (Haas, et al. 2013). In addition, we performed a second, reference-based assembly of the RNAseq data using Trinity’s reference-guide mode together with the fungal genome assembly. Both assemblies, together with the raw read sets, were used to identify transcripts with PASA (Haas, et al. 2008).

The nuclear genome of Trebouxia sp. was annotated with Maker v2.31.8 (Holt and Yandell 2011), utilizing GeneMark (Besemer and Borodovsky 2005), AUGUSTUS v3.1 (Stanke, et al. 2006), and SNAP v2006-07-28 (Korf 2004). CEGMA (Parra, et al. 2007), RNAseq data (Dal Grande, et al. 2017) and the proteome of Asterochloris glomerata (JGI Astpho2 v2.0) were used for model training. The organelle genomes were annotated using MFannot via the web service provided at http://megasun.bch.umontreal.ca/RNAweasel/ (last accessed Feb. 27 2020).

BLAST2GO (Gotz, et al. 2008) and BlastKOala (Kanehisa, et al. 2016) were used to assign Gene Ontology terms and KEGG identifiers to the predicted genes. The graphic representation of the organellar genomes were generated with OGDraw (https://chlorobox.mpimp-golm.mpg.de/OGDraw.html, last accessed Feb. 27 2020).

Manual curation of gene loss
To assess whether the absence of evolutionary old genes from the *U. pustulata* draft genome sequence is likely a methodological artefact or indeed indicates a gene loss, we performed a gene neighborhood analysis (see Supplementary Material for more detailed methods). In brief, we determined the ortholog to the missing LCA$_{Lec}$ gene in the close relative, *Umbilicaria hispanica* (Dal Grande, et al. 2018), and identified its flanking genes. Next, we searched for the orthologs of these flanking *U. hispanica* genes in *U. pustulata*. We decided on a methodological artefact, if any of these orthologs reside at the terminus of either a contig or a scaffold. Otherwise, we extracted the genomic regions flanking the *U. pustulata* orthologs and used it as a query of a BlastX search (Altschul, et al. 1997) against NCBI nr-prot. In addition, we used the *U. hispanica* protein as query for a tBlastN search in the *U. pustulata* genome assembly. Only when both searches provided no evidence of the missing gene, we inferred gene loss.

**Data accessibility**

The raw Illumina and PacBio sequence reads have been deposited in the NCBI Sequence Read Archive (SRR8446862-SRR8446881). The assemblies have been deposited at GenBank under the accession numbers VXIT00000000 (*U. pustulata* A1-1) and VXIU00000000 (*Trebouxia* sp. A1-2), respectively. The orthologous groups representing the LCA$_{Lec}$ gene set together with the gene annotation of *U. hispanica* are available via [https://applbio.biologie.uni-frankfurt.de/download/lichen/](https://applbio.biologie.uni-frankfurt.de/download/lichen/) (last accessed Feb. 27 2020).

**Results and Discussion**

*Reconstructing the holo-genome sequence of U. pustulata*
*Umbilicaria pustulata* is a rock-dwelling lichen (Figure 1), for which all attempts to cultivate the mycobiont in isolation have failed so far. This leaves a metagenomic approach as currently the only option to reconstruct the genome sequences of the lichen symbionts. qPCR revealed an average ratio of fungal to algal genomes in the lichen thallus of 16.2, with individual replicates varying from a minimum of 13 to a maximum of 24 (supplementary table S2, Supplementary Material online). The heterogeneity between the replicates most likely reflects natural variation in the thickness of the algal layer, and thus algal cell number, within and between lichen thalli (Kummerova, et al. 2006). Such skewed data challenge individual assemblers to an extent that no single tool is capable to faithfully reconstruct all genomes (Bradnam, et al. 2013; Greshake, et al. 2016). We therefore devised a sequencing and assembly scheme to reconstruct the lichen holo-genome at high contiguity (for details on the workflow see supplementary figure S1 and supplementary text, Supplementary Material online). In brief, we used both Illumina short reads and PacBio long read data, and integrated three assemblers: FALCON (Chin, et al. 2016) for assembling uncorrected full length PacBio data; the Celera assembler (Berlin, et al. 2015) for assembling the extrinsically error corrected—and thus often fragmented—PacBio reads; and SPAdes (Bankevich, et al. 2012) for a hybrid assembly of both Illumina and PacBio reads (supplementary figure S1, Supplementary Material online). No individual method sufficed to reconstruct all genomes. A taxonomic assignment of the contigs revealed, however, that the tools complement each other in assembling different parts of the holo-genome at different contiguities (Table 2). Interestingly, SPAdes performed substantially better on the low coverage algal reads than on the more abundant fungal data, both with respect to N50 and number of scaffolds. The difference in N50 reproduced findings from a previous study where NG(A)50 values produced by SPAdes from a simulated lichen holo-
genome were consistently about an order of magnitude smaller for the fungal than for the algal parts of the assembly (Greshake, et al. 2016). Because reads from both species were simulated with the same software, ART (Huang, et al. 2012), this performance difference must be due to an intrinsic characteristic of the fungal genome, most likely its considerably high content of interspersed repeats (25%; see below). The average read coverage of 360x for the fungal genome (Table 3) might represent an additional confounding factor. Anecdotal evidence exists that a too high read coverage impairs the performance of SPAdes. To follow up this point, we used ART (Huang, et al. 2012) to simulate MiSeq whole genome shotgun read sets with average read coverages ranging between 10x and 450x using the U. pustulata scaffolds as template. The corresponding read sets were then individually assembled with SPAdes, and we determined assembly size, number of scaffolds and the scaffold N50 (supplementary table 3, Supplementary Material online). This revealed that coverages around 50x allow excellent genome reconstructions, which only very modestly improve upon increase of the read coverage. More importantly, increasing the coverage beyond 100x results in a constant increase of the number of scaffolds without increasing either assembly size or scaffold N50.

