

Independent mechanisms of benzimidazole resistance across *Caenorhabditis* nematodes

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28 **ABBREVIATIONS**

29	ABZ	Albendazole
30	BZs	Benzimidazoles
31	<i>Cbr-ben-1</i>	<i>ben-1</i> in <i>Caenorhabditis briggsae</i>
32	<i>Cbr-BEN-1</i>	BEN-1 in <i>Caenorhabditis briggsae</i>
33	<i>Cbr-tbb-1</i>	<i>tbb-1</i> in <i>Caenorhabditis briggsae</i>
34	<i>Cbr-TBB-1</i>	TBB-1 in <i>Caenorhabditis briggsae</i>
35	<i>Cbr-tbb-2</i>	<i>tbb-2</i> in <i>Caenorhabditis briggsae</i>
36	<i>Cbr-TBB-2</i>	TBB-2 in <i>Caenorhabditis briggsae</i>
37	<i>Cel-ben-1</i>	<i>ben-1</i> in <i>Caenorhabditis elegans</i>
38	<i>Cel-BEN-1</i>	BEN-1 in <i>Caenorhabditis elegans</i>
39	<i>Cel-tbb-1</i>	<i>tbb-1</i> in <i>Caenorhabditis elegans</i>
40	<i>Cel-TBB-1</i>	TBB-1 in <i>Caenorhabditis elegans</i>
41	<i>Cel-tbb-2</i>	<i>tbb-2</i> in <i>Caenorhabditis elegans</i>
42	<i>Cel-TBB-2</i>	TBB-2 in <i>Caenorhabditis elegans</i>
43	<i>Ctr-ben-1</i>	<i>ben-1</i> in <i>Caenorhabditis tropicalis</i>
44	<i>Ctr-BEN-1</i>	BEN-1 in <i>Caenorhabditis tropicalis</i>
45	<i>Ctr-tbb-1</i>	<i>tbb-1</i> in <i>Caenorhabditis tropicalis</i>
46	<i>Ctr-TBB-1</i>	TBB-1 in <i>Caenorhabditis tropicalis</i>
47	<i>Ctr-tbb-2</i>	<i>tbb-2</i> in <i>Caenorhabditis tropicalis</i>
48	<i>Ctr-TBB-2</i>	TBB-2 in <i>Caenorhabditis tropicalis</i>
49	HSB	Horvitz Super Broth
50	HTA	High-throughput larval development assay
51	LoF	Loss-of-function
52	NGMA	Nematode growth media agar
53	SNV	Single nucleotide variants

SV Structural variants

55 **ABSTRACT**

56 Benzimidazoles (BZs), a widely used class of anthelmintic drugs, target beta-tubulin proteins,
 57 disrupt microtubule formation, and cause nematode death. In parasitic nematode species,
 58 mutations in beta-tubulin genes (e.g., isotype-1 beta-tubulin) are predicted to inhibit BZ binding
 59 and are associated with BZ resistance. Similarly, in the free-living nematode *Caenorhabditis*
 60 *elegans*, mutations in an isotype-1 beta-tubulin ortholog, *ben-1*, are the primary drivers of BZ
 61 resistance. The recurrent association of BZ resistance with beta-tubulins suggests that BZ
 62 resistance is repeatedly caused by mutations in beta-tubulin genes, an example of repeated
 63 evolution of drug resistance across nematode species. To evaluate the hypothesis of repeated
 64 evolution of BZ resistance mediated by beta-tubulin, we identified predicted resistance alleles in
 65 beta-tubulin genes across wild strains from three *Caenorhabditis* species: *C. elegans*,
 66 *Caenorhabditis briggsae*, and *Caenorhabditis tropicalis*. We hypothesized that, if these species
 67 experienced similar selective pressures, they would evolve resistance to BZs by mutations in any
 68 of three beta-tubulin genes (*ben-1*, *tbb-1*, and *tbb-2*). Using high-throughput development assays,
 69 we tested the association of predicted beta-tubulin alleles with BZ resistance. We found that a
 70 heterogeneous set of variants identified in *C. elegans ben-1* were associated with BZ resistance.
 71 In *C. briggsae*, only two variants in *ben-1*, predicted to encode a premature stop codon (W21stop)
 72 and a missense substitution (Q134H), were associated with BZ resistance. In *C. tropicalis*, two
 73 missense variants were identified in *ben-1*, but neither was associated with BZ resistance. *C.*
 74 *briggsae* and *C. tropicalis* might have evolved BZ resistance by mutations in other beta-tubulin
 75 genes, but we found that variants in *tbb-1* or *tbb-2* in these species were not associated with BZ
 76 resistance. Our findings reveal a lack of repeated evolution of BZ resistance across the three
 77 *Caenorhabditis* species and highlight the importance of defining BZ resistance mechanisms
 78 outside of beta-tubulins.

1. INTRODUCTION

Global control of parasitic nematode infections relies on the efficacy of a small arsenal of anthelmintic drugs, including benzimidazoles (BZs) (A. C. Kotze et al. 2020). BZs are a widely used class of anthelmintic drugs that target beta-tubulin proteins (Lubega and Prichard 1990), disrupt microtubule formation (Neff et al. 1983; Laclette, Guerra, and Zetina 1980; Ireland et al. 1979), and lead to nematode death. Although BZs are essential in human and veterinary medicine, resistance is prominent in parasitic nematode populations (Theodorides, Scott, and Lademan 1970; Roos, Kwa, and Grant 1995). To effectively screen for and manage the emergence of BZ resistance, it is necessary to understand the underlying genetics that contribute to the evolution of BZ resistance in nematode species.

Historically, BZ resistance in parasite populations (*i.e.*, *Haemonchus contortus*, *Teladorsagia circumcincta*, and *Trichostrongylus colubriformis*) has been associated with three canonical missense variants (F167Y, E198A, and F200Y) in orthologs of the *H. contortus* *isotype-1* (Silvestre and Cabaret 2002; Ghisi, Kaminsky, and Mäser 2007; de Lourdes Mottier and Prichard 2008; Kwa, Veenstra, and Roos 1994) and *isotype-2 beta-tubulin* genes (A. C. Kotze and Prichard 2016; Avramenko et al. 2019). These three missense variants have been used to track and manage BZ resistance across nematode species globally (Kwa, Veenstra, and Roos 1994; Silvestre and Cabaret 2002; Ghisi, Kaminsky, and Mäser 2007). However, these three missense variants do not explain all of the BZ resistance observed in parasitic nematode populations (Andrew C. Kotze et al. 2014; Krücken et al. 2017). Recently, additional novel missense variants have been associated with BZ resistance (*e.g.*, E198I, E198K, E198T, E198stop, and Q134H) (Venkatesan et al. 2023; Mohammedsalih et al. 2020). The three canonical missense variants, along with novel missense variants of these parasitic nematode beta-tubulin alleles, conferred BZ resistance when introduced in the free-living nematode *Caenorhabditis elegans* (Dilks et al. 2020, 2021). Unlike parasitic species, *C. elegans* wild strains

have a heterogeneous set of variants in one of the *isotype-1 beta-tubulin* orthologs, *ben-1*, responsible for much of BZ resistance in the species (Hahnel et al. 2018). The recurrent association of BZ resistance with alleles predicted to impact beta-tubulin function across nematode species suggests that BZ resistance repeatedly evolves by standing variation or recurrent mutations in beta-tubulin genes and provides compelling evidence to predict the emergence of BZ resistance across nematode species.

Repeated evolution is the development of a similar phenotype (e.g., BZ resistance) and genotype (e.g., the same beta-tubulin alleles confer BZ resistance) in response to similar environmental pressures (e.g., BZ exposure) (Cerca 2023). Because parasitic and free-living nematodes have evolved BZ resistance by variants in conserved beta-tubulin genes, we hypothesized that nematodes acquire BZ resistance by repeated variants or mutations in beta-tubulin genes. However, this hypothesis is difficult to test directly in parasitic nematode species because of their host-dependent life cycles, poorly annotated reference genomes, and limited molecular and genetic tools (Hahnel et al. 2020; Mariene and Wasmuth 2025). By contrast, the availability of high-quality genomic data for hundreds of wild strains (Crombie et al. 2023) and the laboratory tractability of the free-living *Caenorhabditis* nematode species (*C. elegans*, *Caenorhabditis briggsae*, and *Caenorhabditis tropicalis*) provide an opportunity to test for the repeated evolution of BZ resistance in the *Caenorhabditis* genus.

Using the global natural diversity of *C. elegans*, *C. briggsae*, and *C. tropicalis*, we assessed variation in conserved beta-tubulin genes to identify predicted BZ resistance alleles. We identified single nucleotide variants (SNVs), small insertions or deletions (INDELs), and structural variants (SVs) predicted to impact function in conserved beta-tubulin genes, herein called “high-impact” variants. Because high-impact variants in the *Cel-ben-1* gene are known to account for much of the BZ resistance in *C. elegans*, we first used an established high-throughput larval development assay (HTA) to expose *C. elegans* strains with novel *ben-1* variants to the prominently used BZ, albendazole (ABZ), to identify new resistance alleles. We found eight novel

variants in *Cel-ben-1*, of which six were associated with ABZ resistance in the HTAs, a result that further confirms the role *Cel-ben-1* plays in ABZ resistance. Next, we used the HTA to expose *C. briggsae* and *C. tropicalis* strains with novel *ben-1* variants to ABZ to identify if these two species confer ABZ resistance by the same mechanism as in *C. elegans*. In *C. briggsae*, two of the eleven BEN-1 variants (W21stop and Q134H) were correlated with ABZ resistance. In *C. tropicalis*, neither of the two BEN-1 variants (P80T and R121Q) were associated with ABZ resistance. Because we found a lack of repeated evolution of BZ resistance by variants in *ben-1*, we tested whether *C. briggsae* and *C. tropicalis* evolved BZ resistance by variants in the beta-tubulin genes *tbb-1* and *tbb-2*, but found no high-impact variants in these genes associated with ABZ resistance. Overall, our results indicate a lack of repeated evolution of ABZ resistance across *Caenorhabditis* species, suggesting that beta-tubulin cannot underlie BZ resistance across all nematode species.

2. MATERIALS AND METHODS

2.1 Identification of beta-tubulin loci

Amino acid sequences for all six *C. elegans* beta-tubulin proteins were obtained from WormBase (WS283) (Sternberg et al. 2024) and used as queries in a BLASTp search (Version 2.12.0) (Camacho et al. 2009). The search was performed against protein sequence databases constructed using gene models for *C. briggsae* (Moya et al. 2023) and *C. tropicalis* (Noble et al. 2021). To construct the protein sequence database, we extracted gene model transcript features from the gene feature file with *gffread* (Version 0.9.11) (Pertea and Pertea 2020) and processed them using the *makeblastdb* function from BLAST (Version 2.12.0). From the BLASTp search, we identified *C. briggsae* and *C. tropicalis* protein sequences with the highest percent identity (PID) to each *C. elegans* beta-tubulin protein. Only protein sequences with the highest PID in both searches were considered orthologs (**S1 Table**). For some *C. elegans* beta-tubulin orthologs,

multiple *C. briggsae* or *C. tropicalis* gene models contained multiple splice isoforms. All gene models for all beta-tubulin transcripts were manually inspected, and isoforms that were not fully supported by short-read RNA sequencing data were removed.

2.2 Single nucleotide variant (SNV) and indel calling and annotation

To identify single nucleotide variants (SNVs) or indels (insertions and deletions) in the beta-tubulin genes across the selfing *Caenorhabditis* species, we used the Variant Annotation Tool from the *Caenorhabditis* Natural Diversity Resource (CaeNDR) (Release IDs: *C. elegans* - 20231213, *C. briggsae* - 20240129, *C. tropicalis* - 20231201) (Crombie et al. 2023). The identified SNVs and indels (**S2, S3, and S4 Tables**) included small insertions and deletions, frameshifts, altered stop and start codons, nonsynonymous changes, and splice variants.

2.3 Structural variant (SV) calling and annotation

Structural variant (SV) calling was performed using *DELLY* (Version 0.8.3), a SV caller optimized to detect large insertions, deletions, and other complex structural variants such as inversions, translocations, and duplications in paired-end short-read alignments (Rausch et al. 2012) and shown to perform well on *C. elegans* short-read sequence data (Lesack et al. 2022). SVs that overlapped with beta-tubulin genes were extracted using *bcftools* (Version 1.10.1) (Danecek et al. 2021). Insertions, deletions, inversions, and duplications that passed the *DELLY* (Version 0.8.3) default quality threshold (greater than three supporting read pairs with a median MAPQ > 20), filtered to high-quality genotypes (genotype quality > 15), and had at least one alternative allele were retained. For complex variants (inversions and duplications), the identification of at least one split-read pair was required (variants flagged as a precise SV by *DELLY*). To validate SVs that passed quality filtering, each SV was manually inspected for breakpoints in the raw-read alignments (*Wally*, Version 0.5.8) and for impacts on the beta-tubulin coding sequence (CaeNDR Genome Browser) (Crombie et al. 2023) (**S5 Table**). We retained

SVs where raw read alignments suggested that the SV impacted the beta-tubulin coding sequence. We compared the *Cel-ben-1* SVs called by *DELLY* to those SVs identified previously (Hahnel et al. 2018). *DELLY* successfully recalled structural variants in several strains, including deletions in JU751, JU830, JU1395, JU2582, JU2587, JU2593, JU2829, and QX1233, as well as an inversion in MY518. However, *DELLY* did not detect a previously reported transposon insertion in strain JU3125. To assess if other SVs could have been missed by *DELLY*, we manually inspected the read alignments for all strains that had not been previously phenotyped to check if any other SVs were not detected by *DELLY*. We confirmed the presence of novel *Cel-ben-1* SVs in multiple strains, including deletions in ECA706 and NIC1832, and a duplication in NIC1107. Additionally we identified a previously undetected deletion in JU4287.

2.4 Association of low *Cel-ben-1* expression with ABZ response

Two previous assays measured developmental responses of wild *C. elegans* strains after ABZ exposure (Hahnel et al. 2018; Shaver et al. 2023). For 180 of these wild *C. elegans* strains, expression levels (transcripts per million estimates [TPM]) were collected at the young-adult stage (Zhang et al. 2022). Of the 180 wild *C. elegans* strains, 105 strains were shared between both assays. A linear model was built using the *lm* function in R to account for assay effects. Subsequently, the residuals of the linear model were used to normalize previous measures of ABZ response. We evaluated the linear fit between each strain's expression of *ben-1*, *tbb-1*, or *tbb-2* and the developmental delay following ABZ exposure.

2.5 Phylogenetic analysis

We characterized the relatedness of isotype reference strains (genetically unique strains) with beta-tubulin variants using species trees downloaded from CaeNDR and generated by the 'post-gatk-nf' pipeline (<https://github.com/AndersenLab/post-gatk-nf>) (Release IDs: *C. elegans* - 20231213, *C. briggsae* - 20240129, *C. tropicalis* - 20231201) (Crombie et al. 2023). Briefly, the

trees were generated with high-quality SNVs in isotype reference strains retained in the hard-filtered variant call format (VCF) file. *vcf2phyliip* (Ortiz 2019) and the *bioconvert* (Caro et al. 2023) function *phyliip2stockholm* were used to prepare inputs for *quicketree*, which was used to construct a tree using a neighbor-joining algorithm (Saitou and Nei 1987). All versions of these software can be accessed from the ‘post-gatk’ docker container (<https://hub.docker.com/r/andersenlab/tree>), used by the ‘post-gatk-nf’ pipeline. We visualized the trees for each species using the *ggtree* function from the *ggtree* (v3.6.2) R package (Yu 2020).

2.6 Strain selection and maintenance

Fifteen *C. elegans* strains, 38 *C. briggsae* strains, and seven *C. tropicalis* strains from the CaeNDR (Crombie et al. 2023) were used in this study (**S2, S3, and S4 Tables**). Isolation details for each strain are included in CaeNDR. For each species, we selected strains that had unique high-impact consequences (SNV or SV) in *ben-1*, *tbb-1*, or *tbb-2* that had not been previously phenotyped. Strains with high-impact consequences in a beta-tubulin gene are herein referred to as “predicted resistant” strains. Strains with no high-impact consequences in beta-tubulin genes that were closely related to predicted resistant strains were also included for *C. briggsae* and *C. tropicalis* and herein classified as “predicted susceptible” strains (**S3 and S4 Figures**) (**Table S6**). The reference strain(s) for all three species was included. Finally, for *C. elegans*, a strain with a loss of *ben-1* (ECA882) was also included.

Before measuring ABZ responses, *C. elegans* and *C. briggsae* animals were maintained at 20°C and *C. tropicalis* animals were maintained at 25°C. All animals were maintained on 6 cm plates with modified nematode growth medium (NGMA), which contains 1% agar and 0.7% agarose to prevent animals from burrowing (Andersen et al. 2014). The NGMA plates were seeded with the *Escherichia coli* strain OP50 as a nematode food source. All strains were grown for three generations without starvation on NGMA plates before anthelmintic exposure to reduce the transgenerational effects of starvation stress (Andersen et al. 2015).

