

Nuclear and Mitochondrial Genome Assemblies for the Endangered Wood-Decaying Fungus *Somion occarium*

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Accepted: December 20, 2024

Abstract

Somion occarium is a wood-decaying bracket fungus belonging to an order known to be rich in useful chemical compounds. Despite its widespread distribution, *S. occarium* has been assessed as endangered on at least 1 national Red List, presumably due to loss of old-growth forest habitat. Here, we present a near-complete, annotated nuclear genome assembly for *S. occarium* consisting of 31 Mbp arranged in 11 pseudochromosomes—9 of which are telomere-to-telomere—as well as a complete mitochondrial genome assembly of 112.9 Kbp. We additionally performed phylogenomic analysis and annotated carbohydrate-active enzymes (CAZymes) to compare gene and CAZyme content across closely related species. This genome was sequenced as the representative for Kingdom *Fungi* in the European Reference Genome Atlas Pilot Project.

Key words: saprotroph, white-rot, cerrenaceae, polyporales, fungal conservation.

Significance

Wood-decaying fungi are not only the foundation of nutrient cycling in our forests, but also known to produce many medically relevant chemical compounds. As *Somion occarium* is also endangered in at least 1 country, it is doubly important to produce high-quality genomic resources to facilitate study of this species. The sole fungal representative sequenced as part of the European Reference Genome Atlas (ERGA) Pilot Project, this genome assembly leads the way for future sequencing efforts of fungi within ERGA.

Introduction

The species *Somion occarium* (Cerrenaceae, Polyporales, Basidiomycota) is a bracket fungus and polypore (Fig. 1a), both of which are polyphyletic groupings based on growth

form rather than ancestry. Usually found on dead wood of hardwood trees—including oaks (*Quercus* spp.) and beech (*Fagus* spp.) (GBIF 2024)—*S. occarium* is predominantly a white-rot decayer, or saprotroph. Its occasional presence