A joint scaffolding of all fungal contigs resulted in a U. pustulata mycobiont genome sequence of 33 Mbp comprising 43 scaffolds with a scaffold N50 of 1.8 Mbp. Merging and scaffolding of the algal contigs generated 217 scaffolds with an N50 of 0.8 Mbp and a total assembly length of 53 Mbp. The assembly lengths for both the fungal and the algal genomes fall well in the diversity of other lichenized fungi and members of the Trebouxiophyceae, respectively (Table 1). Merging and scaffolding the bacterial fraction of the three assemblies resulted in 483 contigs amounting up to 35 Mbp. Two bacterial scaffolds with lengths of 3.6 and 3.4 Mbp represent major parts of two
genomes from the genus *Acidobacterium*. We refer to them as *Acidobacterium BS 16* and *Acidobacterium BS 35*, respectively.

No scaffold in the final assembly represented the full-length genomes of the fungal and algal mitochondria, or of the algal chloroplast. We therefore used the organellar genome sequences of *Cladonia grayi* and of *Asterochloris glomerata* as baits to identify PacBio reads originating from the organellar genomes. The baited reads were assembled individually for each genome, resulting in a circular, gap-free sequence for each of the three organelles (supplementary figures S2-S4, Supplementary Material online). The fungal mitochondrial genome (mt genome) comprises 95.4 kb. It ranks third in length among 23 mt genomes from lecanoromycete lichens (Armaleo, et al. 2019; Pogoda, et al. 2018), superseded only by *Leptogium hirsutum* (120 kbp) and *Parmotrema stuppeum* (109 kbp). The algal mitochondrion and chloroplast have lengths of 99.9 kbp and 272.0 kbp, respectively. They are larger than the organellar genomes in other *Trebouxiophyceae*, both symbiotic and free living (Fan, et al. 2017), with the exception of *Asterochloris glomerata*, which has an even larger mitochondrial genome of 110 kbp in length (Armaleo, et al. 2019).

**Taxon abundance in the lichen holo-genome**

The metagenomic reconstruction of the lichen holo-genome allows, for the first time, to infer average genome copy numbers in a lichen thallus from the read coverage distribution (Table 3, Figure 2; supplementary table S4, Supplementary Material online). The coverage for the fungal nuclear genome assembly, and thus the genomic copy number, is on average about 20 times higher than that of the algal nuclear genome assembly. Similar to the results from the qPCR analysis, the individual estimates vary from a minimum of 9.6 to a maximum of 29.7, which is expected when the thickness
of the algal layer varies within and between lichen thalli (Kummerova, et al. 2006).

Since both symbionts are haploid, this translates into an average abundance of 20 (SD: 7.2) fungal nuclei per algal nucleus. In the mycobiont, there are 15.4 (SD: 4.5) copies of the mitochondrial genome per nuclear genome. This value is substantially lower than the around 60 mtGenome copies per nucleus reported for *Aspergillus fumigatus* (*Eurotiomycetes*) (Neubauer, et al. 2015). It is tempting to speculate that the small number of mitochondrial genomes in the mycobiont is connected to its slow growth.

Yet, too little is known about temporal fluctuations and inter-individual differences in mtGenome content in either species to draw conclusions from this difference. In each *Trebouxia* sp. cell, there are 20 (SD: 7.9) copies of the mitochondrial genome. *Trebouxia* sp. possesses only a single chloroplast. Thus, similar to many other green microalgae (Gallaher, et al. 2018), the *Trebouxia* sp. chloroplast genome is polyploid and contains, on average, 20 (SD: 7.5) copies. To our knowledge, this is the first report of ploidy level for the chloroplast in a lichenized green alga. The two *Acidobacterium* spp. are each represented with about one cell per algal cell.

**Characterization of the bacterial community**

In a first, high resolution approach to characterize individual members of the bacterial community, we identified 21 bacterial scaffolds harboring a 16S rRNA gene. Phylogenetic analyses integrating the 21 16S rRNAs with the most similar sequences represented in the NCBI 16S rRNA database (supplementary S6, Supplementary Material online) grouped the sequences into five major clades, representing *Rhizobiales*, *Rhodospirillales*, *Actinobacteria*, *Chitinophagaceae*, and *Acidobacteria*, respectively (Figure 3). Notably, the *Rhizobiales* tree reveals that the *U. pustulata* microbiome harbors a close relative of *Lichenibacter ramalinae*, which has been previously
identified as an endophytic bacterium in the thalli of subarctic lichens (Pankratov, et al. 2019). Moreover, we found eight 16S rRNA genes that stem from Acidobacteria closely related to Edaphobacter lichenicola, Granulicella aggregans, Acidipila rosea, and Acidobacterium capsulatum. All taxa have been described to inhabit thalli of tundran lichens (Belova, et al. 2018; Pankratov 2012; Pankratov and Dedysh 2010). The remaining 16S rRNA genes represent members of the Rhodospirillales (Alphaproteobacteria; 8 sequences), the Actinobacteria (3 sequences), and the Chitinophagaceae (1 sequence). To our knowledge, neither of these taxa has so far been associated with lichen microbiomes.

To obtain a more comprehensive overview of the bacterial community that is associated with U. pustulata, we performed a taxonomic assignment at the read level (Figure 4 and supplementary figure S5, Supplementary Material online). Acidobacteriaceae, Actinobacteria, and Alphaproteobacteria are the three most abundant bacterial phyla. This is in line with the findings from the 16S rRNA analysis, and it is similar to what has been observed for Antarctic lichens (Park, et al. 2016). In general, the taxonomic composition resembles closely typical rock-inhabiting bacterial communities (Choe, et al. 2018). Yet, other studies suggested that Alphaproteobacteria and not Acidobacteria dominate lichen microbiomes (e.g. (Aschenbrenner, et al. 2014; Bates, et al. 2011; Grube, et al. 2009)), with abundances of up to 32% for the Rhizobiales in the lichen Lobaria pulmonaria (Erlacher, et al. 2015). This indicates that microbiome compositions can vary considerably between lichen species. However, differences in the methodology for assessing taxon frequencies can also result in substantially deviating results (Nayfach and Pollard 2016). The microbiome analyses by Erlacher, et al. (2015) were performed at the level of assembled contigs. While this eases the taxonomic assignment, due to