2.7 Nematode food preparation for NGMA 6 cm plates

A batch of OP50 *E. coli* was grown and used as a nematode food source for NGMA plates. A frozen stock of OP50 *E. coli* was streaked onto a 10 cm Luria-Bertani (LB) agar plate and incubated overnight at 37°C. The following day, a single bacterial colony was transferred into two culture tubes that contained 5 mL of 1x LB. The starter cultures and two negative controls (1X LB without *E. coli*) were incubated for 18 hours at 37°C shaking at 210 rpm. The OD₆₀₀ value of the starter cultures were measured using a spectrophotometer (BioRad, SmartSpec Plus) to calculate how much starter culture was needed to inoculate a 1 L culture at an OD₆₀₀ value of 0.005. For each assay, one culture containing one liter of pre-warmed 1X LB inoculated with the starter culture grew for approximately 4 - 4.5 hours at 37°C at 210 rpm to an OD₆₀₀ value between 0.45 and 0.6. Cultures were transferred to 4°C to slow growth. OP50 was spotted on NGMA test plates (two per culture) and grown at 37°C overnight to ensure no contamination.

2.8 Nematode food preparation for HTAs

One batch of HB101 *E. coli* was used as a nematode food source for all HTAs in this study. A frozen stock of HB101 *E. coli* was streaked onto a 10 cm LB agar plate and incubated overnight at 37°C. The following day, a single bacterial colony was transferred into three culture tubes that contained 5 mL of 1x Horvitz Super Broth (HSB). The starter cultures and two negative controls (1X HSB without *E. coli*) were incubated for 18 hours at 37°C shaking at 180 rpm. The OD₆₀₀ value of the starter cultures were measured using a spectrophotometer (BioRad, SmartSpec Plus) to calculate how much starter culture was needed to inoculate a 1 L culture at an OD₆₀₀ value of 0.001. A total of four cultures each containing 1 L of pre-warmed 1X HSB inoculated with the starter culture grew for 15 hours at 37°C while shaking at 180 rpm. After 15 hours, flasks were removed from the incubator and transferred to 4°C to slow growth. The 1X HSB was removed from the cultures by performing three rounds of centrifugation, where the

supernatant was removed, and the bacterial cells were pelleted. Bacterial cells were washed with K medium, resuspended in K medium, pooled, and transferred to a 2 L glass beaker. The OD₆₀₀ value of the bacterial suspension was measured and diluted to a final concentration of OD₆₀₀ 100 with K medium, aliquoted to 15 mL conicals, and stored at -80°C for use in the HTAs.

2.9 ABZ dose-response assays for *C. briggsae* and *C. tropicalis*

Because ABZ response has been minimally characterized in *C. briggsae* (Zamanian et al. 2018) and has not yet been described in *C. tropicalis*, we first measured dose-response curves for both species after exposure to ABZ to assess developmental delay. Before performing HTAs, ABZ (Sigma-Aldrich, Catalog # A4673-10G) stock solutions were prepared in dimethyl sulfoxide (DMSO) (Fisher Scientific, Catalog # D1281), aliquoted, and stored at -20°C for use in the assays. For the dose-response assays, animals were exposed to ABZ at the following concentrations (μM): 0 (1% DMSO), 0.12, 0.23, 0.47, 0.94, 1.88, 3.75, 7.5, 15, 30, 60, and 120. Animals were allowed to develop in the presence of ABZ as described in HTAs to assess nematode development.

Dose-response model estimation and statistics were performed as described previously (Widmayer et al. 2022; Shaver et al. 2023). Briefly, a four-parameter log-logistic dose-response curve was fit independently for a genetically diverse set of 11 *C. briggsae* strains (**S5 Figure**) and seven *C. tropicalis* strains (**S6 Figure**), where normalized median animal length was used as a metric for phenotypic response (see *Methods*). For each strain-specific dose-response model, slope (*b*) and concentration (*e*) were estimated with strain as a covariate. We calculated EC₁₀ as we have previously found EC₁₀ response to be more heritable than half maximal effective concentration (EC₅₀) estimates and were therefore used in our analysis (Shaver et al. 2023; Widmayer et al. 2022). A dosage of 30 μM ABZ was closest to the EC₁₀ for *C. briggsae* and *C. tropicalis*, consistent with ABZ concentrations used in past *C. elegans* assays (Dilks et al. 2020, 2021; Shaver et al. 2024) and in all HTAs in this study.

2.10 HTAs to assess nematode development

Populations of each strain were amplified and bleach-synchronized in three independent assays. Independent bleach synchronizations controlled for variation in embryo survival and subsequent effects on developmental rates. After bleach synchronization, approximately 30 embryos were dispensed into the wells of a 96-well microplate in 50 μ L of K medium. Forty-eight wells were prepared per bleach for each strain. Each 96-well microplate was prepared, labeled, and sealed using gas-permeable sealing films (Fisher Scientific, Catalog # 14-222-043). Plates were placed in humidity chambers to incubate for 24 hours at 20°C while shaking at 170 rpm (INFORS HT Multitron shaker). After 24 hours, every plate was inspected to ensure that all embryos hatched and animals were developmentally arrested at the first larval (L1) stage so all strains started each assay at the same developmental stage. Next, food was prepared to feed the developmentally arrested L1 animals using the required number of OD₆₀₀100 HB101 aliquots (see *Nematode food preparation for HTAs*). The HB101 aliquots were thawed at room temperature, combined into a single conical tube, and diluted to an OD₆₀₀30 with K medium. To inhibit further bacterial growth and prevent contamination, 150 μ L of kanamycin was added to the HB101. An aliquot of 100 μ M ABZ stock solution was thawed at room temperature and added to an aliquot of OD₆₀₀30 K medium at a 3% volume/volume ratio. Next, 25 μ L of the food and ABZ mixture was transferred into the appropriate wells of the 96-well microplates to feed the arrested L1 animals at a final HB101 concentration of OD₆₀₀10 and expose L1 animals to ABZ. Immediately afterward, the 96-well microplates were sealed using a new gas permeable sealing film, returned to the humidity chambers, and incubated for 48 hours at 20°C (*C. elegans* and *C. briggsae*) or 25°C (*C. tropicalis*) while shaking at 170 rpm. After 48 hours (*C. elegans* and *C. briggsae*) or 42 hours (*C. tropicalis*) of incubation and shaking in the presence of food and either DMSO or ABZ, the 96-well microplates were removed from the incubator and treated with 50 mM sodium azide in M9 for 10 minutes to paralyze and straighten nematodes. After 10 minutes, images of nematodes in

the microplates were immediately captured using Molecular Devices ImageXpress Nano microscope (Molecular Devices, San Jose, CA) using a 2X objective. The ImageXpress Nano microscope acquires brightfield images using a 4.7 megapixel CMOS camera and stores images in a 16-bit TIFF format. The images were used to quantify the development of nematodes in the presence of DMSO or ABZ as described below (see *High-throughput imager assays [HTA] data collection and cleaning*). A full step-by-step protocol for the HTA has been deposited in protocols.io.

2.11 HTA data collection and data cleaning

CellProfiler (Version 24.10.1) was used to characterize and quantify biological data from the image-based assays. Custom software packages designed to extract animal measurements from images collected on the Molecular Devices ImageXpress Nano microscope were previously described (Nyaanga et al. 2021). *CellProfiler* modules and *Worm Toolbox* were developed to extract morphological features of individual animals from images from the HTA (Wählby et al. 2012). Worm model estimations and custom *CellProfiler* pipelines were written using the *WormToolbox* in the GUI-based instance of *CellProfiler* (Widmayer et al. 2022). Next, a Nextflow pipeline (Version 24) was written to run command-line instances of *CellProfiler* in parallel on the Rockfish High-Performance Computing Cluster (Johns Hopkins University). The *CellProfiler* workflow can be found at <https://github.com/AndersenLab/cellprofiler-nf>. The custom *CellProfiler* pipeline generates animal measurements by using four worm models: three worm models tailored to capture animals at the L4 larval stage, in the L2 and L3 larval stages, and the L1 larval stage, as well as a “multi-drug high dose” (MDHD) model, to capture animals with more abnormal body sizes caused by extreme anthelmintic responses. These measurements comprised our raw dataset. Two *C. briggsae* strains (NIC1052 and VX34) were not fully paralyzed and straightened at the time of imaging, which created some misclassification of animal measurements. Thus, the animal lengths for strains NIC1052 and VX34 measured by *CellProfiler* are shorter than the actual

animal lengths. However, the difference discrepancy in animal lengths does not affect the classification of a strain as resistant or sensitive to ABZ. Data cleaning and analysis steps were performed using a custom R package, *easyXpress* (Version 2.0) (Nyaanga et al. 2021) and followed methods previously reported (Shaver et al. 2024). All analyses were performed using the R statistical environment (Version 4.2.1) unless stated otherwise.

3. RESULTS

3.1 In three free-living *Caenorhabditis* species, strains with high-impact variants in beta-tubulin genes or low levels of beta-tubulin expression are predicted to confer ABZ resistance

To investigate if the mechanisms of ABZ resistance are repeated across the three selfing *Caenorhabditis* species, we identified “predicted resistant” strains that have high-impact variants (*i.e.*, SNVs, INDELs, or SVs) (Crombie et al. 2023) likely to affect the function of the five conserved beta-tubulin genes (*i.e.*, *ben-1*, *tbb-1*, *tbb-2*, *tbb-4*, and *mec-7*) (see *Methods*). Although ABZ response has been highly characterized and associated with mutations in *ben-1* across *C. elegans* wild strains (Hahnel et al. 2018; Shaver et al. 2024), less is known about how *C. briggsae* (Zamanian et al. 2018) and *C. tropicalis* respond to ABZ exposure. To minimize the amount of genetic variation outside of the beta-tubulin that could affect the BZ response, we also selected *C. briggsae* and *C. tropicalis* strains that lacked high-impact variants in beta-tubulin genes but were closely related to predicted resistant strains (**S3 and S4 Figures**). The strains with no high-impact variants in any beta-tubulin gene were classified as “predicted susceptible”.

In a set of 611 *C. elegans* strains (Crombie et al. 2023), we identified 65 strains with 33 unique high-impact variants in *Cel-ben-1* (**Table S2**). Of the 33 unique high-impact variants, 24 were previously phenotyped and 23 were associated with ABZ resistance (**S1A Figure**) (Hahnel et al. 2018; Shaver et al. 2024). Here, we report eight novel high-impact variants predicted to impact *Cel-BEN-1* (E3stop, Y50C, P80S, VDN113N, frameshift 319, frameshift 368, stop445S,

and a duplication) that have not previously associated with ABZ resistance in nematodes (**Table S2**). The missense variants predicted to encode Y50C and P80S *Cel*-BEN-1 substitutions have been found in fungus and human beta-tubulins, respectively (Qiu et al. 2011; Dingerdissen et al. 2018). We found no strains with high-impact variants predicted to impact *Cel*-TBB-1 or *Cel*-TBB-2.

Previously, low *ben-1* expression was correlated with ABZ resistance in *C. elegans* strains (Zhang et al. 2022), providing another parameter to investigate BZ resistance across the species. We evaluated if beta-tubulin expression levels were predictive of ABZ responses in *C. elegans*. We used a linear regression model to assess the relationship between the expression of *Cel*-*ben-1*, *Cel*-*tbb-1*, or *Cel*-*tbb-2* (Zhang et al. 2022) and ABZ responses in 180 wild strains (Hahnel et al. 2018; Shaver et al. 2024). We found a significant negative relationship between *Cel*-*ben-1* expression (≥ 3.75 TPM) and ABZ resistance (p -value = 5.16×10^{-16} , $r^2 = 0.344$) (**S1 Figure**). Importantly, 19 of the 23 strains with low *ben-1* expression also had high-impact variants in *ben-1* (SVs, frameshifts, or stop/start altering variants). Next, we identified strains with no high-impact variants in *ben-1* but had low *ben-1* expression that we predicted to cause ABZ resistance. Using these parameters, we found one strain, JU1581, with low *Cel*-*ben-1* expression to test for ABZ resistance and to further define the role *Cel*-*ben-1* plays in ABZ resistance. No significant relationship was found between *Cel*-*tbb-2* and *Cel*-*tbb-1* expression and ABZ response (**S2 Figure**).

In a set of 641 *C. briggsae* strains, we identified 22 strains with eight unique high-impact variants in *Cbr*-*ben-1* (**Table S3**). The eight unique high-impact variants in *Cbr*-BEN-1 included seven amino acid substitutions (V91I, Q94K, D128E, Q134H, S218L, M299V, and R359H) and one premature stop codon (W21stop). Outside of *Cbr*-BEN-1, we identified missense variants in *Cbr*-TBB-1 (T35A, A275T, L377I, and V64I) and *Cbr*-TBB-2 (E441A) predicted to disrupt function. All *C. briggsae* predicted resistant strains, predicted susceptible strains, and the reference strains (AF16 and QX1410) (Stevens et al. 2022; Moya et al. 2023) were phenotyped for ABZ response

(S1 Figure).

In a set of 518 *C. tropicalis* strains, we identified two strains with unique high-impact variants in *Ctr*-BEN-1 (**Table S4**) (P80T and R121Q). Outside of *Ctr*-BEN-1, we identified one missense variant in *Ctr*-TBB-2 (N89S) and no high-impact variants in *Ctr*-TBB-1. All *C. tropicalis* predicted resistant strains, three predicted susceptible strains, and the reference strain (NIC58) (Luke et al. 2021) were phenotyped for ABZ response.

3.2 Natural allelic variation in *ben-1* is associated with ABZ resistance in *C. elegans*

We performed image-based HTAs across all *C. elegans* wild strains with novel high-impact variants in *ben-1* (*i.e.*, predicted resistant strains) to measure nematode length (a proxy for larval development) in drug (ABZ) and control (DMSO) conditions (see *Methods*). The canonical laboratory strain N2 was included as an ABZ-susceptible control (Dilks et al. 2020, 2021; Shaver et al. 2024) and a strain with the loss of *ben-1* in the N2 strain background (ECA882) was included as an ABZ-resistant control. Previously, two high-impact variants (frameshift at position 299 and a deletion) were associated with ABZ resistance (Hahnel et al. 2018; Dilks et al. 2020). With expanded sampling, we identified two novel strains with the same frameshift at position 299 (JR4305) and the same deletion (NIC1832) in *Cel*-*ben-1*. Therefore, we included JR4305 and NIC1832 to validate the role that a frameshift at position 299 and a deletion play in ABZ resistance. The assay included 48 replicates per strain with 5 to 30 animals per replicate in ABZ or DMSO conditions. The reported nematode length of each strain is the delta between animal lengths in DMSO (**S7 Figure**) and ABZ to obtain normalized animal length and assess drug effects. We classified a strain as resistant to ABZ if its nematode length after ABZ exposure was 75% greater than that of the reference strain (see *Methods*). A longer median animal length (*i.e.*, larger animals) indicated resistance to ABZ, and a shorter median animal length (*i.e.*, smaller animals) indicated susceptibility to ABZ.

As expected, when exposed to ABZ, strains ECA882 (loss of *ben-1*), JR4305 (frameshift

at position 299), and NIC1832 (deletion) all recapitulated a resistance phenotype (nematode length $\geq 75\%$ of the reference strain N2). Of the eight predicted resistant *C.elegans* strains, six displayed a resistance phenotype (**Fig. 1**). All six wild strains showed minimal developmental delays after exposure to ABZ, a phenotype similar to loss of *ben-1* (**Fig. 1**). Of all the assayed predicted resistant strains with variants in *Cel-BEN-1*, three (P80S, stop445S, and low *ben-1* expression) were not classified as resistant to ABZ. The P80S variant might partially alter *ben-1* function, causing a moderate resistance phenotype. In stop445S, the normal stop codon was replaced with a serine, likely allowing translation to continue beyond the termination point. This variant likely does not affect *ben-1* function because position 445 is at the end of the BEN-1 protein. Finally, the strain with low *ben-1* expression (JU1581), which did not have any high-impact variants in *ben-1*, was sensitive to ABZ, indicating that the selected threshold of *ben-1* expression (≥ 3.75 TPM) still retained strains with adequate *ben-1* function.