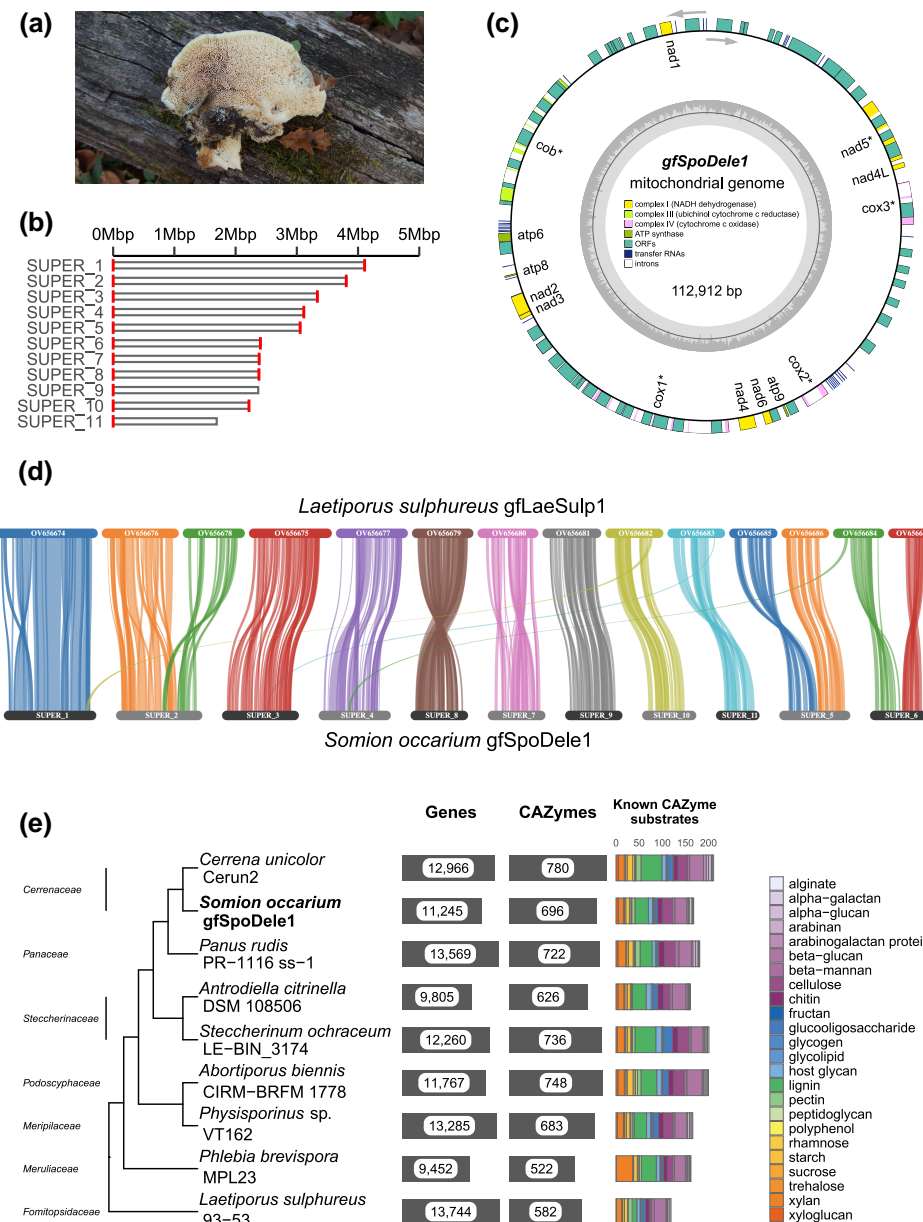


Fig. 1. a) Photograph of a *S. occarium* specimen from the same locality as the strain sequenced in this study, taken in 2018. b) A representation of the assembled pseudo-chromosomal contigs, with red bars at each end indicating predicted telomeres. c) The assembled and annotated mitogenome. d) Synteny between *S. occarium* *gfSpoDole1* and *L. sulphureus* *gfLaeSulp1*. e) Comparison of gene content between *S. occarium* and closely related *Polyporales* species, with an emphasis on CAZymes. Known substrates are only shown for CAZyme genes with no contradictory substrate predictions by run_dbcan.

on living trees suggests it is also capable of facultative parasitic growth, similar to other species in the Cerrenaceae (Hallenberg et al. 2008; Justo et al. 2017). In addition to wood, the fungus has also occasionally been recorded from soil DNA metabarcoding (Větrovský et al. 2020). Such wood-decaying fungi are essential for carbon and nutrient cycling and the formation and stabilization of soil aggregates in forest ecosystems (Miller and Lodge 2007).

Somion occarium belongs to the order *Polyporales*, which contains many, but not all, of the polypore species. Polypores are iconic in forest ecosystems, and have long been culturally valued, including a rich history of their use in folk medicine (Grienke et al. 2014). Fittingly, the *Polyporales* are some of the highest known producers of bioactive compounds within fungi, including antibacterials, antifungals, and drugs or drug leads (Prescott et al. 2023). Generating genomic data for species in the order provides

the groundwork to support further discovery of potentially useful compounds.

The Earth BioGenome Project (EBP) is an ambitious initiative to sequence reference genomes for all eukaryotic species (Lewin et al. 2018). The European Reference Genome Atlas (ERGA) is the European node of EBP, and has coordinated a pilot project to establish and test a decentralized infrastructure to deliver the EBP's aspirations (Mc Cartney et al. 2024), for which *S. occarium* was selected as a representative for fungi. *Somion occarium* was previously known as *Spongipellis delectans* (or the synonym *Hydnum occarium*), but recent division of *Spongipellis* *sensu lato* into several genera following DNA-based and morphological revision saw the restoration of the suppressed older genus name *Somion* for the *Sp. delectans* complex (Miettinen et al. 2023). North American *Sp. delectans* is now *Somion delectans*, while European *Sp. delectans* is *S. occarium*. There is 1 existing assembly for the Cerrenaceae family, which *S. occarium* belongs to, for the species *Cerrena unicolor* (<https://mycocosm.jgi.doe.gov/Cerun2/Cerun2.home.html>), although the *S. occarium* ERGA assembly reported here represents the first EBP-level reference genome for the family.