http://mc.manuscriptcentral.com/gbe
the use of longer sequences (Vollmers, et al. 2017), it is bound to result in distorted abundance estimates. The high read coverage for abundant taxa in a microbiome generally results in more contiguous assemblies comprising only few contigs. In a typical MEGAN analysis, taxon abundance is assessed by the number of sequences that are assigned to that taxon. As a consequence, common taxa with contiguous genome assemblies will receive low counts, and their abundance will be underestimated. Rare taxa, in turn, whose lower read coverage results in more fragmented genome reconstructions with many short contigs will receive high counts. Their abundance will be overestimated (supplementary figure S6, Supplementary Material online). We demonstrate the effect of the chosen methodology on the reconstruction of the U. pustulata microbiome. Applying the method of Erlacher, et al. (2015) increased the estimated abundance of the Rhizobiales to 11%, and decreased that of the Acidobacteriaceae to 18% (supplementary figure S7A, Supplementary Material online). The dominance of the Acidobacteriaceae was restored when pursuing a hybrid approach, in which the taxonomic assignment was done at the contig level and the abundance estimates were based on the reads mapping to the contigs (supplementary figure S7B, Supplementary Material online). We conclude that the methodological impact on the taxon abundance estimates is substantial, and needs to be taken into account when comparing microbiome community composition in different studies.

Annotation of the nuclear genomes

The nuclear genome of U. pustulata (mycobiont) has an average GC content of 51.7%, and interspersed repeats account for 25.5% of the sequence. We identified 9,825 protein-coding genes (Table 2), with on average 3.3 exons, and a mean transcript length of 1,406 bp. A BUSCO analysis (Simao, et al. 2015) revealed that 94.4% of the 1,315
genes in the ‘Ascomycota’ dataset are represented over their full length in the genome sequence. Similarly, FGMP (Cisse and Stajich 2019) found 90% of the 31 highly conserved fungal non-coding elements and 96.8% of the 593 conserved fungal proteins that are represented in the FGMP search set. Both tools indicate a level of assembly completeness that is in the same range of what has been, thus far, achieved only for fungal genomes reconstructed from axenic cultures (Table 1, supplementary table S5, Supplementary Material online). Contrasting to the situation in many other lichens (cf. (Spribille, et al. 2016), we found no evidence for the presence of a second fungus in the lichen thalli (supplementary text, Supplementary Material online).

The genome of *Trebouxia* sp. has an average GC content of 50.0%, and interspersed repeats account for only 4.9% of the sequence. We predicted 13,919 genes with on average 6.7 exons per gene, and a mean transcript length of 1,221 bp. With 13.9%, the fraction of genes from the ‘Chlorophyta’ BUSCO (2,168 genes) that were not found in the genome sequence is considerably high. However, similar results were obtained when analyzing other representatives of the *Trebouxiophyceae* with both free living and symbiotic lifestyles (Table 1). A notable exception, with only 2.4 % missing BUSCOs, is *Coccomyxa subellipsoidea*. This is, however, not surprising since this species was used for the initial compilation of the ‘Chlorophyta’ BUSCO set. We have shown previously that even highly fragmented genome assemblies can recover most of the BUSCO genes (Greshake, et al. 2016). Thus, our results indicate that the plasticity of the algal gene set might be higher than hitherto acknowledged.

No evidence for horizontal gene transfer in *U. pustulata*

The lichen symbiosis, an evolutionarily old, obligate, and stable association of individuals from different species, should provide an optimal basis for the mutual
exchange of genetic material. We therefore screened the fungal genome assembly for indications of horizontal acquisitions of either algal or bacterial genes. Ten fungal genes were classified as of algal, and further 12 as of bacterial origin. All genes are located amidst fungal genes in the genome assembly. However, a subsequent case-by-case curation of these 22 genes revealed that the taxonomic assignments by MEGAN are, in all instances, borderline cases (supplementary table S7, Supplementary Material online). The sequence similarity of the corresponding genes to an algal or bacterial protein, which served as basis for the classification, was low, and only slightly higher than the similarity to the closest fungal gene. Only a slight shift in the parameterization of MEGAN’s taxonomic classification algorithm left these genes essentially taxonomically unassigned. Thus, the true evolutionary origin remains unknown for all 22 genes. Individual examples of genetic exchange between lichenized fungi and their algal partners have been reported before (e.g. (Beck, et al. 2015; Wang, et al. 2014)). Here, we find no convincing evidence for the horizontal acquisition of either algal or bacterial genes by *U. pustulata*.