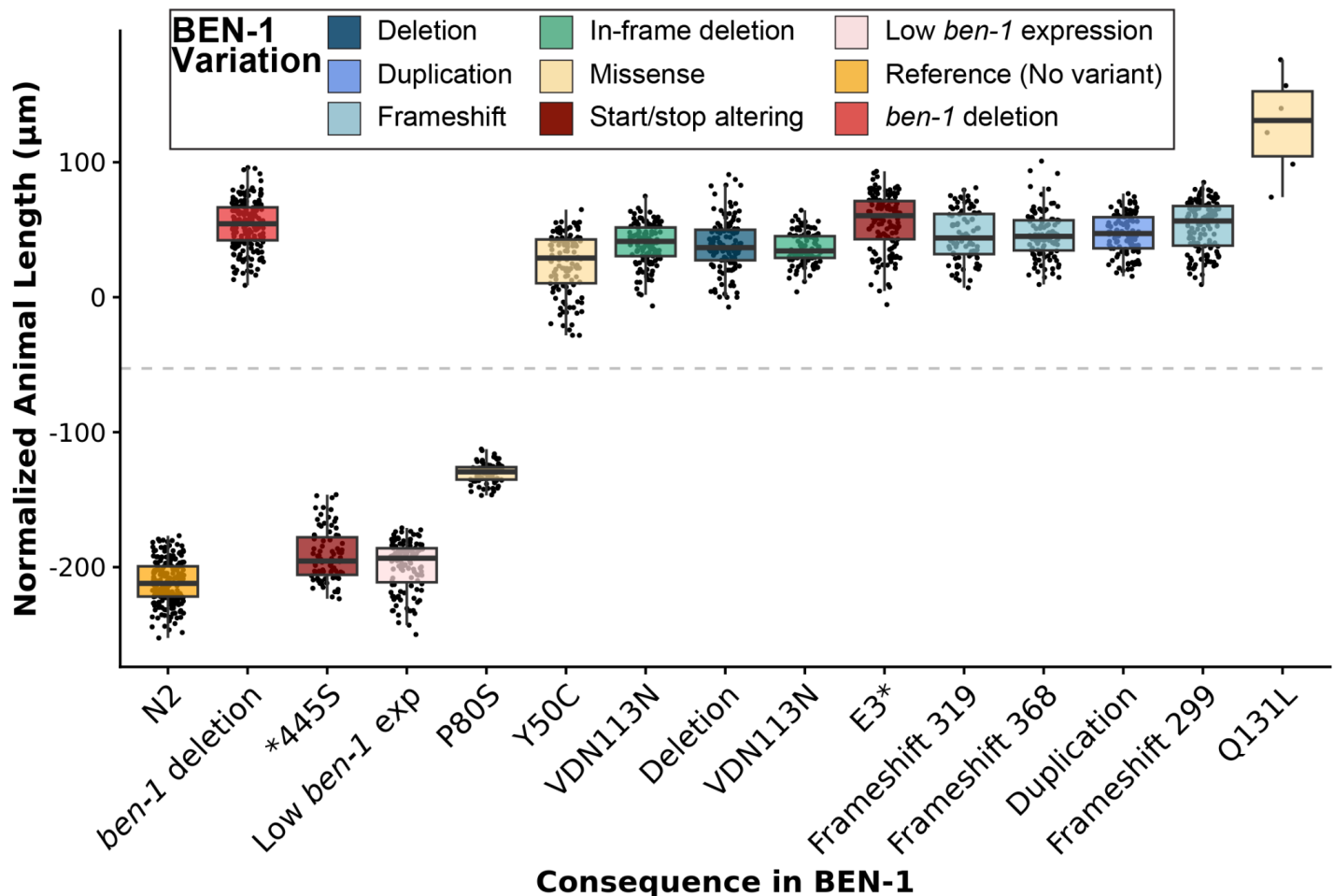


Figure 1. High-throughput assays for each *C. elegans* strain with a high-impact variant in BEN-1 in the presence of albendazole

The regressed median animal length values for populations of nematodes grown in 30 μM albendazole (ABZ) are shown on the y-axis. Each point represents the normalized median animal length value of a well containing approximately 5-30 animals. Strains are sorted by their relative resistance to ABZ based on median animal length. Data are shown as Tukey box plots with the median as a solid horizontal line, and the top and bottom of the box representing the 75th and 25th quartiles, respectively. The top whisker is extended to the maximum point that is within the 1.5 interquartile range from the 75th quartile. The bottom whisker is extended to the minimum point that is within the 1.5 interquartile range from the 25th quartile. Strains are colored by beta-tubulin variant status. The gray dashed line marks the resistance threshold (nematode length ≥ 75% of the reference strain N2).

3.3 Rare missense and nonsense variants in *Cbr-ben-1* are associated with ABZ resistance,

whereas high-impact variants in *Ctr-ben-1* are not

We performed HTAs on 11 predicted resistant *C. briggsae* strains with eight unique variants in *Cbr-ben-1* and on 13 predicted susceptible strains in DMSO (**S8 Figure**) and ABZ conditions. We found that in *C. briggsae*, only two variants in BEN-1 (Q134H and W21stop) conferred ABZ resistance (nematode length $\geq$ 75% of the reference strain AF16) (**Fig. 2**). The Q134H amino acid change has been associated with ABZ resistance in *Ancylostoma caninum* and validated in *C. elegans* (Venkatesan et al. 2023). An early stop gain at position 21 is predicted to cause the premature termination of protein synthesis and *Cbr-ben-1* loss-of-function (LoF). Of all the assayed strains with predicted resistance variants in *Cbr-BEN-1*, six (V91I, Q94K, D128E, S218L, M299V, and R359H) were not resistant to ABZ. Overall, because only one of the seven missense variants was associated with ABZ resistance, we could not reliably predict how a strain responded to ABZ based on the presence of a missense variant alone. These results indicate that ABZ resistance associated with *Cbr-ben-1* variants is uncommon and that accurately predicting ABZ resistance in *C. briggsae* requires additional factors beyond *ben-1* variation.

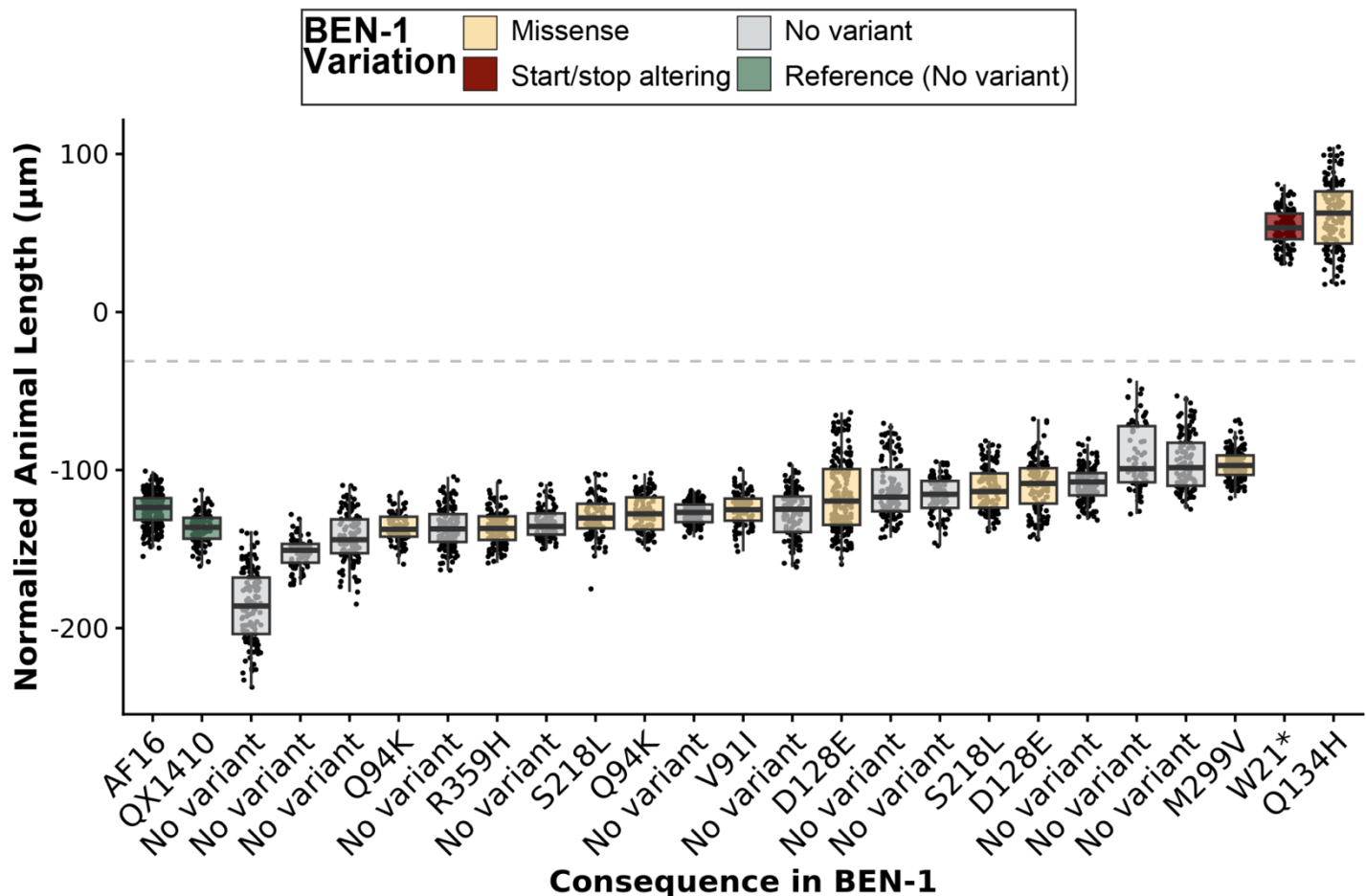
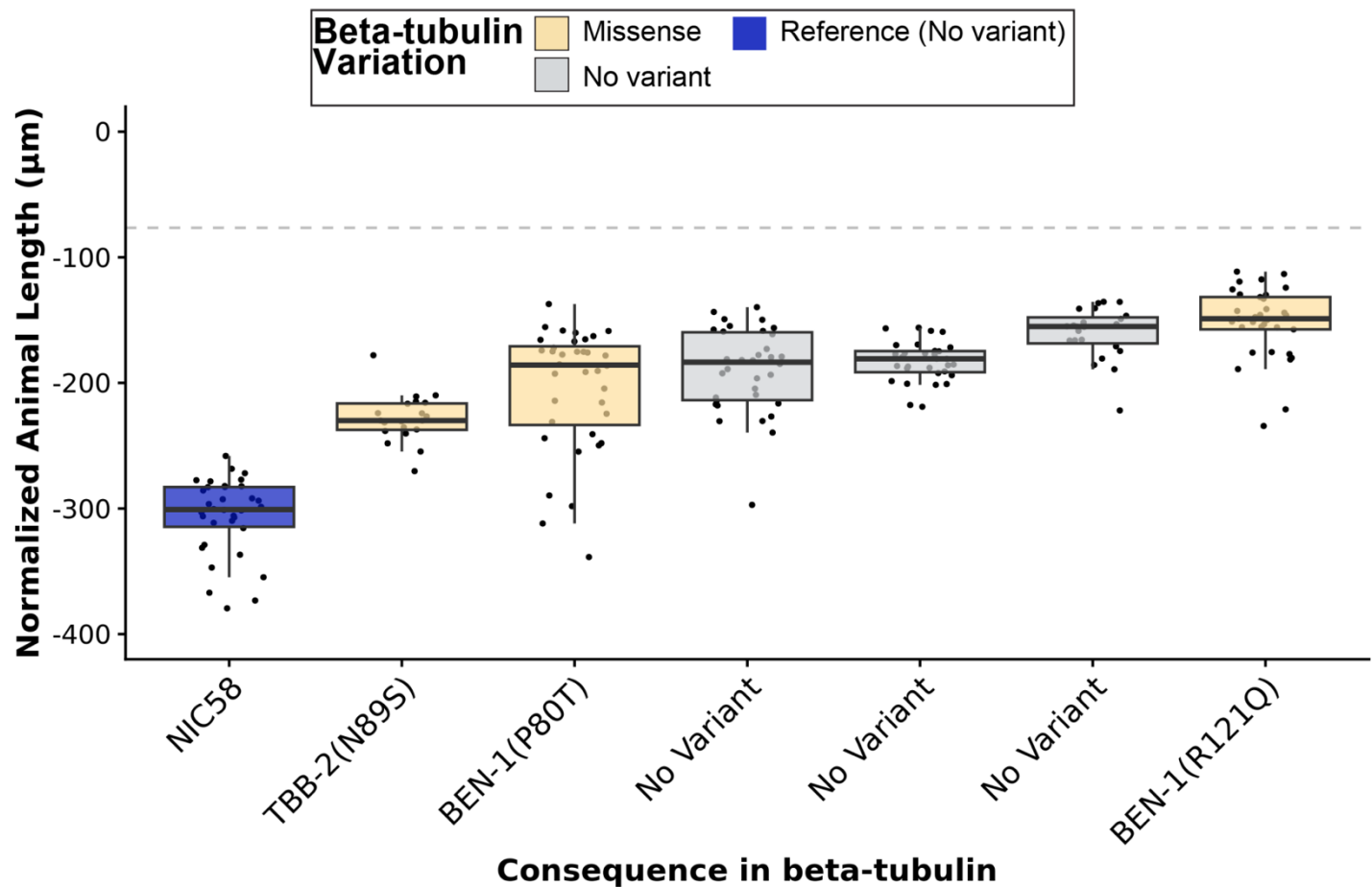


Figure 2. High-throughput assays for each *C. briggsae* strain with a high-impact variant in BEN-1 and paired predicted susceptible strains in the presence of albendazole

The regressed median animal length values for populations of nematodes grown in 30 μM albendazole (ABZ) are shown on the y-axis. Each point represents the normalized median animal length value of a well containing approximately 5-30 animals. Strains are sorted by their relative resistance to ABZ based on median animal length. Data are shown as Tukey box plots with the median as a solid horizontal line, and the top and bottom of the box representing the 75th and 25th quartiles, respectively. The top whisker is extended to the maximum point that is within the 1.5 interquartile range from the 75th quartile. The bottom whisker is extended to the minimum point that is within the 1.5 interquartile range from the 25th quartile. Strains are colored by beta-tubulin variant status. The gray dashed line marks the resistance threshold (nematode length ≥ 75% of the reference strain AF16).

We performed HTAs on two *C. tropicalis* predicted resistant strains with high-impact variants in *Ctr*-BEN-1 (P80T and R121Q) and two predicted susceptible strains in DMSO (**S9 Figure**) and ABZ conditions. We found that both predicted resistant strains had a similar ABZ response as the predicted susceptible strains, where no strains met the resistance threshold (nematode length $\geq$ 75% of the reference strain NIC58), indicating that P80T and R121Q are not associated with ABZ resistance (**Fig. 3**). These results indicate that *Ctr*-BEN-1 variants do not confer ABZ resistance and that predicting ABZ resistance in *C. tropicalis* requires considering factors beyond *ben-1* variation. Additionally, to evaluate if the predicted resistant variants in *Cbr*-BEN-1 or *Ctr*-BEN-1 were less deleterious compared to *Cel*-BEN-1, we assessed the BLOSUM and Grantham scores of each high-impact variant (Crombie et al. 2023). The BLOSUM and Grantham scores measure the evolutionary likelihood of observing a particular missense substitution, and we found that these scores do not accurately predict ABZ resistance (**Table S7**). We performed a linear regression analysis of strain ABZ responses by the BLOSUM or Grantham scores of the beta-tubulin variant in each strain. We found no significant relationship between either BLOSUM or Grantham scores and ABZ response (**S10, S11, S12 Figures**). For example, the only missense substitution to cause ABZ resistance in *Cbr*-BEN-1, Q134H, had substantially less radical BLOSUM and Grantham scores (BLOSUM: 0, Grantham: 24) compared to other missense substitutions, such as S218L (BLOSUM: -2, Grantham: 145).



484 **Figure 3. High-throughput assays for each *C. tropicalis* strain with a high-impact variant**
 485 **in BEN-1 or TBB-2 and paired predicted susceptible strains in the presence of**
 486 **albendazole**

487 The regressed median animal length values for populations of nematodes grown in 30 μM
 488 albendazole (ABZ) are shown on the y-axis. Each point represents the normalized median animal
 489 length value of a well containing approximately 5-30 animals. Strains are sorted by their relative
 490 resistance to ABZ based on median animal length. Data are shown as Tukey box plots with the
 491 median as a solid horizontal line, and the top and bottom of the box representing the 75th and
 492 25th quartiles, respectively. The top whisker is extended to the maximum point that is within the
 493 1.5 interquartile range from the 75th quartile. The bottom whisker is extended to the minimum
 494 point that is within the 1.5 interquartile range from the 25th quartile. Strains are colored by beta-
 495 tubulin variant status. The gray dashed line marks the resistance threshold (nematode length ≥
 496 75% of the reference strain NIC58).

3.4 Natural allelic variants in *tbb-1* and *tbb-2* are not associated with ABZ resistance across three free-living *Caenorhabditis* species

To identify if high-impact variants in other beta-tubulin genes beyond *ben-1* play a role in ABZ resistance in the three free-living *Caenorhabditis* species, we categorized variants in the two genes most highly expressed in *C. elegans* (*tbb-1* and *tbb-2*). Because the loss of *tbb-1* or *tbb-2* is deleterious in *C. elegans* (Collins et al. 2024), LoF mutations in either gene would likely be purged by purifying selection from natural populations. Therefore, as expected, we found no variants in TBB-1 or TBB-2 in *C. elegans* wild strains. Unlike *C. elegans*, *C. briggsae* had four missense variants in TBB-1 (T35A, V64I, A275T, and L377I) and one missense variant in TBB-2 (E441A). In *C. tropicalis*, we identified one rare missense variant in TBB-2 (N89S). Performing the same HTA to assess nematode development as described previously, we found that none of the missense variants in TBB-1 or TBB-2 in *C. briggsae* (**Fig. 4**) or *C. tropicalis* conferred ABZ resistance (**Fig. 3**). These results suggest that missense variants in TBB-1 and TBB-2 are not associated with ABZ resistance in *Caenorhabditis* species.