While *S. occarium* is found across the northern temperate zone in Europe and northernmost Africa (Miettinen et al. 2023), the species is classed as endangered on the national Red List for the Czech Republic (Zíbarová et al. 2024), the locality from which the strain sequenced here was collected. Wood-decaying fungi are predominantly dependent on deadwood, which is more available in old-growth forests compared with younger or managed forests/plantations, and so the ongoing loss of old-growth forest habitat means that wood-decaying fungi are increasingly threatened by niche reduction (Jönsson et al. 2008; Lonsdale et al. 2008). Incidentally, this also means that certain fungal species can be used as indicator species for old-growth forest (Halme et al. 2009). Recognition of fungi in conservation discourse is relatively new (May et al. 2018), but progress is being made thanks to the enduring efforts of mycologists to raise their profile in Red Lists (Dahlberg and Mueller 2011; Mueller et al. 2014), as well as growing public and political awareness of the importance of fungi to ecosystems (IUCN 2021). Producing high-quality genomic resources for endangered species such as *S. occarium* is foundational to understanding and protecting them.

Results and Discussion

Near-complete Genome Assembly for *S. occarium*

We assembled the *S. occarium* genome using hifiasm (Cheng et al. 2021) from ~100-fold coverage of PacBio HiFi reads, with a mean read length of 18.6 Kbp. The final, manually curated assembly (Table 1) exceeded the minimum reference standard defined by EBP assembly quality metrics (EBP 2023). Note that this final version of the

Table 1 Summary statistics for the gfSpoDele1 assembly and annotation

	Nuclear genome	Mitogenome
Number of contigs	11	1
Total size (bp)	30,870,404	112,912
N50 (Mbp)	3.05	...
Kmer QV	66.53	...
Number of protein-coding genes (high confidence/low confidence)	11,245 (9,409/1,836)	14 (35 ORFs, 19 intronic ORFs)
BUSCO completeness (assembly)	C: 94.4% [S: 93.8%, D: 0.6%], F: 0.8%, M: 4.8%, n: 4464	...
BUSCO completeness (annotation)	C: 98.6% [S: 97.7%, D: 0.9%], F: 0.1%, M: 1.3%, n: 4464	...

assembly is a primary assembly and is highly heterozygous (1.69%; [supplementary fig. S1, Supplementary Material online](#)).

We predicted telomeres on both ends of 9 out of the total 11 pseudo-chromosomes, with the remaining 2 contigs (SUPER_9 and SUPER_11) having a telomere at one end (Fig. 1b). A total of 11 chromosomes was slightly lower than the number that has been previously reported for other *Polyporales* species assembled to chromosome level, which ranges from 12 to 14 (Chen et al. 2012, 2022; Wright et al. 2022; Ma et al. 2024), although this was explained by 3 chromosomal "fusions" that were flagged when we examined synteny relative to the *Laetiporus sulphureus* gfLaeSulp1 assembly (Fig. 1d). Each of these fusion events was supported following curation and manual checking of read alignments.

In addition to the nuclear genome assembly, we assembled and annotated the mitochondrial genome (Fig. 1c), which was a single circular contig 112.9 Kbp in length and included all 14 core protein-coding genes typically present in fungal mitogenomes (Sandor et al. 2018).

Gene and CAZyme Content

Gene models were generated using REAT (Robust Eukaryotic Annotation Toolkit; <https://github.com/EI-Core/Bioinformatics/reat>) and functionally annotated with AHRD (<https://github.com/groupschoof/AHRD>; [supplementary file S1, Supplementary Material online](#)). We compared gene content across species with available genome annotations within the *Polyporales* "residual" clade as defined by Justo et al. (2017), including *L. sulphureus* as an outgroup ([supplementary table S1, Supplementary Material online](#)). This first involved inference of a species tree from orthologous genes using OrthoFinder (Emms and Kelly 2019), where we found that the divergence of families in the phylogenomic tree corresponded with previous multilocus phylogenetic analysis within the order (Justo et al. 2017).

We additionally predicted carbohydrate-active enzyme (CAZyme) genes using run_dbcan (Zheng et al. 2023; [supplementary file S2, Supplementary Material](#) online), as CAZymes are a major component of the gene repertoire necessary for white-rot (Hage et al. 2021). The total number of genes and CAZymes was slightly lower than either the closest available relative within Cerrenaceae, *Cerrena unicolor*, or *Panus rufus* in the sister family of Panaceae (Fig. 1e). The proportions of CAZymes known to act on various substrates were similar across all taxa, however.