*Lineage specific absence of evolutionarily old genes in U. pustulata*

We subsequently increased the resolution of the gene set analysis to search for 9,081 genes that were present in the last common ancestor of the *Lecanoromycetes* (LCA\textsubscript{Lec}; see supplementary text, Supplementary Material online). For 142 LCA\textsubscript{Lec} genes we were missing an ortholog only in the *U. pustulata* gene set, suggesting, on the first sight, an exclusive loss on the *U. pustulata* lineage. On closer scrutiny, however, all but 33 of these genes had been either missed during genome annotation, or reside in assembly gaps since an ortholog could be detected in the transcript data. A corresponding analysis in genes exclusively missing in *C. grayi* and in *U. muehlenbergii* obtained similar
results (supplementary text, supplementary table S8 and supplementary figure S8, Supplementary Material online). Taking the absence of genes in annotated gene sets at face value can, therefore, lead to wrong evolutionary inferences (Deutekom, et al. 2019). However, for 33 LCA\textsubscript{Lec} genes we could find, to this point, no indication of an experimental artefact, and they appear genuinely absent from the \textit{U. pustulata} genome assembled by us (supplementary table S9, Supplementary Material online). Four of these genes are represented by an ortholog in the closely related \textit{Umbilicaria hispanica} (Dal Grande, et al. 2018), dating their putative loss to after the split of the two \textit{Umbilicaria} species. In three cases, a subsequent manual curation found no evidence against the gene loss assumption. The three genes encode an oxidoreductase with a significant sequence similarity to gibberellin-20-oxidases, a putative methyltransferase, and a protein with unknown function. The functional consequences of these alleged losses remain to be determined. Moreover, it is not yet clear whether the absence of these genes is fixed within \textit{U. pustulata}, or whether it represents a copy number variation between different populations of this species (Zhao and Gibbons 2018). For the fourth gene encoding a dihydrofolate reductase (DHFR), however, our curation revealed an error source in the gene identification, which is typically neglected. DHFR encodes a protein, which is involved in the basal nucleotide metabolism. This gene is almost ubiquitously present throughout fungi and animals. Its absence in \textit{U. pustulata} therefore would imply far-reaching changes in metabolism (Huang, et al. 1992). Our manual curation could exclude assembly errors and genomic rearrangements as likely explanations for the absence of DHFR (Figure 5). A tBlastN search with the \textit{Saccharomyces cerevisiae} DHFR (Uniprot-ID: P07807) as query obtained a partial hit in this region, which indicated that the ORF of DHFR is disrupted by several frameshift mutations. Because this region is covered by about 200 PacBio reads, sequencing errors
appeared unlikely suggesting a recent pseudogenization of DHFR in the lineage leading to *U. pustulata*. However, we noted a very low Illumina read coverage at the DHFR locus (Figure 5). This coverage drop coincides with an extraordinary high GC content of up to 79% paired with the presence of extended stretches of self-complementarity (Figure 6). In combination, this can lead to the formation of stable stem-loops that can interfere with both DNA amplification and sequencing (Benjamini and Speed 2012; Ross, et al. 2013; Schirmer, et al. 2016). We suspected that the low Illumina read coverage rendered assembly polishing with Pilon less effective. Indeed, a visual inspection exploiting the few Illumina reads that map to the DHFR locus, identified six of eight frameshift mutations as recurrent sequencing errors in the underlying PacBio reads (supplementary figures S9-S14, Supplementary Material online). The remaining two frameshifts towards the 3’-end of the ORF, which are not covered by any Illumina reads, coincide with runs of Gs. Thus, they are very likely to be also sequencing errors (supplementary figures S15-S16, Supplementary Material online). Correcting all frameshifts resulted in an uninterrupted ORF (supplementary figure S17, Supplementary Material online) encoding a full-length DHFR.

To assess the extent to which GC-rich inverted repeats may interfere in general with the correct identification of genes, we annotated inverted repeats throughout the genome draft sequence of *U. pustulata* with the Inverted Repeat Finder (Warburton, et al. 2004). This revealed 1,464 IR, with a median length of 819.5 bp. The GC content of these repeats follows a bimodal distribution peaking at 51% and 75%. While the number of inverted repeats falls within the values obtained for other genomes of lichenized fungi, IRs with a GC content of over 70% are largely unique to *U. pustulata* (Figure 7). Whether this is due to the fact that only *U. pustulata* was sequenced with a long read technology that is less sensitive to GC rich inverted repeats, or whether the other
genomes are devoid of such repeats remains to be determined. Overlaying the IR regions with the Illumina and the PacBio read coverage information reveals 467 IR with a mean GC content of 67.8% for which the Illumina read coverage drops to <10x while the PacBio coverage remains uniformly high. Any gene residing in such a region has a considerable chance to be either incorrectly predicted or overlooked due to remaining sequencing errors in the genome draft sequence.

**Organellar genome annotation**

Annotation of the *L. pustulata* mitochondrial genome resulted in 15 protein-coding genes, a small subunit rRNA gene, 33 additional ORFs, and 31 tRNA genes encoding 24 distinct tRNAs (supplementary figure 2, Supplementary Material online). All 15 fungal core protein coding genes (Pogoda, et al. 2018) are represented, among them atp9, which was found to be frequently missing in the mt genomes of lichenized fungi (supplementary table S10, Supplementary Material online). While this suggests, on the first sight, a considerably standard layout of the mt genome, a closer look at the annotated genes revealed a number of interesting findings. Most notably, *cox2*, the gene encoding the cytochrome c oxidase subunit II is fused head-to-tail to *cob*, which encodes cytochrome b, into one transcription unit (supplementary figure 18, Supplementary Material online). The corresponding Trinity transcript contains an uninterrupted reading frame, suggesting that it is translated into a single fusion protein. To the best of our knowledge, such a fusion as never been reported before, although at least the lecanoromycete *Usnea ceratina* contains a similar fusion (NCBI Gene ID: 34569213). Future studies will have to reveal when during evolution this gene fusion emerged, and at what stage during gene expression—and via what mechanism—the two proteins are separated. Moreover, we noted that nad6, the gene encoding the NADH
dehydrogenase subunit 6, is disrupted by the integration of a 2.4 kb long segment, most likely a mobile Group II intron (Lambowitz and Belfort 1993) (supplementary figure S19, Supplementary Material online). Eventually, three protein-coding genes do not possess a recognizable stop codon (supplementary table S10). One example is the gene encoding the NADH dehydrogenase subunit 3 (nad3). The predicted ORF is covered by three distinct transcripts, indicating that it is not a single transcription unit (supplementary figure S20, Supplementary Material online). A search against the MitoFun database (http://mitofun.biol.uoa.gr, last accessed Feb. 27 2020) reveals that the CDS encoding nad3 spans approximately the first 396 bp of this ORF. In this region, no canonical stop codon is detected, and the agreement between the about 100 individual RNAseq reads and the genomic sequence suggests that no stop codon is generated post-transcriptionally via RNA editing. BlastP and BlastN searches (Altschul, et al. 1997) against the NCBI databases nr-prot and nr, respectively, revealed no significant hits for the parts of the ORF downstream of nad3. The absence of recognizable stop codons in the gene encoding nad3 can be found in the mt genome annotations of other Lecanoromycetes, e.g. in Usnea mutabilis (NCBI GeneID: 38289161) and Parmotrema ultralucens (NCBI GeneID: 38466336). It remains unclear how lichenized fungi achieve an accurate termination of the translation for such genes.