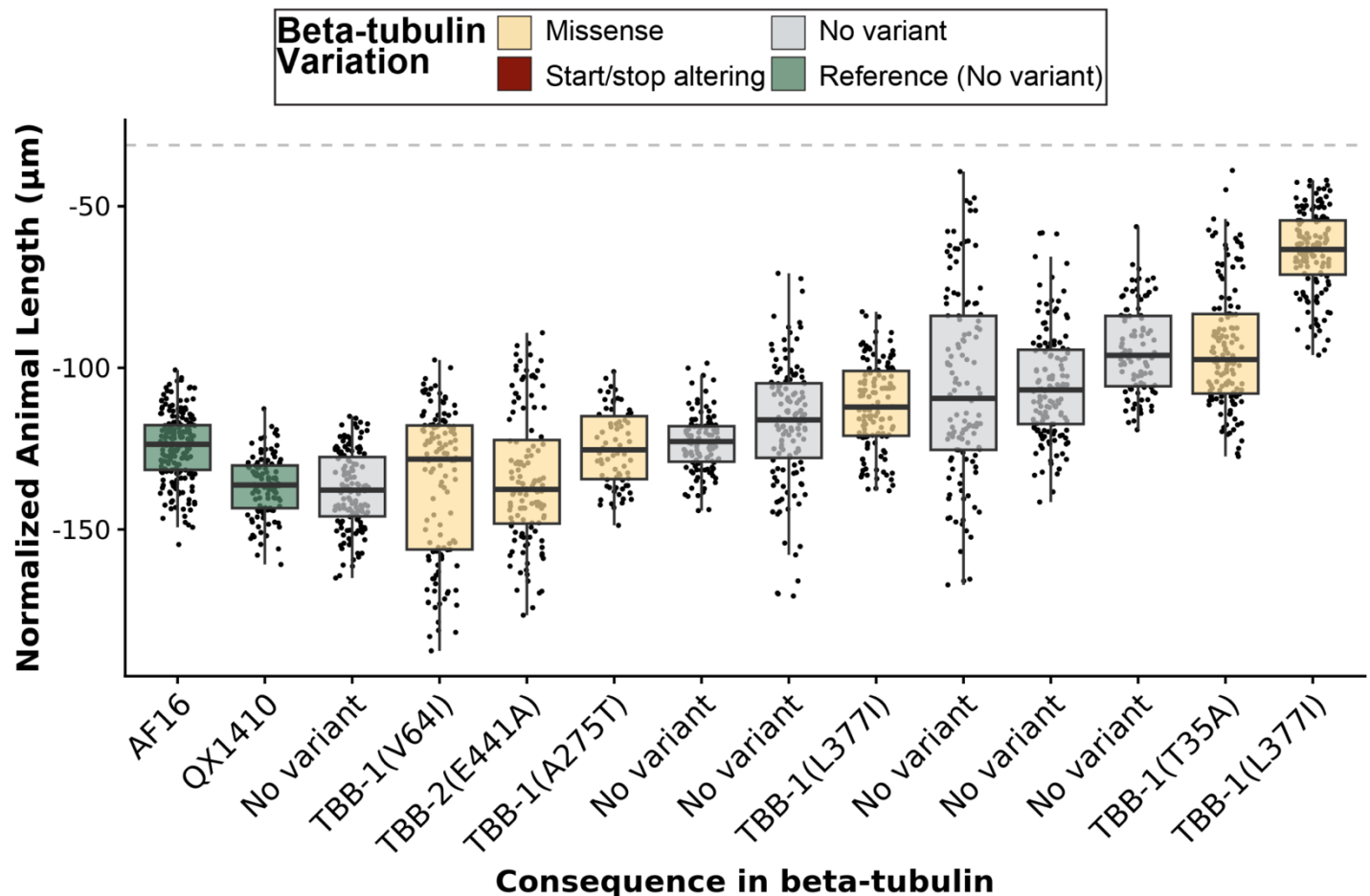


Figure 4. High-throughput assays for each *C. briggsae* strain with a high-impact variant in TBB-1 or TBB-2 in the presence of albendazole

The regressed median animal length values for populations of nematodes grown in 30 μM albendazole (ABZ) are shown on the y-axis. Each point represents the normalized median animal length value of a well containing approximately 5-30 animals. Strains are sorted by their relative resistance to ABZ based on median animal length. Data are shown as Tukey box plots with the median as a solid horizontal line, and the top and bottom of the box representing the 75th and 25th quartiles, respectively. The top whisker is extended to the maximum point that is within the 1.5 interquartile range from the 75th quartile. The bottom whisker is extended to the minimum point that is within the 1.5 interquartile range from the 25th quartile. Strains are colored by beta-tubulin variant status. The gray dashed line marks the resistance threshold (nematode length ≥ 75% of the reference strain AF16.)

3.5 High-impact variants in beta-tubulin genes are rare and not enriched for geography or substrate

To better understand the evolution of BZ resistance alleles in three *Caenorhabditis* species, we assessed the population-wide frequencies of each beta-tubulin consequence, along with the geographic location and substrate of where each strain was isolated from nature. First, to determine the prevalence of high-impact variants in the five conserved beta-tubulin genes (*tbb-1*, *tbb-2*, *mec-7*, *tbb-4*, and *ben-1*) across *Caenorhabditis* species, we quantified the frequency of each consequence (deletion, duplication, frameshift, in-frame deletion, inversion, missense, splice donor, and start/stop altering) in each species. Despite extensive global sampling of the three *Caenorhabditis* species (*C. elegans*: 611 strains, *C. briggsae*: 641 strains, *C. tropicalis*: 518 strains), we found that all consequences in all three *Caenorhabditis* species were rare (< 0.05% of all strains in a species) (**Fig. 5**). The most variation in consequences was found in *Cel*-BEN-1, where deletions, frameshifts, in-frame deletions, inversions, missense, stop/start altering variants, and a duplication were found. However, we found one missense variant in *Cel*-TBB-4 and several missense variants and a splice donor in *Cel*-MEC-7. *Cel-tbb-6* is unique to *C. elegans* and was therefore not assessed. Overall, *C. elegans* has acquired the most diverse set of consequences in BEN-1 and has conferred BZ resistance primarily by variation in *Cel-ben-1*. In *C. briggsae*, we found 21 missense amino-acid substitutions and a single strain with a start/stop altering consequence in *Cbr*-BEN-1. In both *Cbr*-TBB-1 and *Cbr*-TBB-2, we identified rare missense consequences. Finally, we found nine strains with splice variants in *Cbr*-TBB-4 and one strain with a missense variant. In *C. tropicalis*, we found missense consequences in all beta-tubulin genes except *tbb-1*. These findings highlight that *C. elegans* has the most diverse set of beta-tubulin variants, particularly in *ben-1*, reinforcing its role in BZ resistance.

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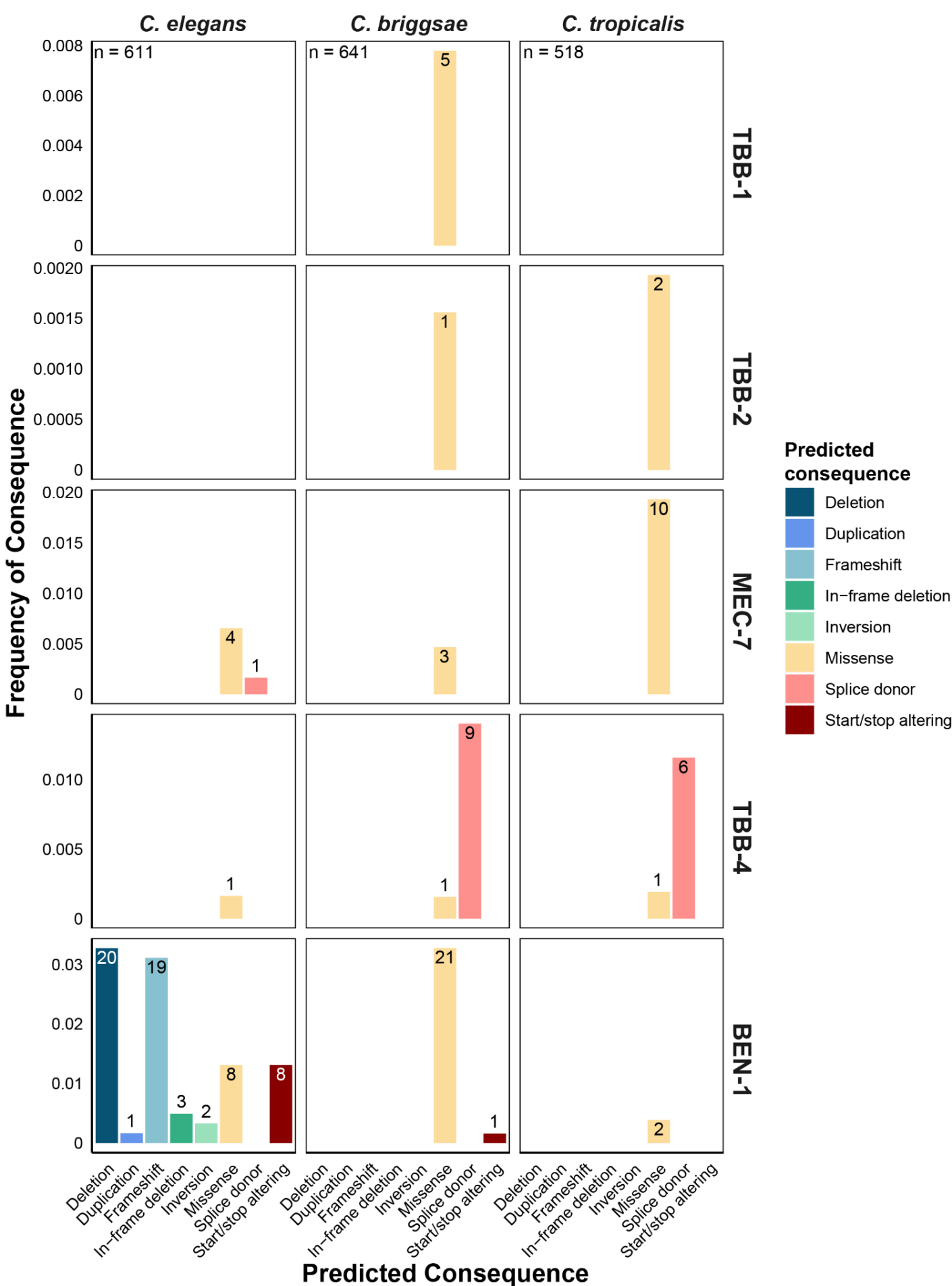


Figure 5. The frequency of predicted high-impact consequences in the five beta-tubulin genes present in natural populations of *C. elegans*, *C. briggsae*, and *C. tropicalis*

The frequency of SNVs present in natural populations of *C. elegans* (n = 611), *C. briggsae* (n = 641), and *C. tropicalis* (n = 518) (y-axis) are shown by their predicted consequence in each beta-tubulin gene (x-axis). The total number of isotype reference strains with a given predicted consequence are displayed on top of each bar plot.

Second, we examined the geographic distribution of strains carrying high-impact variants in BEN-1, TBB-1, and TBB-2 to identify if beta-tubulin variants were associated with geography. Variants in BEN-1 were found globally across the three *Caenorhabditis* species (**Fig. 6A**), with no discernible geographic pattern. In *C. elegans*, variants in BEN-1 were found in more recent clades (*i.e.*, strains that experienced recent selective sweeps) (Andersen et al. 2012; Zhang, Mostad, and Andersen 2021) (**Fig. 6B**). *Cel*-BEN-1 variants in swept clades suggest that these mutations arose as relatively recent evolutionary events and might be maintained in response to BZ-like compounds in the natural niche. By contrast, in *C. briggsae*, variants in BEN-1 were distributed throughout the species tree and found on more ancestral branches (*i.e.*, earlier diverged lineages in the species) (**Fig. 6C**). However, it is unlikely that variants in *Cbr*-BEN-1 emerged from ancestral evolutionary events, because none of the variants identified were shared across strains. For *C. tropicalis*, the limited number of variants in BEN-1 precludes any definitive conclusions regarding their evolutionary patterns (**Fig. 6D**). Finally, because few variants are found in TBB-1 and TBB-2 in *C. briggsae* and *C. tropicalis*, we cannot identify the evolutionary patterns of BZ resistance in these genes (**S13 and S14 Figures**).

Finally, as substrates harbor distinct microbial communities that can influence the evolution of BZ resistance alleles, we determined if strains carrying a high-impact variant in a beta-tubulin gene were associated with specific substrates. We performed substrate enrichment analysis to assess enrichment between 12 substrates and all strains in the three *Caenorhabditis* species (**Table S8**). However, no significant enrichment was observed between a high-impact variant in a beta-tubulin gene and any given substrate (Fisher's Exact Test, $p=1$) (**S15 Figure**). Because no geographic or substrate enrichment was observed, evolutionary pressures driving

576 beta-tubulin variation are likely not strongly tied to substrate. Instead, it is likely that species-
577 specific evolutionary trajectories (e.g., selective pressures or gene family redundancy) shape
578 differential susceptibility to BZs.

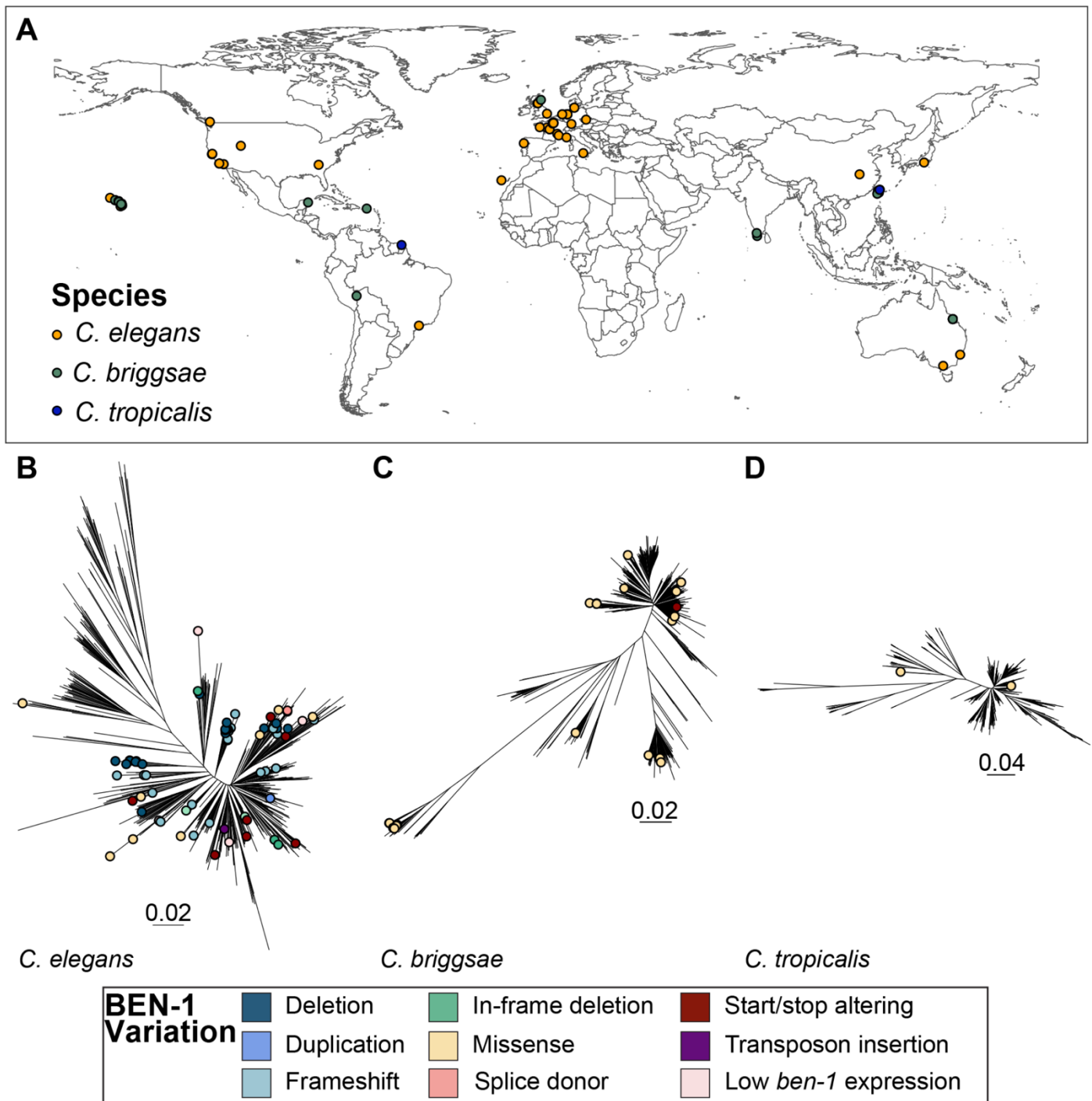


Figure 6. The global distribution of *Caenorhabditis* strains that contain predicted high-impact variation in BEN-1

(A) Each point corresponds to the sampling location of an individual *C. elegans* (orange), *C. briggsae* (green), or *C. tropicalis* (blue) isotype reference strain with a predicted high-impact consequence in BEN-1. A genome-wide phylogeny of (B) 611 *C. elegans*, (C) 641 *C. briggsae*, and (D) 518 *C. tropicalis* isotype reference strains where each point denotes an isotype reference strain with a predicted high-impact consequence in BEN-1 is shown.