Conclusion

Here, we present the first near-complete, annotated genome assembly for the wood-decaying bracket fungus *S. occarium* (previously *Sp. delectans*). The first representative for Kingdom *Fungi* under the ERGA initiative, this new high-quality genome resource will enable further exploration of the genetic basis of saprotrophy in an ecologically and chemically important lineage of fungi, the *Polyporales*.

Materials and Methods

Sample Collection and Isolation

The *S. occarium* strain sequenced here was collected as a fresh basidiome growing on dead wood of *Quercus cerris*, on 2005 October 20 at the Rendez-vous nature monument in Czech Republic (Valtice, Břeclav, South Moravian region, 48.7499006N, 16.7939872E). This is the same locality from which the epitype of *S. occarium* was collected (Miettinen et al. 2023). The basidiome was placed on a petri dish of 2% malt extract agar, and the resulting culture was deposited in the culture collection of MENDELU (accession MUcc 838) and simultaneously kept in the Culture Collection of Basidiomycetes of the Institute of Microbiology, Prague (accession CCBAS136). Its identity was confirmed using phylogenetic analysis by Tomšovský (2012), under the previous name of *Sp. delectans*.

Nucleic Acid Isolation and Sequencing

DNA and RNA Extractions

Fresh mycelia were obtained after 2 week of stationary cultivation in 2% malt extract liquid media (24 °C, dark), washed in deionized water, hand-squeezed, and stored at –80 °C prior to DNA or RNA extraction. High molecular weight (HMW) DNA was isolated in 8 aliquots, each consisting of up to 200 mg of frozen biomass, using a modified phenol–chloroform extraction (Sagova-Mareckova et al. 2008). Frozen biomass was homogenized in liquid nitrogen to a fine powder. All vortexing steps in the workflow were replaced by 50x repeated inversions of tubes to prevent DNA fragmentation. DNA was eluted in 10 mM Tris-Cl, pH 8.5, and stored at –80 °C. RNA was extracted in 6

aliquots of at least 50 mg of mycelia using the Quick RNA Fungal/Bacterial Miniprep kit (Zymo Research) according to the manufacturer's protocol, excepting DNase treatment, which was carried out separately using the Turbo DNA-free kit (Invitrogen) and stored at –80 °C. The quality of the isolated DNA/RNA was checked with gel electrophoresis, where samples with HMW DNA or RNA showing 3 separate bands for RNA subunits were considered acceptable.

PacBio HiFi Genome Sequencing

Four HWM DNA extractions were combined to construct a PacBio HiFi library at the Earlham Institute, Norwich, UK using the SMRTbell Express Template Prep Kit 2.0 (PacBio, P/N 100-983-900). In total, 18 µg of HWM DNA was manually sheared with the Megaruptor 3 instrument (Diagenode, P/N B06010003) according to the operations manual. After shearing, the sample underwent AMPure PB bead (PacBio, P/N 100-265-900) purification and concentration before undergoing library preparation using the SMRTbell Express Template Prep Kit 2.0 (PacBio, P/N 100-983-900). The library was prepared according to the HiFi protocol v03 (PacBio, P/N 101-853-100) and the final library was size fractionated using the SageELF system (Sage Science, P/N ELF0001) and a 0.75% cassette (Sage Science, P/N ELD7510). The library was quantified by fluorescence (Invitrogen Qubit 3.0, P/N Q33216) and the size of the library fractions was estimated from a smear analysis performed on the FEMTO Pulse System (Agilent, P/N M5330AA). The loading calculations for sequencing were completed using the PacBio SMRTLink Binding Calculator 10.1. Sequencing primer v2 was annealed to the adapter sequence of the HiFi library. The library was bound to the sequencing polymerase with the Sequel II Binding Kit v2.0 (PacBio, P/N 101-842-900). Calculations for primer and polymerase binding ratios were kept at default values for the library type. Sequel II DNA internal control 1.0 was spiked into the library at the standard concentration prior to sequencing. The sequencing chemistry used was Sequel II Sequencing Plate 2.0 (PacBio, P/N 101-820-200) and the Instrument Control Software v10.1.0.125432. The library was sequenced on the Sequel IIe on 1 Sequel II SMRTcell 8 M. The parameters for sequencing were diffusion loading, a 30-h movie, a 4-h pre-extension time, a 2-h immobilization, and 70 pM on plate loading concentration.