Of the remaining 36 ORFs annotated in the U. pustulata mt genome, 9 encode homing endonucleases that have been proposed to act as selfish genetic elements driving changes in both mt genome size and gene order (Aguileta, et al. 2014; Kanzi, et al. 2016).

The annotation of the Trebouxia sp. mitochondrial genome revealed 32 protein coding genes, 20 additional ORFs, and 26 tRNAs, which agrees with previous findings in the Trebouxiophyceae (Fan, et al. 2017). Similar to other plant and algal species (Ko and
Kim 2016), we found a nuclear copy of the mtGenome (NUMT), which was identified via a local increase of the read coverage in the Anvio’o plot shown in figure 2. In the chloroplast genome, we could annotate 78 protein coding genes, three ribosomal RNAs, 52 additional ORFs, and 31 tRNA. The set of annotated genes comprises all green algal core genes, and additionally 15 out of 16 common algal chloroplast genes showing sporadic lineage specific gene loss (Turmel, et al. 2015). Interestingly, the missing ribosomal protein, rps4, is encoded on scaffold 44 of the algal nuclear genome assembly. Here, it is flanked by two genes, whose counterparts in other green algae are located in the nucleus (supplementary figure S21, Supplementary Material online), and the read coverage pattern provides no hint for any assembly error. This indicates a relocation of rps4 from the chloroplast to the nucleus in *Trebouxia* sp.. Recently, it was hypothesized that a fission of the tRNA-Ile lysidine synthase encoding gene, *tilS* (Suzuki and Miyauchi 2010), observed in mutualistic or parasitic species of the *Trebouxiophyceae* might be connected to symbiosis (Armaleo, et al. 2019). The corresponding gene ycf62 in the chloroplast genome of *Trebouxia* sp. encodes a 725 aa long polypeptide (supplementary figure S4, Supplementary Material online). It harbours the full Pfam domain ATP_bind_3 (PF1171.20) representing the TilS/TtcA_N domain (IPR011063) (supplementary figure S22, Supplementary Material online), similar to the situation in most chlorophyte and streptophyte *tilS* proteins. The two further domains of bacterial tRNA-Ile lysidine synthases described by Suzuki and Miyauchi (2010), *tilS* (PF09179.11) and *tilS-C* (PF11734.8) (supplementary figure S23, Supplementary Material online), are absent from all eukaryotic *tilS* proteins described thus far. In essence, we found no evidence for a fission of this gene in *Trebouxia* sp..

**Conclusion**
Here, we have shown that the reconstruction of the holo-genome for an obligate symbiotic community purely from metagenomic sequence reads at contiguitities comparable to assemblies for single-species samples is feasible. The greatly varying coverage ratios for the individual genomes, spanning three orders of magnitude, emerged as the most challenging task. Key to success was the combination of short Illumina and long PacBio reads with a comprehensive assembly scheme. In particular, we had to (i) target different components of the holo-genome with different assembly methodologies, (ii) include taxonomic assignments on the contig level, (iii) perform a merging of contigs from different assembly approaches that were assigned to the same taxonomic group, and (iv) perform a final scaffolding step. Numerous benchmark studies have indicated that there is no general gold standard for a genome assembly procedure (Dominguez Del Angel, et al. 2018). Thus, our workflow should be considered a template that can be adapted to the needs of the precise symbiotic community under study. The initial analysis of the *U. pustulata* holo-genome already revealed a number of genetic changes both in the nuclear and in the organellar genomes whose functional relevance for this obligate lichen symbiosis will be interesting to determine. However, we encountered also a number of pitfalls that, if remain unnoticed, lead to wrong conclusions. One of the main advantages of metagenomic approaches is that holo-genome reconstruction, relative genomic copy number assessment, taxonomic classification and relative taxon abundance estimation will be performed on the same data. It is tempting to use the assembled contigs for the taxonomic assignments, because longer sequences will allow a classification with greater confidence. If the aim is, however, to assess the abundance of individual taxa in microbial community, the analysis has to take the read data into account. Either by performing the taxonomic assignment at the read level—bearing the risk that a fraction of reads will remain
unclassified—or by taking the read coverage of the taxonomically assigned contigs into account, which will miss rare taxa covered by only few reads. From an evolutionary perspective, the availability of genome sequences for an obligate symbiotic community is the relevant starting point for determining the genetic changes underlying the dependency of the symbionts. A comprehensive gene annotation is essential for such analyses, which have a strong focus on detecting loss of individual genes. BUSCO and FGMP analyses provide an initial indication for the completeness of gene annotations. However, the number of genes in both BUSCO and FGMP sets are, compared to the gene set of a species, typically small, and they are often not designed for the phylogenetic clade in focus, i.e. Lecanoromycetes and Trebouxiophyceae in this study. On the example of the Trebouxiophyceae, we showed that the latter aspect makes it difficult to differentiate between the absence of BUSCO genes due to an incomplete gene set reconstruction, or due to lineage specific losses of BUSCO genes in higher than expected numbers. The use of tailored core gene sets for the clade of interest, paired with targeted ortholog searches both in the annotated gene set and in the assembled transcriptome data is an alternative that substantially increases resolution. Genes that then remain undetected are good candidates for a lineage-specific loss with all its consequences for the symbionts’ metabolism. Still, this does not exclude an artefact. It was only the suspicious deviation in coverage between the PacBio reads and the Illumina reads, which eventually revealed that the gene encoding the dihydrofolate reductase was not lost in U. pustulata. Ultima ratio remains, therefore, expert candidate curation considering all evidences that can hint towards an artefact mimicking gene loss.