4. DISCUSSION

4.1 A lack of repeated evolution of beta-tubulin mediated BZ resistance across *Caenorhabditis* nematodes

This study provides new insights into the mechanisms of ABZ resistance across three *Caenorhabditis* species. No comprehensive survey of variation in the beta-tubulin gene family or BZ response had been conducted across the three *Caenorhabditis* species to date. First, we found that each *Caenorhabditis* species had unique predicted high-impact beta-tubulin alleles that were not shared among the three species. Next, with an expanded sampling of *C. elegans*, we confirmed that a heterogeneous set of rare high-impact alleles in *ben-1* were associated with ABZ resistance (Hahnel et al. 2018; Dilks et al. 2021). Additionally, we identified a heterogenous set of rare high-impact alleles in *Cbr-ben-1*, but only two (W21stop and Q134H) were associated with ABZ resistance. The lack of ABZ resistance associated with *Cbr-ben-1* alleles suggested that other beta-tubulin genes might play a role in BZ resistance. Unlike in *C. elegans*, we identified high-impact alleles in *Cbr-tbb-1* and *Cbr-tbb-2*, but alleles in these genes were not associated with ABZ resistance. Finally, although we identified high-impact alleles in *ben-1*, *tbb-1*, and *tbb-2* in *C. tropicalis*, we did not observe ABZ resistance. The absence of repeated evolution of ABZ resistance linked to variation in beta-tubulin genes across the three *Caenorhabditis* species suggests that each species either employs distinct strategies to overcome ABZ exposure, possibly arising from distinct ecological niches with unique selection pressures, or they are not exposed to BZ compounds in the natural niche.

4.2 What drives the evolution of BZ resistance across *Caenorhabditis* species?

The lack of shared resistance alleles among the three species highlights the complexity of BZ resistance. Across clade V nematodes, BZ resistance alleles come in two types (Collins et al. 2024). First, when redundant beta-tubulin genes are present, LoF alleles of a beta-tubulin gene can cause resistance. Second, without redundant beta-tubulin genes, LoF alleles are

hypothesized to reduce fitness or even cause lethality, so only alleles that alter BZ binding cause resistance. Across *Caenorhabditis* nematode species, either type could be present so additional criteria must be assessed to understand the evolution of BZ resistance. We must clearly determine (1) the specific role of each beta-tubulin in ABZ resistance, (2) the tissue-specific expression patterns of beta-tubulins, (3) the unique selective pressures acting on each *Caenorhabditis* species, and (4) non-beta-tubulin genes involved in the ABZ response.

In *C. elegans*, the contribution of each beta-tubulin gene to BZ response has been characterized in a controlled genetic background (*i.e.*, LoF beta-tubulin alleles in the N2 strain background) (Collins et al. 2024). Similarly, we must first definitively identify the role each beta-tubulin gene plays in BZ resistance in *C. briggsae* and *C. tropicalis*. CRISPR-Cas9 genome editing should be used to delete each beta-tubulin gene in the reference strain background of each species to determine the gene's contribution to resistance. Because the reference strains for *C. briggsae* (AF16 and QX1410) and *C. tropicalis* (NIC58) are sensitive to ABZ, a resistance phenotype following gene deletion would confirm the role of the gene in ABZ resistance. Additionally, it is possible that disrupting the function of *tbb-1* and *tbb-2* in *C. briggsae* and *C. tropicalis* could be highly detrimental to animal development, as seen in *C. elegans* (Collins et al. 2024). Altogether, the creation of LoF alleles will allow us to define the role each gene plays on nematode fitness and BZ response, defining the likelihood that BZ resistance will evolve through naturally occurring LoF alleles.

In *C. elegans*, significant BZ susceptibility was identified in animals that express *ben-1* in cholinergic neurons, suggesting that *ben-1* function in this cell type underlies susceptibility to BZ (Gibson, Ness-Cohn, and Andersen 2022). The identification of beta-tubulin specific expression improves our understanding of the cellular mode of action of BZs in *C. elegans*. Similarly, we must characterize the temporal and tissue-specific expression patterns of beta-tubulins in *C. briggsae* and *C. tropicalis*. Tissue-specific differences in beta-tubulin expression among the three *Caenorhabditis* species could influence susceptibility and the evolution of BZ resistance. To

compare the expression patterns of *ben-1*, *tbb-1*, and *tbb-2*, we analyzed a publicly available dataset that estimated the conservation of expression between *C. elegans* and *C. briggsae* orthologs in homologous cell types (Large et al. 2024). The expression patterns of *ben-1* (Jensen-Shannon Distance of expression [JSDgene] = 0.21), *tbb-1* (JSDgene = 0.24), and *tbb-2* (JSDgene = 0.27) were highly conserved between *C. elegans* and *C. briggsae* (Large et al. 2024), indicating that these genes maintain similar transcriptional profiles across species. Lower JSD values indicate higher conservation, suggesting that beta-tubulin expression is largely preserved in homologous cells, which might imply that the beta-tubulins have a conserved functional role in BZ response. However, the homologous cells are not cholinergic neurons, so conserved activities in this specific neuronal cell type might be divergent.

However, some neuronal cell-types (e.g., PVQ, AVD, AVJ) expressed *Cel-ben-1* at higher levels than in the *Cbr-ben-1*, whereas two cell types, ADL and interior arcade cells, showed higher *ben-1* expression in *C. briggsae* than in *C. elegans*. (Large et al. 2024). Therefore, differences in cell-specific expression patterns of beta-tubulins might influence BZ susceptibility in each species and should be further investigated. One approach involves introducing *ben-1* into a *C. briggsae* *ben-1* knockout strain background using transgenesis, where multi-copy arrays express *ben-1* in specific tissues, as previously done in *C. elegans* (Gibson, Ness-Cohn, and Andersen 2022). Plasmids containing the coding sequence of *ben-1* are fused to tissue-specific promoters to form extrachromosomal arrays in transgenic animals, allowing precise tissue-specific expression. Investigating beta-tubulin expression patterns in specific tissues will clarify the role of each beta-tubulin and identify the tissues that BZs target in *C. briggsae*. The same approach can be applied to other beta-tubulin genes in *C. briggsae* and *C. tropicalis*. Additionally, understanding the tissue-specific roles of each beta-tubulin across *Caenorhabditis* species can inform strategies for managing resistance in parasitic nematodes by identifying key sites of BZ action. For instance, if *ben-1* function in specific tissues is critical for BZ sensitivity, it might be possible to identify similar tissues in parasitic species where beta-tubulins play a pivotal role in BZ resistance. This

information could lead to targeted approaches for enhancing drug efficacy, such as developing treatments that specifically affect these tissues or identifying novel drug targets within them to combat resistance.

Next, the lack of shared resistance variants across *C. elegans*, *C. briggsae*, and *C. tropicalis* suggests that each species experiences distinct selective pressures in the evolution of BZ resistance. Although *Caenorhabditis* species share overlapping niches, differences in microbial communities that produce natural BZs and proximity to synthetic BZs in agriculture could lead to differences in BZ response. Despite the fact that our substrate enrichment tests indicate no significant association between the substrate where a strain was collected and the probability of having a predicted resistance variant in a beta-tubulin gene, our broad substrate categories and small number of BZ resistant *C. briggsae* and *C. tropicalis* strains might obscure finer-scale ecological patterns. For instance, the microbial community composition on each substrate, such as the presence or absence of BZ producing bacterial species, could shape the evolution of BZ resistance. Future studies characterizing microbial communities associated with each substrate might clarify the selective pressures on *Caenorhabditis* nematodes.

Finally, although beta-tubulins are the primary targets of BZs in Clade V nematodes, other mechanisms, such as variation in detoxification pathways or drug efflux transporters, could drive the evolution of BZ resistance outside the beta-tubulin gene family. In *C. elegans*, variation in BZ responses among wild strains has been leveraged to identify alleles in a cytochrome P450 that cause BZ resistance independent of *Cel-ben-1* (Collins et al. 2025). Similarly, variation in ABZ response among *C. briggsae* and *C. tropicalis* strains can be investigated with quantitative genetic mapping techniques to discover sources of variation in ABZ response among wild strains. Expanding genomic resources to better characterize additional sources of variation in ABZ response among wild strains (e.g., SVs, transcriptomic variation) is critical to disentangle the lack of repeated evolution in BZ resistance.

DATA AVAILABILITY STATEMENT

All code and data used to replicate the data analysis and figures are available on GitHub at: https://github.com/AndersenLab/ce_cb_ct_betatubulin. Table S1 contains all beta-tubulin transcript IDs in the three *Caenorhabditis* species. Tables S2, S3, and S4 contain the list of *C. elegans*, *C. briggsae*, and *C. tropicalis* isotype strains, collection location, substrate type, and their beta-tubulin variant status, respectively. Table S5 contains the manual curation of SVs found in beta-tubulin genes. Table S6 contains all *C. briggsae* and *C. tropicalis* isotype strains with beta-tubulin variants (*i.e.*, predicted resistant strains) and their corresponding predicted susceptible strains. Table S7 contains the BLOSUM and Grantham scores for amino acid changes in the beta-tubulin genes in the three *Caenorhabditis* species. Table S8 contains results from the substrate enrichment analysis.

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CRedit AUTHOR STATEMENT

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722 **DECLARATION OF COMPETING INTERESTS**

723 The authors have declared that no competing interests exist.

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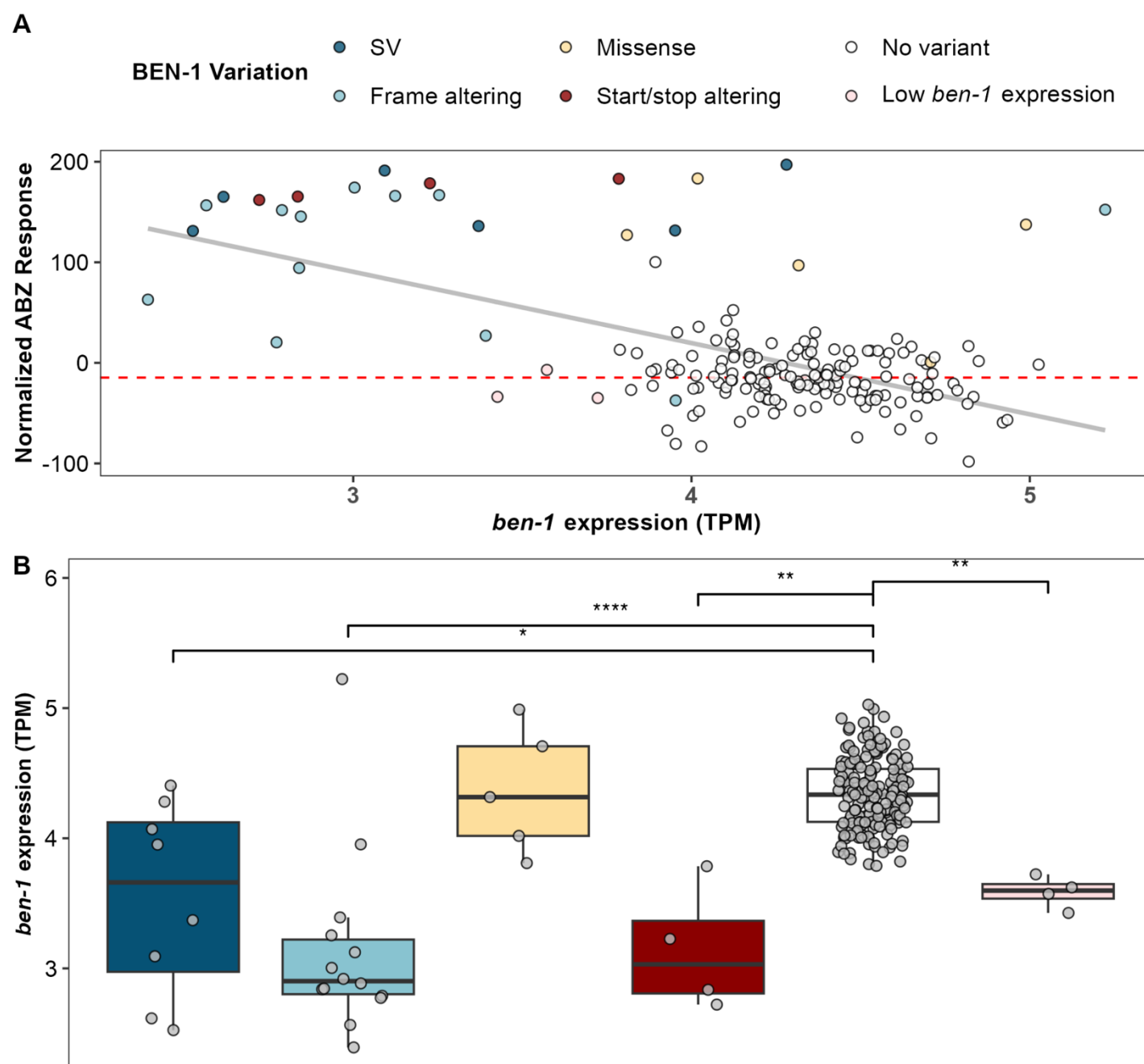
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908 **SUPPORTING INFORMATION**

- 909 **Supplemental Table 1.** Beta-tubulin transcript IDs
- 910 **Supplemental Table 2.** *C. elegans* isotype variant table
- 911 **Supplemental Table 3.** *C. briggsae* isotype variant table
- 912 **Supplemental Table 4.** *C. tropicalis* isotype variant table
- 913 **Supplemental Table 5.** Manual curation of SVs
- 914 **Supplemental Table 6.** *C. briggsae* and *C. tropicalis* strain pairs
- 915 **Supplemental Table 7.** BLOSUM and Grantham scores for amino acid changes in beta-tubulin
- 916 genes in the three *Caenorhabditis* species
- 917 **Supplemental Table 8.** Location and substrate where each isotype reference strain was
- 918 collected

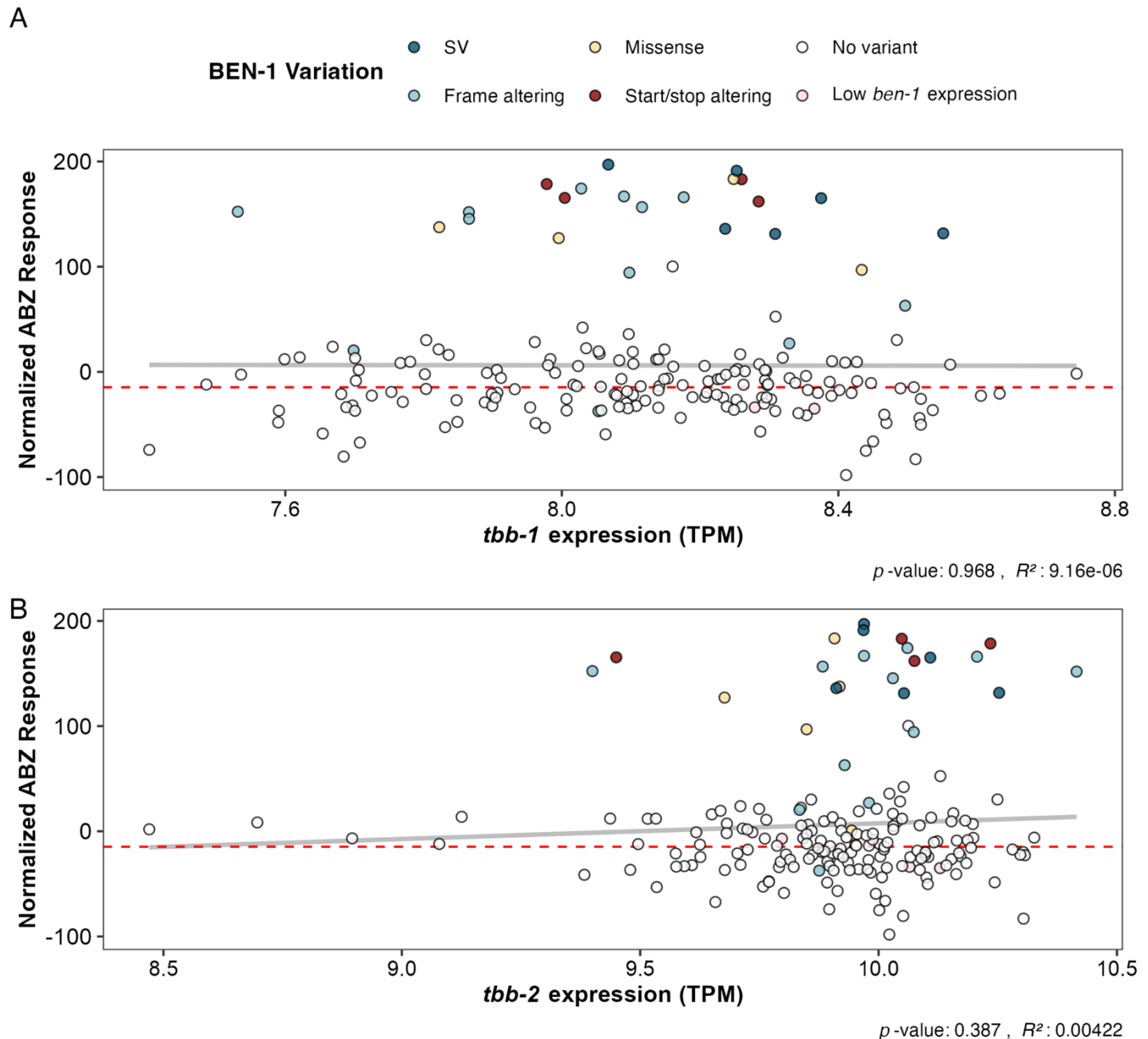
919 SUPPLEMENTAL FIGURES



Supplemental Figure 1. The relationship between *ben-1* expression levels and albendazole response in *C. elegans* strains