PacBio Iso-Seq

One PacBio Iso-Seq library was constructed starting from 315 ng of total RNA according to the guidelines laid out in the Iso-Seq protocol v02 (PacBio, 101-763-800), using SMRTbell express template prep kit 2.0 (PacBio, 102-088-900). Please see the [Supplementary Material](#) for full details. The sequencing chemistry used was Sequel II

Sequencing Plate 2.0 (PacBio, 101-820-200) and the Instrument Control Software v10.1.0.119549. The Iso-Seq library was sequenced on the Sequel IIe instrument with 1 Sequel II SMRTcell 8 M cell. The parameters for sequencing were diffusion loading, a 30-h movie, a 2-h immobilization time, a 2-h preextension time, and an 80 pM on plate loading concentration.

Illumina RNA-Seq

One RNA-Seq library was constructed using the NEBNext Ultra II RNA Library prep for Illumina kit (NEB#E7760L), NEBNext Poly(A) mRNA magnetic isolation module (NEB#7490), and NEBNext Multiplex Oligos for Illumina (E6440S) at a concentration of 10 µM. Please see the [Supplementary Material](#) for full details. The library was loaded onto a v1.5 NovaSeq SP flow cell using the NovaSeq Xp flow cell dock. The flow cell was then loaded onto the NovaSeq 6000 along with a NovaSeq 6000 v1.5 SP cluster cartridge, a buffer cartridge, and a 300-cycle SBS cartridge (Illumina). The NovaSeq was run with NVCS v1.7.5 and RTA v3.4.4, and was set up to sequence 150 bp paired-end reads. The data were demultiplexed and converted to fastq using bcl2fastq2.

Genome Assembly

PacBio HiFi reads were randomly subsampled to ~100× coverage and assembled using hifiasm v0.16.1 (Cheng et al. 2021). Alternative haplotypes were removed using purge_dups v1.0.1 (Guan et al. 2020). The assembly was manually curated by loading it into the Gap5 sequence editor (Bonfield and Whitwam 2010), and each contig's integrity was assessed. Four breaks were made due to erroneously assembled reads at contig ends, and 1 join was made between 2 contigs based on repeat analysis and genome synteny with the *Trametes hirsuta* reference genome (accession GCA_001302255.2).

Assembly completeness was estimated using BUSCO v5.4.7 with the Polyporales_odb10 dataset (Manni et al. 2021). The assembly consensus quality value (QV) was estimated using Merquery v1.3 (Rhee et al. 2020). Telomeric repeats (TTAGGG) at the ends of fragments were identified using tick v0.2.31 (Brown 2023), with at least 5 repeats required for positive telomere prediction. The mitochondrial genome was assembled separately using IPA (<https://github.com/PacificBiosciences/pipa>).

Genome Annotation

Annotation of repetitive elements was performed using the El-Repeat pipeline v1.1.0 (<https://github.com/El-CoreBioinformatics/eirepeat>), which masked the genome assembly using RepeatMasker v4.0.7 (Smit et al. 2015) with a repeat library from RepBase and a *de novo* library of repeats

constructed using RepeatModeler v1.0.11 (Smit and Hubley 2015).

Gene models were annotated using REAT (<https://github.com/El-CoreBioinformatics/reat>) and Minos (<https://github.com/El-CoreBioinformatics/minos>), incorporating transcript assemblies from Illumina RNA-Seq and PacBio Iso-Seq data, alignment of protein sequences from related species, and evidence-guided gene prediction with AUGUSTUS (Stanke and Morgenstern 2005). Please see the [Supplementary Material](#) for full details. Gene models were functionally annotated using El-FunAnnot pipeline v1.3 (<https://github.com/El-CoreBioinformatics/eifunannot>) utilizing AHRD v3.3.3 (<https://github.com/groupschoof/AHRD>) with hits against fungal proteins from both Swiss-Prot and TrEMBL (downloaded 2022 June 15; The UniProt Consortium 2021) generated with BLAST v2.6.0 (Camacho et al. 2009) as well as results from InterProScan v5.22.61 (Jones et al. 2014).

The mitochondrial genome was annotated using MFannot v1.35 (Lang et al. 2023) and visualized using OGDRAW (Greiner et al. 2019).