Acknowledgements
The authors thank the LOEWE-Centre TBG funded by the Hessen State Ministry of Higher Education, Research and the Arts (HMWK), Anjuli Calchera (Frankfurt) for technical assistance, and Pavel Škaloud (Dpt. Botany, Charles University, Prague) for useful discussion on *Trebusxia* organellar genomics and ontogeny. Moreover, they acknowledge Daniele Armaleo and Basil Britto for providing access to the organelle genomes of *Cladonia greyi* & *Asterochloris glomerata*, and Olafur S. Andresson for sharing the *Lobaria pulmonaria* data. Moreover, we acknowledge two anonymous reviewers for their constructive comments.
# Tables

Table 1. Genome assembly characteristics of a selection of lichenized fungi, and of green algae from the class *Trebouxiophyceae*. The species are sorted by descending scaffold N50. The lichen symbionts sequenced for this study are highlighted in grey.

| Species                        | Size (Mbp) | Scaffolds | N50 (Mbp) | Genes | Missing BUSCO (%) | FGMP: HCE (%) | FGMP: Proteins (%) |
|-------------------------------|------------|-----------|-----------|-------|-------------------|---------------|-------------------|
| *U. muehlenbergii*            | 34.6       | 7         | 7.0       | 8,822 | 1.3               | 90.3          | 94.9              |
| *A. radiata*                  | 33.5       | 17        | 2.2       | na    | 3.0               | 87.1          | 97.8              |
| *U. pustulata*                | 33.5       | 43        | 1.8       | 9,825 | 3.6               | 90.3          | 96.8              |
| *G. flavorubescens*           | 34.5       | 36        | 1.7       | 10,460 | 1.5*              | 77.4          | 97.3              |
| *X. parietina*                | 31.9       | 39        | 1.7       | 11,065 | 1.4*              | 77.4          | 96.6              |
| *C. metacorallifera*          | 36.7       | 30        | 1.6       | 10,497 | 3.0*              | 83.9          | 97.3              |
| *C. macilenta*                | 37.1       | 240       | 1.5       | 10,559 | 2.7*              | 80.6          | 96.3              |
| *P. furfuracea*               | 37.8       | 46        | 1.2       | 8,842  | 1.8               | 93.5          | 97.1              |
| *R. intermedia*               | 26.2       | 198       | 0.3       | na    | 3.3               | 87.1          | 97.3              |
| *E. prunastri*                | 40.3       | 277       | 0.3       | 10,992 | 1.3*              | 87.1          | 96.6              |
| *C. rangiferina*              | 35.7       | 1,069     | 0.3       | na    | 2.5               | 80.6          | 98.0              |
| *C. grayi*                    | 34.6       | 414       | 0.2       | 11,388 | 3.0*              | 87.1          | 96.8              |
| *E. psilium*                  | 36.8       | 908       | 0.2       | 9,238  | 3.9*              | 80.6          | 96.0              |
| *L. hispanica*                | 41.2       | 1,619     | 0.1       | 8,488  | 1.6               | 90.3          | 97.3              |
| *R. peruvianum*               | 27.0       | 1,657     | <0.1      | 9,338* | 6.7*              | 80.6          | 95.4              |
| *L. pulmonaria*               | 56.1       | 1,911     | <0.1      | 15,607 | 1.5*              | 83.9          | 97.0              |
| *C. uncialis*                 | 32.9       | 2,124     | <0.1      | 10,902* | 5.3*            | 87.1          | 97.1              |
| *C. lineartis*                | 19.5       | 2,703     | <0.1      | na    | 25.0              | 51.6          | 83.8              |
| *A. sarmentosum*              | 40.0       | 915       | <0.1      | na    | 21.9              | 58.1          | 83.3              |
| *Alga*                        |           |           |           |       |                   |               |                   |
| *T. gelatinosa*               | 61.7       | 848       | 3.5       | na    | 68.7              | na            | na                |
| *C. subellipsoidea*           | 48.8       | 29        | 2.0       | 9,851  | 2.4               | na            | na                |
| *Chlorella* sp. A99*          | 40.9       | 82        | 1.7       | 8,298  | 18.4              | na            | na                |
| *Treboxxia* sp. L,M           | 52.9       | 217       | 0.8       | 13,919 | 13.9              | na            | na                |
| *A. glomerata*                | 55.8       | 151       | 0.8       | 10,025 | 12.4              | na            | na                |
| *A. protothecoides*           | 22.9       | 374       | 0.3       | 7,016  | 12.2              | na            | na                |
| *Treboxxia* sp. TZW2008*      | 69.3       | 677       | 0.2       | na    | 14.8              | na            | na                |
| *Helicosporidium* sp.*        | 12.4       | 5666      | <0.1      | 6,035  | 50.8              | na            | na                |

§Genome accession numbers are provided in supplementary table S5, Supplementary Material online

$BUSCO$ analysis was performed on the assembly level, *Values taken from Calchera et al. (2019)

*FGMP assembly completeness was determined using 31 highly conserved non-coding elements (HCE) and 593 conserved fungal proteins

*free-living algae, L* lichen photobionts, *other symbiotic algae, M* assemblies resulting from metagenomic sequencing projects
Table 2. Metrics of the metagenome assembly

| Assembly method | Taxonomic classification | Number of Scaffolds | Total length (Mbp) | N50 (kbp) |
|-----------------|--------------------------|---------------------|--------------------|-----------|
| FALCON          | All                      | 2,343               | 62                 | 323       |
|                 | Fungal                   | 120                 | 32                 | 551       |
|                 | Algal                    | 709                 | 9                  | 17        |
|                 | Bacterial                | 790                 | 15                 | 56        |
| SPAdes          | All                      | 21,900              | 123                | 225       |
|                 | Fungal                   | 5,736               | 35                 | 159       |
|                 | Algal                    | 257                 | 47                 | 461       |
|                 | Bacterial                | 1,193               | 26                 | 91        |
| Celera          | All                      | 22,216              | 216                | 11        |
|                 | Fungal                   | 12,230              | 113                | 10        |
|                 | Algal                    | 3,557               | 52                 | 17        |
|                 | Bacterial                | 2,804               | 17                 | 8         |
| Merged (Minimus)| Fungal                   | 43                  | 33                 | 1,808     |
|                 | Algal                    | 217                 | 53                 | 848       |
|                 | Bacterial                | 483                 | 35                 | 251       |