(A) Scatterplot of the relationship between *ben-1* expression levels and normalized albendazole (ABZ) response across *C. elegans* wild strains. Each point represents a strain phenotyped for ABZ response in previous publications (Hahnel *et al.*, 2018; Shaver *et al.*, 2024) with *ben-1* expression data (Zhang *et al.* 2022). The *ben-1* expression level measured in transcripts per million (TPM) is displayed on the x-axis. The normalized ABZ response values adjusted for assay-specific effects are displayed on the y-axis. The gray line represents the linear regression fit between *ben-1* expression and normalized response ($R^2 = 0.34$, p -value = 5.16×10^{-18}), with the

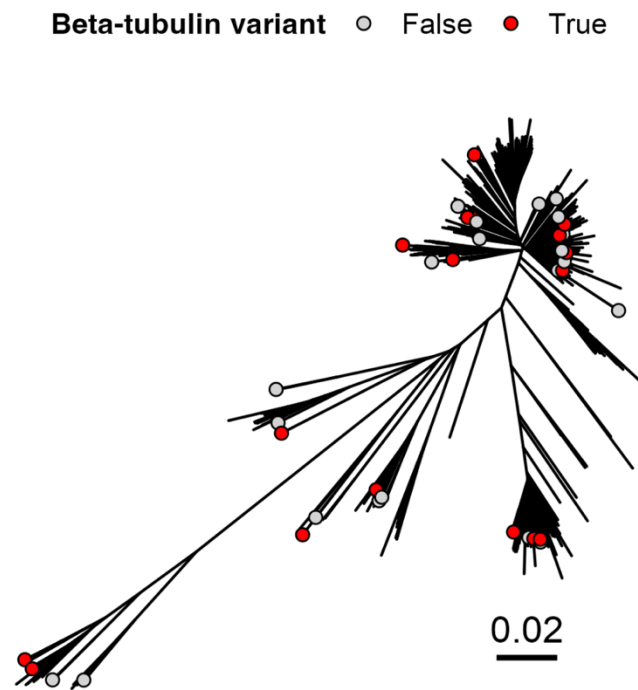
linear model's coefficient of determination (R^2). Data points are colored based on the predicted functional consequence of their *ben-1* allele (e.g., large structural variant (SV), Frame Altering, Missense substitution, Disrupted Start/Stop sequence, No high-impact variant, Low *ben-1* expression). A horizontal red line represents a resistance threshold calculated from the normalized phenotypes of all strains from previous publications set at 75% of the reference strain N2. Strains with normalized responses above this threshold are considered resistant to ABZ. **(B)** Boxplots of *ben-1* expression levels among individuals grouped by the predicted functional consequences of their *ben-1* alleles. Each point represents the *ben-1* expression level of an individual within each group. We tested for statistically significant differences in the expression between each consequence type and strains without a high-impact *ben-1* allele with an unpaired Wilcoxon test. Significance levels are indicated by symbols: '*' ($p < 0.05$), '**' ($p < 0.01$), '***' ($p < 0.001$), '****' ($p < 0.0001$).



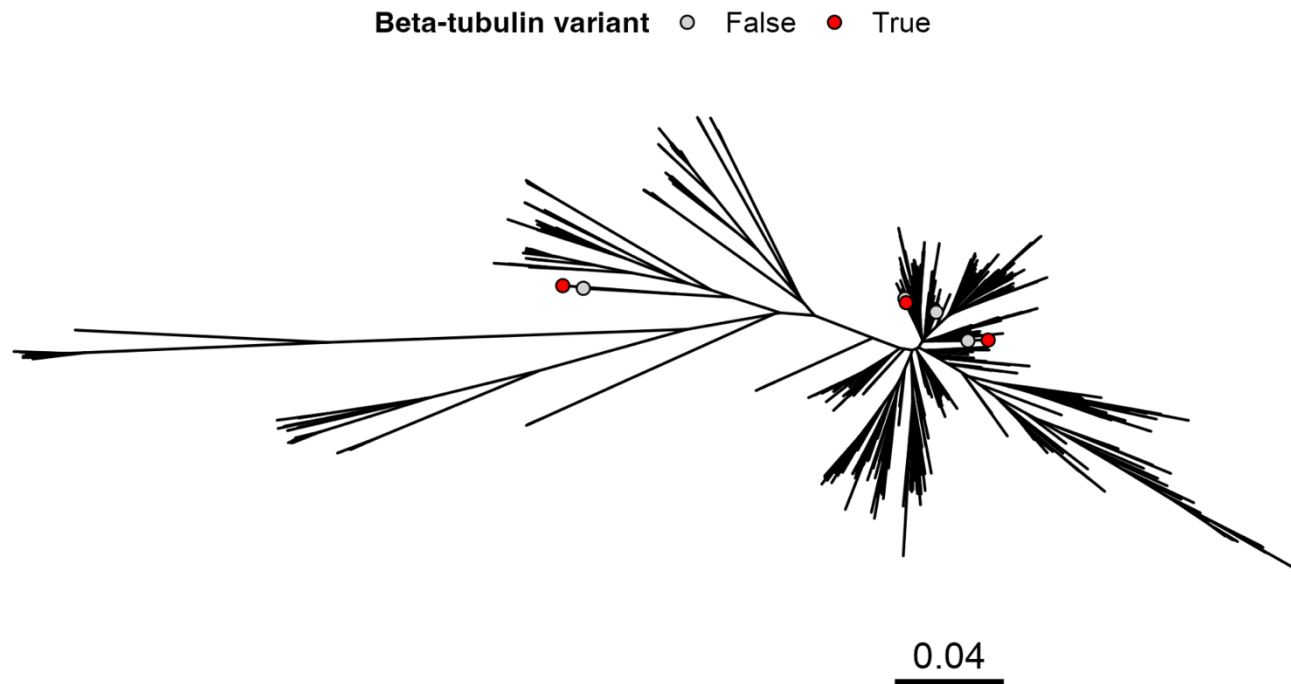
Supplemental Figure 2. The relationship between *C. elegans* beta-tubulin expression and albendazole response

Scatterplot of the relationship between (A) *tbb-1* and (B) *tbb-2* expression levels and normalized albendazole (ABZ) response across *C. elegans* wild strains. Each point represents a strain phenotyped for ABZ response in previous publications (Hahnel *et al.*, 2018; Shaver *et al.*, 2024) with *tbb-1* and *tbb-2* expression data (Zhang *et al.* 2022). The *tbb-1* and *tbb-2* expression levels measured in transcripts per million (TPM) are displayed on the x-axis. The normalized ABZ response values adjusted for assay-specific effects are displayed on the y-axis. The gray line represents the linear regression fit between beta-tubulin gene expression and normalized response, with the linear model's coefficient of determination (R^2). Data points are colored based

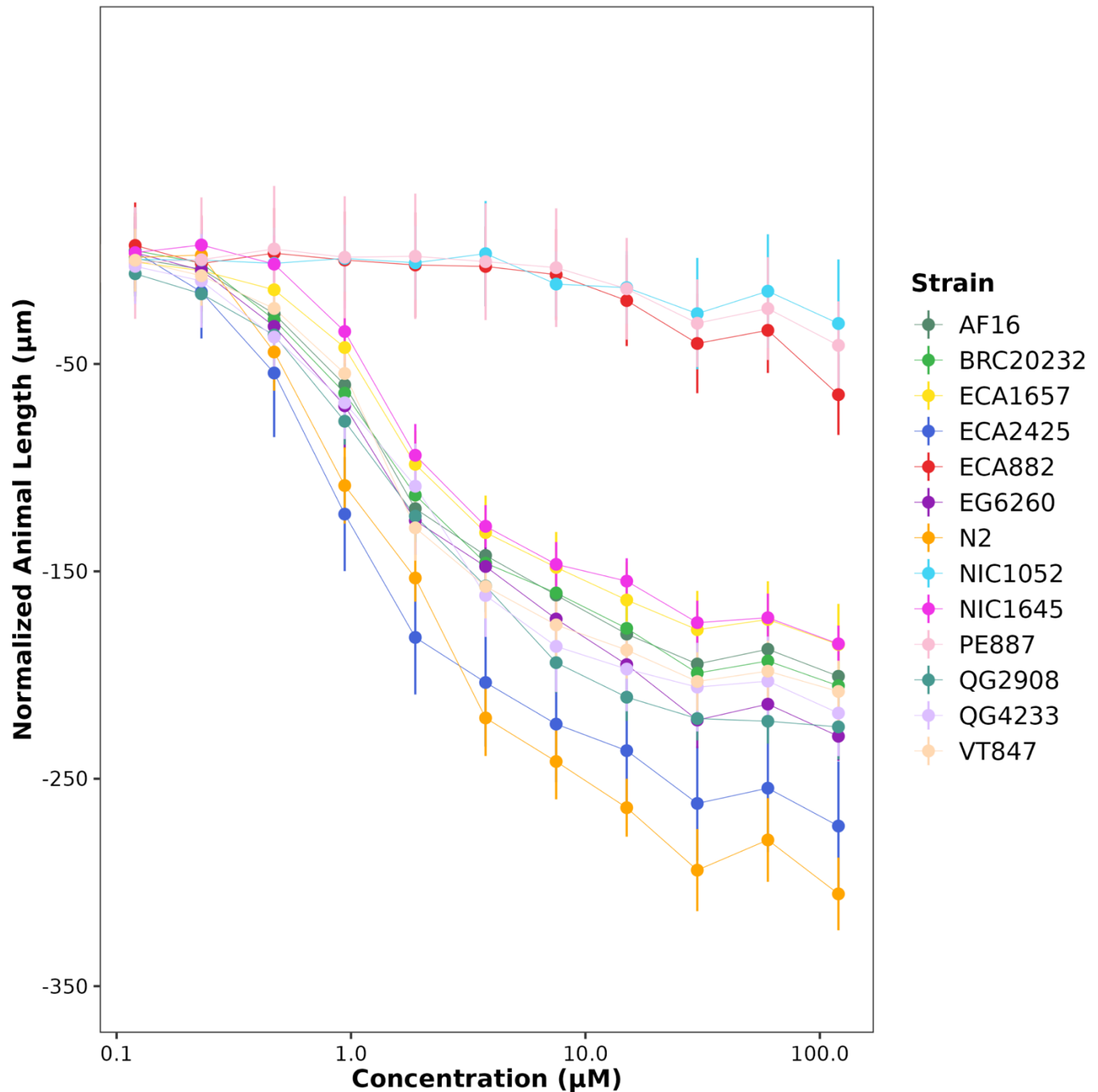
952 on the predicted functional consequence of their *ben-1* alleles (e.g., large structural variant (SV),
 953 Frame Altering, Missense substitution, Disrupted Start/Stop sequence, No high-impact variant,
 954 Low *ben-1* expression). A horizontal red line represents a resistance threshold calculated from
 955 the normalized phenotypes of all strains from previous publications set at 75% of the reference
 956 strain N2. Strains with normalized responses above this threshold are considered resistant to
 957 ABZ.



958 **Supplemental Figure 3. *C. briggsae* species tree highlighting isotype reference strains**
 959 **tested for ABZ resistance.** *C. briggsae* strains with predicted high-impact variants are colored
 960 red, and strains with no predicted variants in beta-tubulin genes are colored gray.

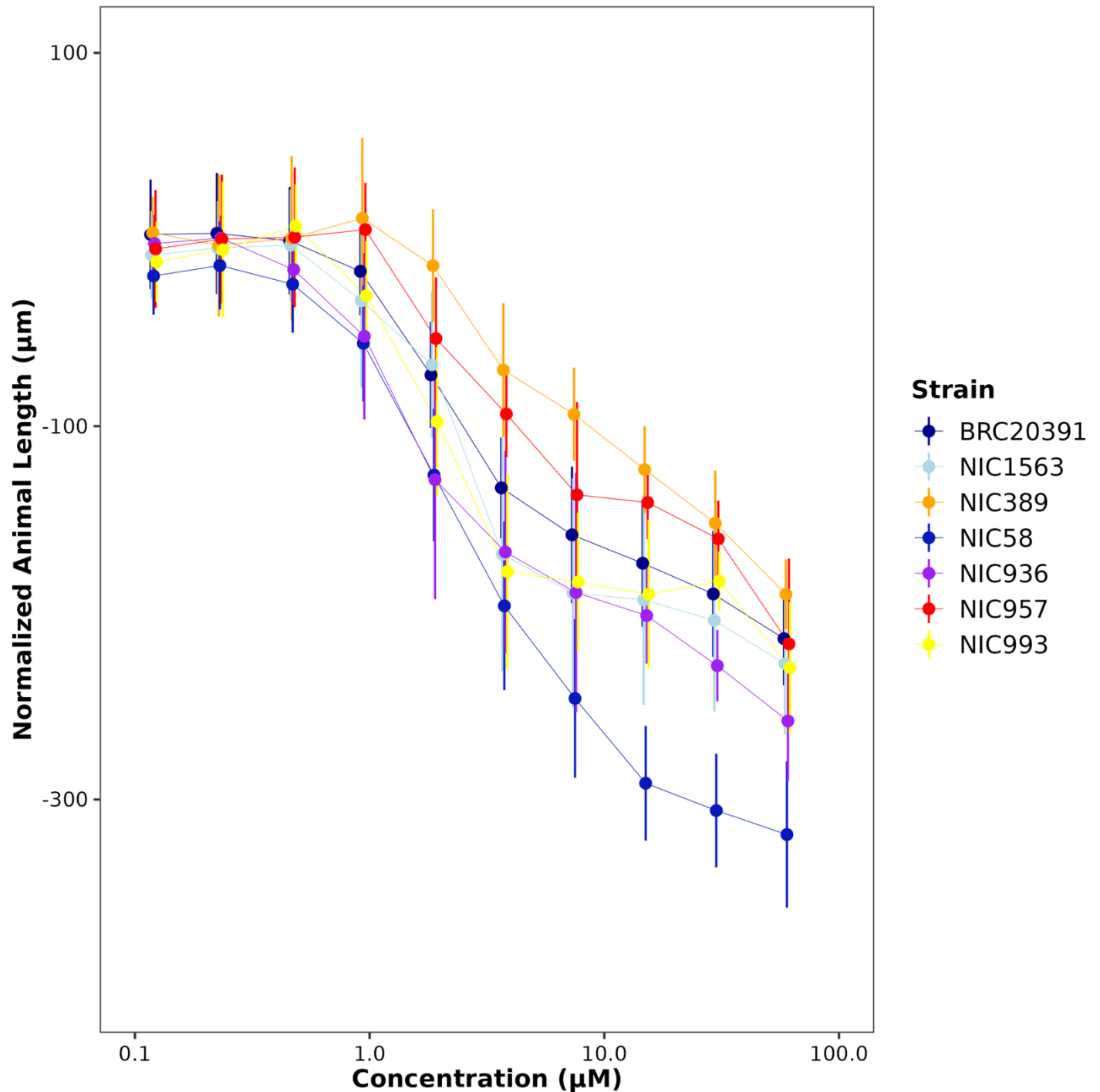


961 **Supplemental Figure 4. *C. tropicalis* species tree highlighting isotype reference strains**
 962 **tested for ABZ resistance.** *C. tropicalis* strains with predicted high-impact variants are colored
 963 red, and strains with no predicted variants in beta-tubulin genes are colored gray.