Assembly and annotation data were submitted to the European Nucleotide Archive using COPO (Shaw et al. 2020).

Phylogenomics and Comparative Genomics

Assessment of syntenic conservation and collinearity with the chromosome-scale genome assembly of *L. sulphureus* gfLaeSulp1 (Wright et al. 2022) was performed using MCScanX v1.1 (Wang et al. 2012) and visualized using SynVisio (Bandi and Gutwin 2020).

For phylogenomic reconstruction of the *Polyporales* "residual" clade *sensu* Justo et al. (2017), we inferred orthologous gene families by running OrthoFinder v2.5.4 (Emms and Kelly 2019) on the *S. occarium* proteome alongside proteomes from other residual *Polyporales* taxa with available data and *L. sulphureus* as an outgroup ([supplementary table S1, Supplementary Material](#) online). CAZyme genes for all taxa were predicted using run_dbcan v4 (Huang et al. 2018; Zhang et al. 2018; Zheng et al. 2023) and visualized alongside the STAG species tree produced by OrthoFinder.

Supplementary Material

[Supplementary material](#) is available at *Genome Biology and Evolution* online.

Acknowledgments

We thank the essential work of the ERGA Pilot Project coordinators, Ann Mc Cartney, Giulio Formenti and Alice Mouton, in making this work possible with their coordination effort as part of the ERGA Pilot Project daily activities.

Many thanks to Felix Shaw, Debby Ku and Aaliyah Providence for their assistance with the COPO data upload. We thank Steven Singer for permitting the use of unpublished data for *Cerrena unicolor* from Mycosom. The authors acknowledge the work delivered via the Laboratory Managers and Research Computing Groups at Earlham Institute who manage and deliver High Performance Computing at EI. We thank Thomas Brabbs and James Lipscombe for assistance and advice with laboratory automation, and Neil Shearer for LIMS support.

Author Contributions

Rowena Hill: formal analysis, investigation, methodology, visualization, writing—original draft, writing—review & editing. Jamie McGowan: formal analysis, investigation, methodology, visualization, writing—original draft, writing—review & editing. Vendula Brabcová: methodology, investigation, writing—original draft, writing—review & editing. Seanna McTaggart: project administration, writing—review & editing. Naomi Irish: methodology, investigation, writing—original draft. Tom Barker: investigation, writing—original draft. Vanda Knitlhofer: investigation. Sacha Lucchini: methodology. Kendall Baker: project administration. Leah Catchpole: supervision. Chris Watkins: project administration, supervision. Karim Gharbi: resources, supervision, writing—review & editing. Gemy Kaithakottil: formal analysis, resources, software, writing—review & editing. Alan Tracey: data curation. Jonathan M.D. Wood: data curation, supervision. Michal Tomšovský: resources, writing—original draft, writing—review & editing. Petr Baldrian: resources, writing—original draft, writing—review & editing. David Swarbreck: resources, software, supervision, writing—review & editing. Neil Hall: resources, supervision, writing—review & editing

Funding

This work was supported by funding from the Biotechnology and Biological Sciences Research Council (BBSRC), part of UK Research and Innovation, Core Capability Grant (BB/CCG1720/1). Part of this work was delivered via the BBSRC National Capabilities in Genomics and Single Cell Analysis grant (BBS/E/T/000PR9816) by members of the Technical Genomics and the Core Bioinformatics Groups, and in e-Infrastructure (BBS/E/T/000PR9814) by members of the e-Infrastructure group at the Earlham Institute. Part of this work was supported by the Earlham Institute Strategic Programme Grant Decoding Biodiversity (BBX011089/1) and its constituent work package—Decode WP2 Genome Enabled Analysis of Diversity to Identify Gene Function, Biosynthetic Pathways, and Variation in Agri/Aquacultural Traits (BBS/E/ER/230002B). P.B. and V.B. were supported by the

Czech Science Foundation (21-09334J) and by the Ministerstvo Školství, Mládeže a Tělovýchovy (Ministry of Education, Youth and Sports of the Czech Republic; Talking microbes—understanding microbial interactions within One Health framework; CZ.02.01.01/00/22_008/0004597).

Data Availability

Genome assembly and annotation data are available in the European Nucleotide Archive under BioProject PRJEB75241 and accession GCA_964035595.1.

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Associate editor: Li-Jun Ma