Table 3. Mean read coverages for the fungal and algal components of the *U. pustulata* holo-genome.

| Sequencing technology | Library | *U. pustulata* (mycobiont) | *Trebouxia* sp. |
|-----------------------|---------|----------------------------|-----------------|
|                       |         | Nuclear mtGenome           | Nuclear mtGenome cpGenome |
| Illumina MiSeq        | Mate pair | 40.7 573.3 | 2.5 25.1 | 48.6 |
|                       | Paired End | 123.4 2472.4 | 12.8 239.5 | 214.2 |
| PacBio RS II          | 16 SMRT cells | 195.5 4685.5 | 20.1 754.8 | 776.7 |
Fig. 1.—The lichen *Umbilicaria pustulata*
Fig. 2.—Read coverages and GC content distribution across the genomes in the lichen holo-genome, and the three whole genome shotgun libraries. The genome assemblies were split into non-overlapping bins of 20 Kbp in length and were subsequently clustered according to their tetra-nucleotide frequency. Bins representing the same taxon share the same color. The bar height indicates mean read coverage (black) or mean GC content (green) for each bin. Read coverages are represented on a log scale. The arrows indicate 10x, 100x, and 1000x read coverage, respectively. The mitochondrial genome of the fungus (*U. pustulata* (mt)) is, with a mean read coverage (PacBio) of 3,713, the most abundant component of the holo-genome. The read coverages across the nuclear genome reconstructions of the alga and the fungus are considerably even with only few notable coverage variations (CV). CV1 represents a GC rich (>70%) repetitive region at the terminus of scaffold 8 paired with an assembly gap in this scaffold. The local increase in read coverage of the algal genome
assembly combined with a drop in GC content (CV2) represents a nuclear copy of the algal mitochondrial genome (NUMT).

**Fig. 3.**—16S rRNA phylogenies for bacterial taxa represented in the *U. pustulata* microbiome. A) *Rhizobiales*; B) *Rhodospirillales*; C) *Actinobacteria*; D) *Chitinophagaceae*; E) *Acidobacteria*. 16S rRNA genes for the taxa in blue were extracted from the bacterial fraction of the *U. pustulata* holo-genome reconstruction. The trees reveal that the *U. pustulata* microbiome harbors close relatives to bacterial taxa that have been previously associated with microbiomes of tundran and subarctic lichens (red). Branch labels denote percent bootstrap support. NCBI accession numbers of the sequences are provided in the supplementary table S6, Supplementary Material online.
Fig. 4.—Composition of the bacterial fraction represented in the *U. pustulata* metagenomic reads. Reads from the two Illumina whole genome shotgun libraries and the PacBio reads were pooled and taxonomically assigned with MEGAN (Huson, et al. 2016). *Acidobacteria*, uniting 35% of the read counts, *Proteobacteria* (27%), and *Actinobacteria* (8%) are the three most abundant phyla. Notably, a single family, the *Acidobacteriaceae* (32%), dominates the microbiome. Its most abundant genera are *Granulicella*, *Terriglobus*, and *Acidobacterium* to which the two largest bacterial contigs belong to. Among the *Proteobacteria*, *Rhodospirillales* (6%), and therein the *Acetobactereaceae* (4%) take the largest share, followed by the *Rhizobiales* (4%). Within the *Actinobacteria*, *Actinomycetales* are the dominant
family (3%). See supplementary figure 5, Supplementary Material online, for a species-level resolution of the microbiome.

Fig. 5.—Read coverage distribution in the DHFR locus. Coverage pattern at the DHFR locus (Scaffold 3: 2,310,756-2,313,256). While the read coverage is consistently high for PacBio (~200x), there is a marked decrease for the two Illumina whole genome shotgun libraries towards the center of this region. This decrease coincides with a marked increase of the GC content up to 79%. A tBlastN search using the Dihydrofolate Synthase of *S. cerevisiae* (Uniprot-ID: P07807) obtains a partial hit in the central part of region. Eight frameshift mutations in the CDS of DHFR were manually corrected (supplementary figures S9-S19, Supplementary Material online) resulting in a curated putative protein of 210 aa in length.
Fig. 6.—Inverted repeats in the DHFR locus. We assessed the potential of the DHFR locus to form secondary structures that may interfere with the Illumina sequencing technology. The plot shows self-complementarity predicted by ProbKnot (Bellaousov and Mathews 2010) as black arcs. The pattern reveals that the DHFR gene in *U. pustulata* is embedded in an inverted repeat spanning approximately 800 bp.
Fig. 7.—The distribution of inverted repeats in the draft genome sequences of five lichenized fungi. Inverted repeats with a GC content above 70% are observed only in *U. pustulata*. 
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Gene prediction

tBLASTN

RNAseq

PairedEnd

MatePair

PacBio

GC content

2,477 bp

[0 - 808]

[0 - 154]

[0 - 77]

[45 - 100%]

Scaffold 3:

2310756 - 2313256 bp

Gene prediction

FUN_02688

DHFR S. cerevisiae

FUN_02687

DHFR L. pustulata curated

http://mc.manuscriptcentral.com/gbe/advance-article-abstract/doi/10.1093/gbe/evaa049/5803651 by guest on 15 March 2020
Table 1. Genome assembly characteristics of a selection of lichenized fungi, and of green algae from the class Trebouxiophyceae. The species are sorted by descending scaffold N50. The lichen symbionts sequenced for this study are highlighted in grey.