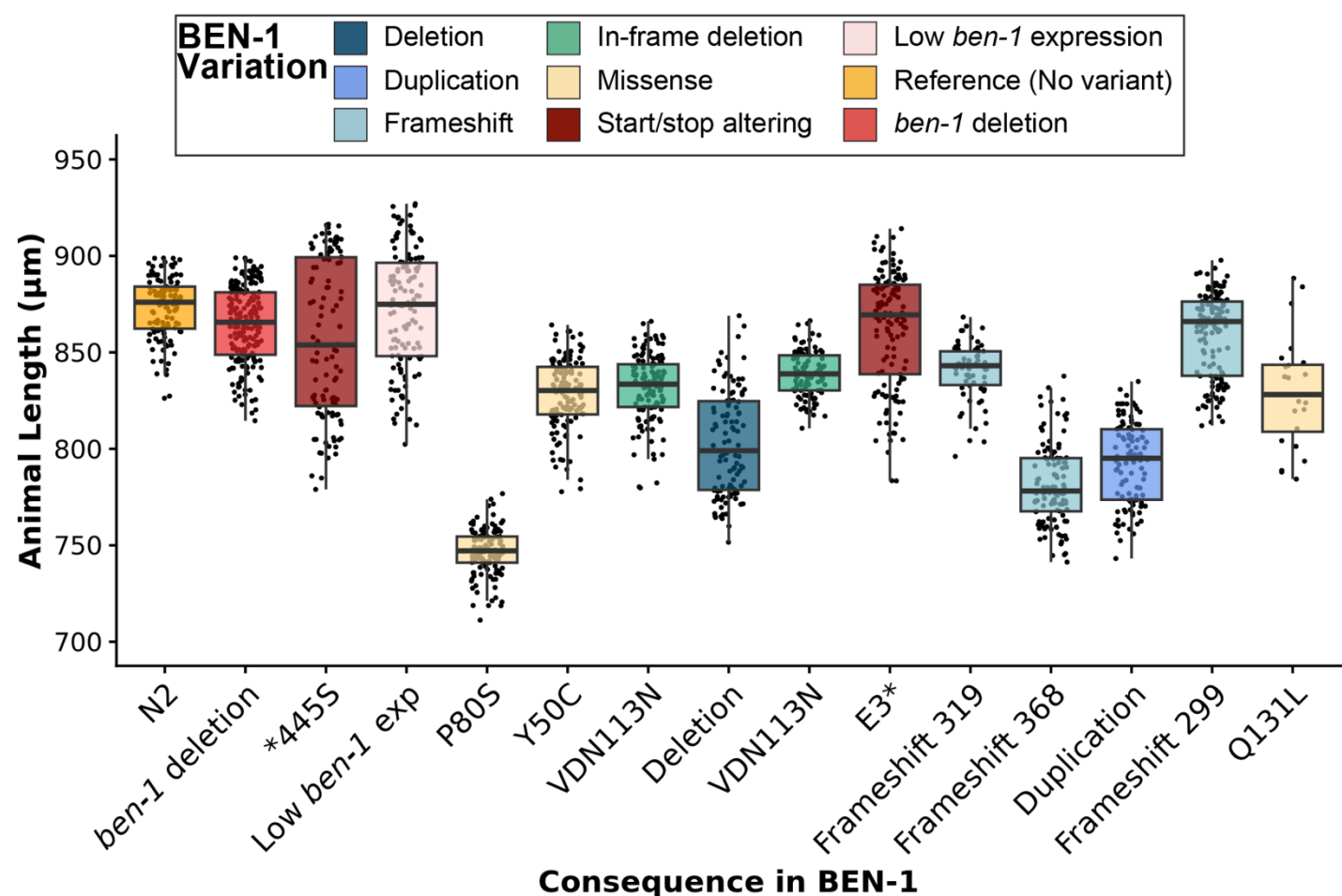
Supplemental Figure 5. Dose-response curves for *C. briggsae* strains in albendazole

Normalized animal lengths (y-axis) are plotted for each strain as a function of the dose of albendazole (ABZ) in the high-throughput development assay (x-axis). Strains are denoted by color. Lines extending vertically from points represent the standard deviation from the mean response. Statistical normalization of animal lengths is described in *Methods*. *C. elegans* strains N2 and ECA882 were added for ABZ-susceptible and ABZ-resistant controls, respectively.



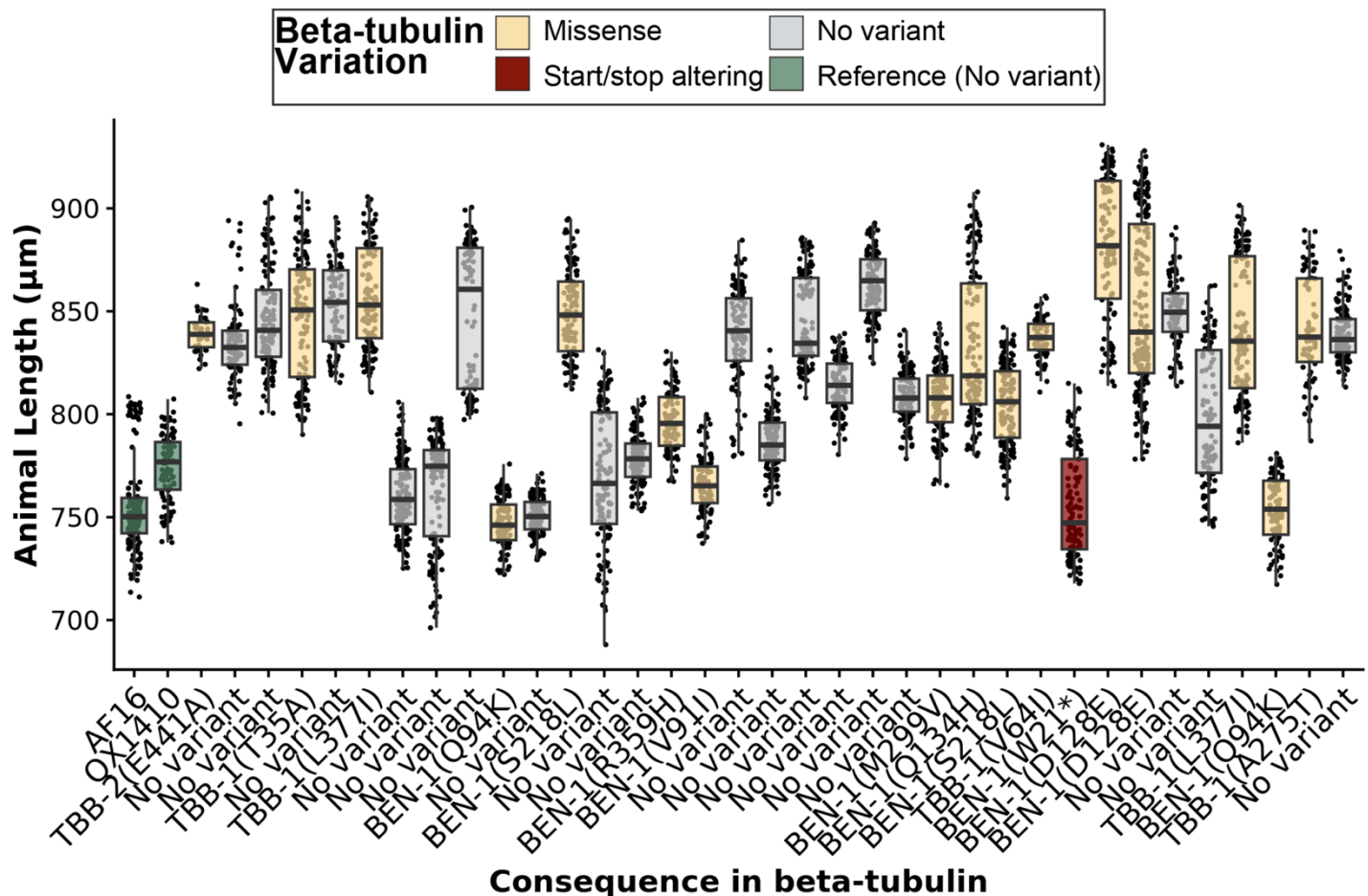
Supplemental Figure 6. Dose-response curves for *C. tropicalis* strains in albendazole

Normalized animal lengths (y-axis) are plotted for each strain as a function of the dose of albendazole (ABZ) in the high-throughput development assay (x-axis). Strains are denoted by color. Lines extending vertically from points represent the standard deviation from the mean response. Statistical normalization of animal lengths is described in *Methods*.



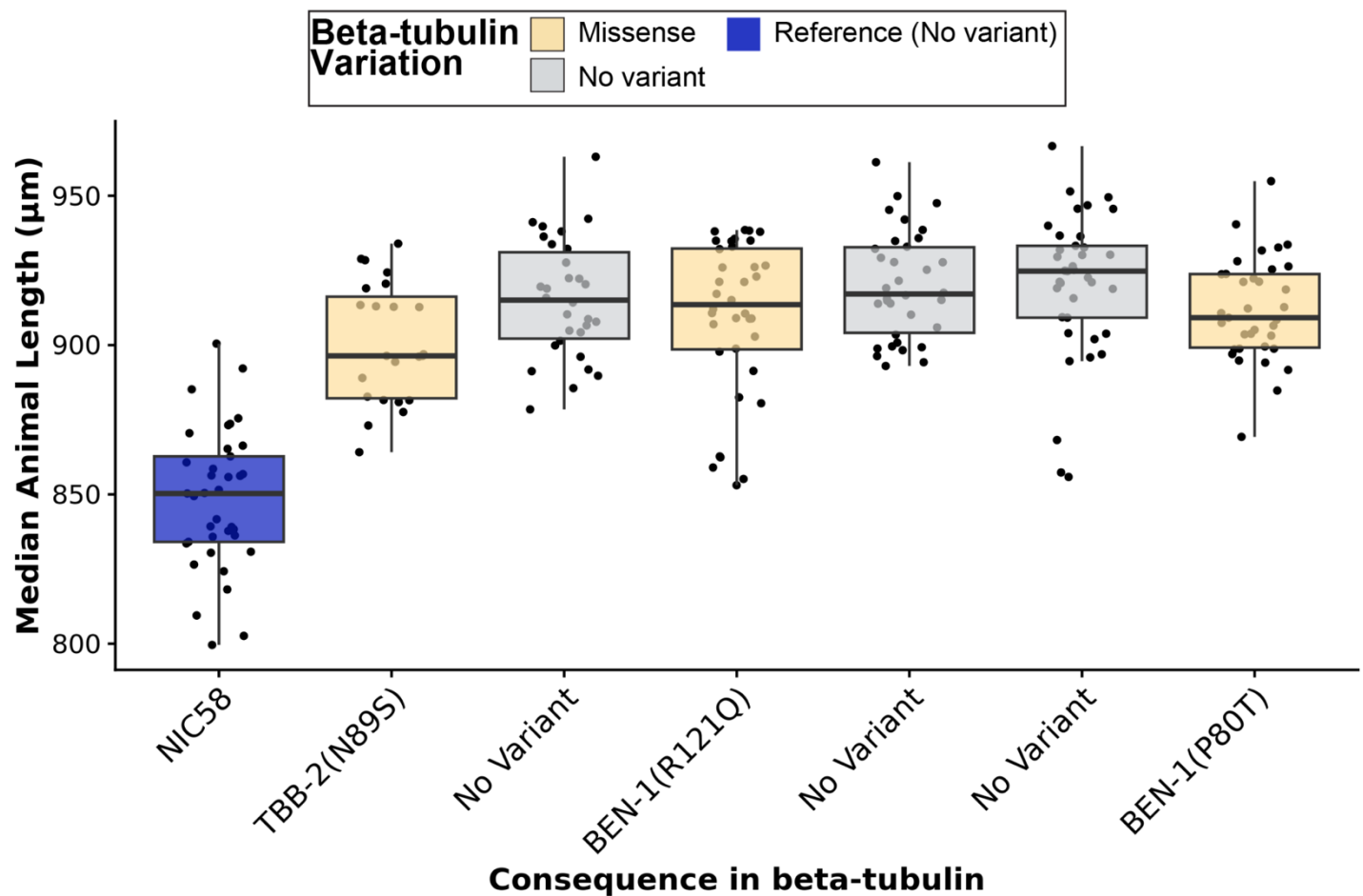
Supplemental Figure 7. High-throughput assays for each *C. elegans* strain in control conditions

Median animal length values from populations of nematodes grown in DMSO are shown on the y-axis. Each point represents the median animal length from a well containing approximately 5-30 animals. Data are shown as Tukey box plots with the median as a solid horizontal line, the top and bottom of the box representing the 75th and 25th quartiles, respectively. The top whisker is extended to the maximum point that is within a 1.5 interquartile range from the 75th quartile. The bottom whisker is extended to the minimum point that is within the 1.5 interquartile range from the 25th quartile.



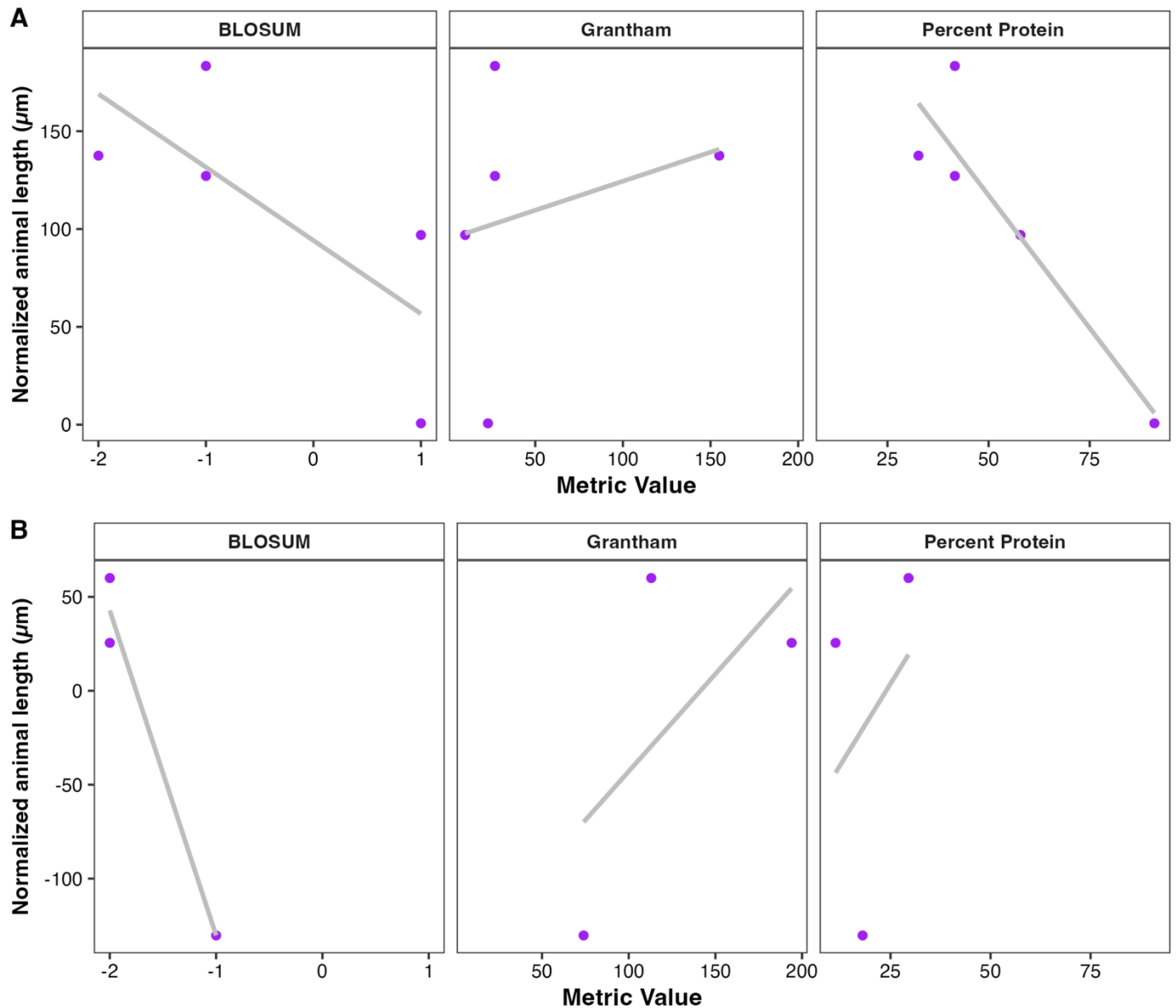
Supplemental Figure 8. High-throughput assays for each *C. briggsae* strain in control conditions.

Median animal length values from populations of nematodes grown in DMSO are shown on the y-axis. Each point represents the median animal length from a well containing approximately 5-30 animals. Data are shown as Tukey box plots with the median as a solid horizontal line, the top and bottom of the box representing the 75th and 25th quartiles, respectively. The top whisker is extended to the maximum point that is within a 1.5 interquartile range from the 75th quartile. The bottom whisker is extended to the minimum point that is within the 1.5 interquartile range from the 25th quartile. Strains are colored by beta-tubulin variant status.