| Species       | Size (Mbp) | Scaffolds | N50 (Mbp) | Genes | Missing BUSCO (%) | FGMP: HCE (%) | FGMP: Proteins (%) |
|---------------|------------|-----------|-----------|-------|-------------------|---------------|-------------------|
| **Fungus**    |            |           |           |       |                   |               |                   |
| U. muehlenbergii | 34.6       | 7         | 7.0       | 8,822 | 1.3               | 90.3          | 94.9              |
| A. radiata    | 33.5       | 17        | 2.2       | 8,822 | na                | 3.0           | 87.1              |
| U. pustulataM | 33.5       | 43        | 1.8       | 9,825 | 3.6               | 90.3          | 96.8              |
| G. flavorubescens | 34.5       | 36        | 1.7       | 10,460* | 1.5*             | 77.4          | 94.9              |
| X. paretina    | 31.9       | 39        | 1.7       | 11,065 | 1.4*             | 77.4          | 96.6              |
| C. metacorallfera | 36.7       | 30        | 1.6       | 10,497* | 3.0*             | 83.9          | 97.3              |
| C. macilenta   | 37.1       | 240       | 1.5       | 10,559* | 2.7*             | 80.6          | 96.3              |
| P. furfuracea  | 37.8       | 46        | 1.2       | 8,842  | 1.8               | 93.5          | 97.1              |
| R. intermedia  | 26.2       | 198       | 0.3       | na     | 3.3               | 87.1          | 97.3              |
| E. prunastri   | 40.3       | 277       | 0.3       | 10,992 | 1.3*             | 87.1          | 96.6              |
| C. rangiferina | 35.7       | 1,069     | 0.3       | na     | 2.5               | 80.6          | 98.0              |
| C. grayi       | 34.6       | 414       | 0.2       | 11,388 | 3.0*             | 87.1          | 96.8              |
| E. pusillum    | 36.8       | 908       | 0.2       | 9,238  | 3.9*             | 80.6          | 96.0              |
| L. hispanica   | 41.2       | 1,619     | 0.1       | 8,488  | 1.6               | 90.3          | 97.3              |
| R. peruviana   | 27.0       | 1,657     | <0.1      | 9,338* | 6.7*             | 80.6          | 95.4              |
| L. pulmonaria  | 56.1       | 1,911     | <0.1      | 15,607 | 1.5*             | 83.9          | 97.0              |
| C. uncialis    | 32.9       | 2,124     | <0.1      | 10,902* | 5.3*             | 87.1          | 97.1              |
| C. linearisM  | 19.5       | 2,703     | <0.1      | na     | 25.0             | 51.6          | 83.8              |
| A. sarmentosaM | 40.0       | 915       | <0.1      | na     | 21.9             | 58.1          | 83.3              |
| **Alga**      |            |           |           |       |                   |               |                   |
| T. gelatinosaL | 61.7       | 848       | 3.5       | na     | 68.7             | na           | na                |
| C. subellipsoideaF | 48.8       | 29        | 2.0       | 9,851  | 2.4              | na           | na                |
| Chlorella sp. A99S | 40.9       | 82        | 1.7       | 8,298  | 18.4             | na           | na                |
| Trebouxia sp. L,M | 52.9       | 217       | 0.8       | 13,919 | 13.9             | na           | na                |
| A. glomerataL | 55.8       | 151       | 0.8       | 10,025 | 12.4             | na           | na                |
| A. protococcoidesF | 22.9       | 374       | 0.3       | 7,016  | 12.2             | na           | na                |
| Trebouxia sp. TZW2008L | 69.3       | 677       | 0.2       | na     | 14.8             | na           | na                |
| Helicosporidium sp. S | 12.4       | 5666      | <0.1      | 6,035  | 50.8             | na           | na                |

§Genome accession numbers are provided in supplementary table S5, Supplementary Material online

$BUSCO analysis was performed on the assembly level, *Values taken from Calchera et al. (2019)

*FGMP assembly completeness was determined using 31 highly conserved non-coding elements (HCE) and 593 conserved fungal proteins

Ffree-living algae, Llichen photobionts, Sother symbiotic algae, Massemblies resulting from metagenomic sequencing projects

http://mc.manuscriptcentral.com/gbe
Table 2. Metrics of the metagenome assembly

| Assembly method | Taxonomic classification | Number of Scaffolds | Total length (Mbp) | N50 (kbp) |
|-----------------|--------------------------|---------------------|--------------------|-----------|
| **FALCON**      |                           |                     |                    |           |
| All             | 2,343                    | 62                  | 323                |           |
| Fungal          | 120                      | 32                  | 551                |           |
| Algal           | 709                      | 9                   | 17                 |           |
| Bacterial       | 790                      | 15                  | 56                 |           |
| **SPAdes**      |                           |                     |                    |           |
| All             | 21,900                   | 123                 | 225                |           |
| Fungal          | 5,736                    | 35                  | 159                |           |
| Algal           | 257                      | 47                  | 461                |           |
| Bacterial       | 1,193                    | 26                  | 91                 |           |
| **Celera**      |                           |                     |                    |           |
| All             | 22,216                   | 216                 | 11                 |           |
| Fungal          | 12,230                   | 113                 | 10                 |           |
| Algal           | 3,557                    | 52                  | 17                 |           |
| Bacterial       | 2,804                    | 17                  | 8                  |           |
| **Merged (Minimus)** |                     |                     |                    |           |
| Fungal          | 43                       | 33                  | 1,808              |           |
| Algal           | 217                      | 53                  | 848                |           |
| Bacterial       | 483                      | 35                  | 251                |           |
Table 3. Mean read coverages for the fungal and algal components of the *U. pustulata* holo-genome

| Sequencing technology | Library   | *U. pustulata* (mycobiont) | *Trebouxia* sp. |
|-----------------------|-----------|-----------------------------|-----------------|
|                       |           | Nuclear | mtGenome | Nuclear | mtGenome | cpGenome |
| Illumina MiSeq        | Mate pair| 40.7    | 573.3    | 2.5     | 25.1     | 48.6     |
|                       | Paired End| 123.4   | 2472.4   | 12.8    | 239.5    | 214.2    |
| PacBio RS II          | 16 SMRT cells| 195.5 | 4685.5 | 20.1    | 754.8    | 776.7    |