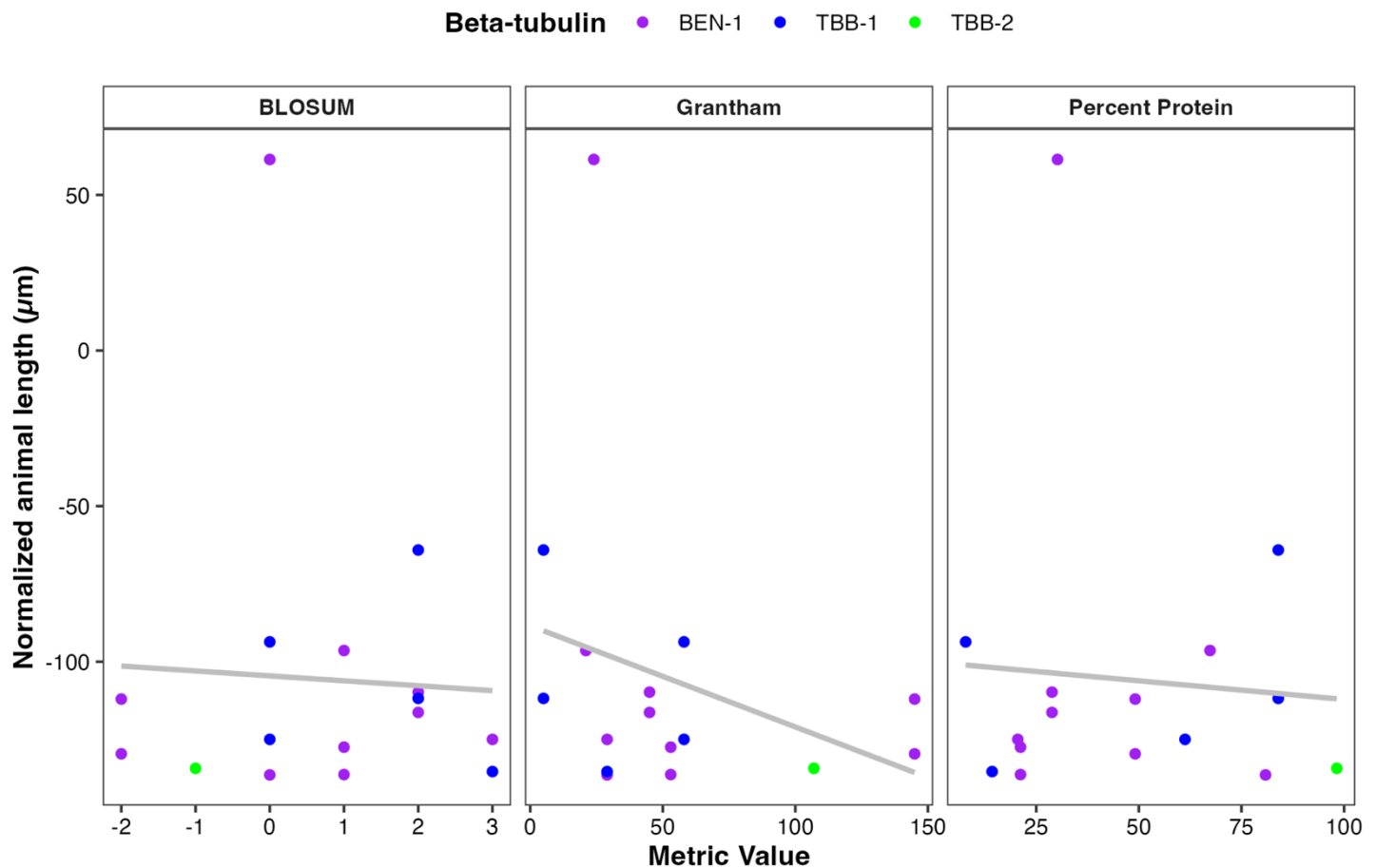


Supplemental Figure 9. High-throughput assays for each *C. tropicalis* strain in control conditions

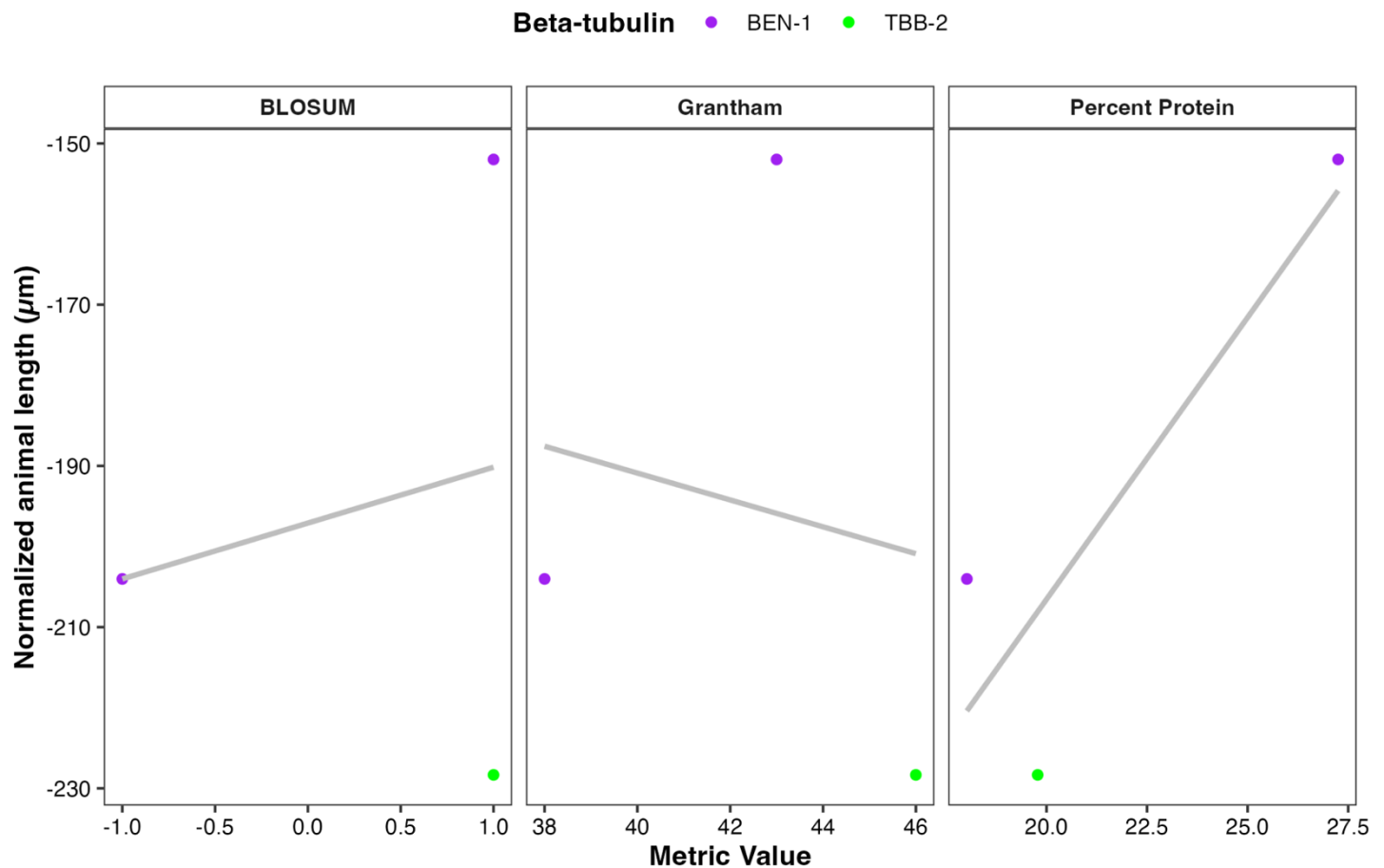
Median animal length values from populations of nematodes grown in DMSO are shown on the y-axis. Each point represents the median animal length from a well containing approximately 5-30 animals. Data are shown as Tukey box plots with the median as a solid horizontal line, the top and bottom of the box representing the 75th and 25th quartiles, respectively. The top whisker is extended to the maximum point that is within a 1.5 interquartile range from the 75th quartile. The bottom whisker is extended to the minimum point that is within the 1.5 interquartile range from the 25th quartile. Strains are colored by beta-tubulin variant status.



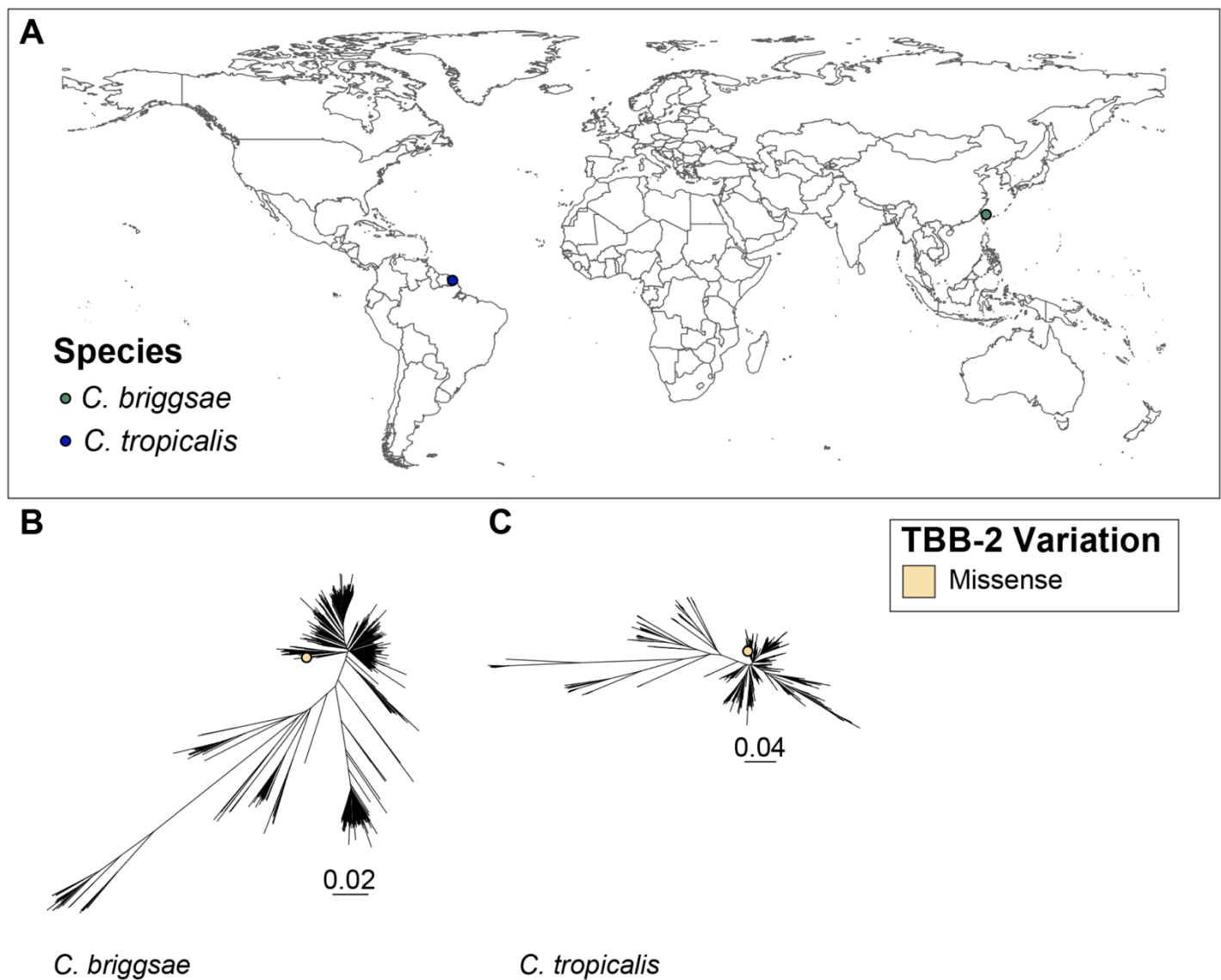
Supplemental Figure 10. Relationship between missense substitutions in BEN-1 and albendazole response in *C. elegans* strains. Scatterplots show the relationship between normalized median animal length (y-axis) and three amino acid substitution scoring metrics (x-axis): BLOSUM ($R^2 = 0.97$, p -value = 0.11), Grantham ($R^2 = 0.39$, p -value = 0.57), and percent protein ($R^2 = 0.1$, p -value = 0.8). Each point represents a *C. elegans* strain with a missense substitution in BEN-1 that was (A) phenotyped for ABZ response in previous assays (Hahnel et al. 2018; Shaver et al. 2024) or (B) in the high-throughput assays performed for this manuscript are reported. Gray lines indicate the linear regression fit for these models.



1017 **Supplemental Figure 11. Relationship between missense substitutions in BEN-1, TBB-1,**
1018 **TBB-2 and albendazole response in *C. briggsae* strains.** Scatterplots show the relationship
1019 between normalized median animal length (y-axis) and three amino acid substitution scoring
1020 metrics (x-axis): BLOSUM ($R^2 = 0.0$, p -value = 0.85), Grantham ($R^2 = 0.08$, p -value = 0.28),
1021 percent protein ($R^2 = 0.01$, p -value = 0.79). Each point represents a *C. briggsae* strain with a
1022 missense substitution in either BEN-1, TBB-1, or TBB-2. Gray lines indicate the linear regression
1023 fit, and the R^2 and p -values of these models are displayed in each panel

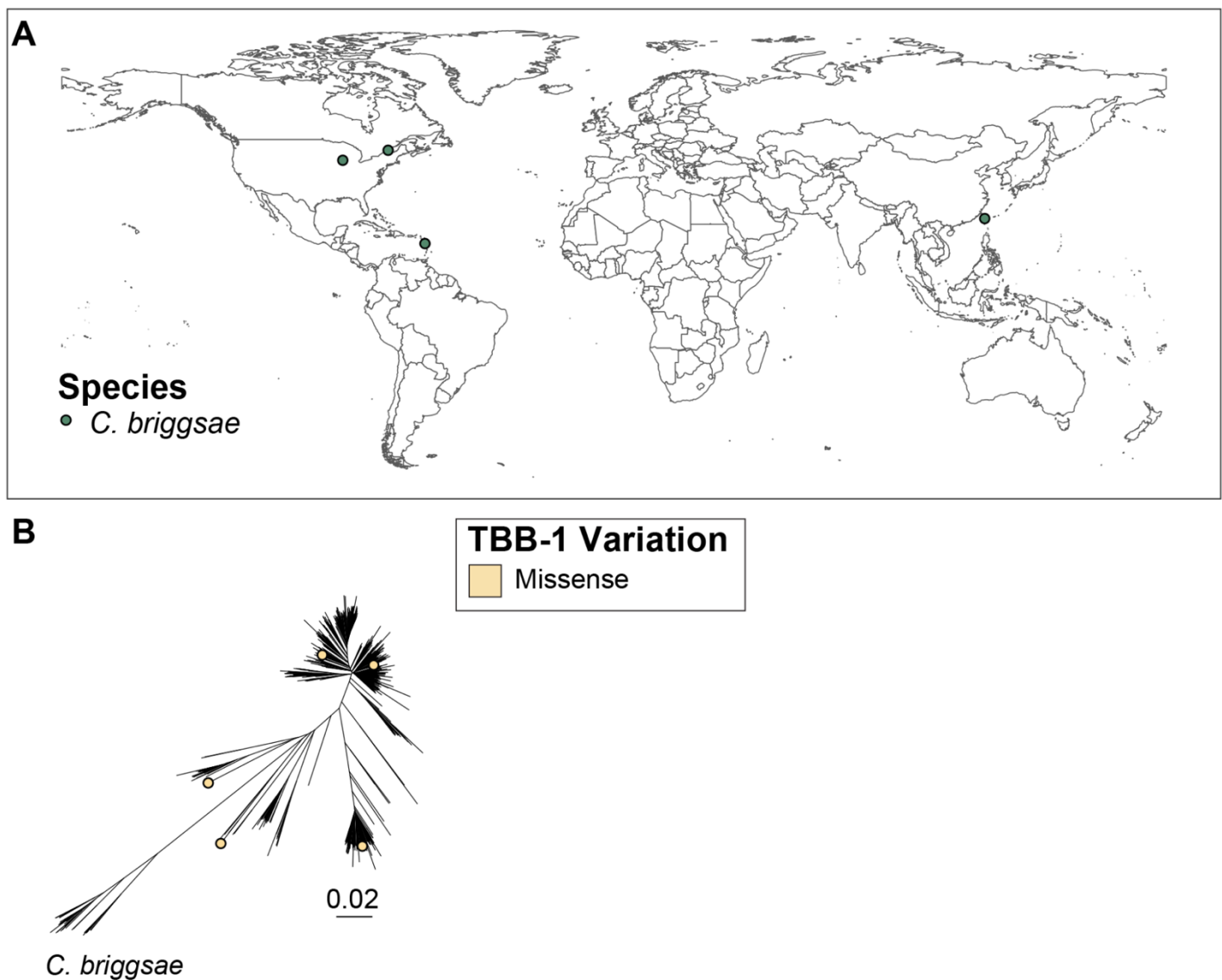


Supplemental Figure 12. Relationship between missense substitutions in BEN-1 or TBB-2 and albendazole response in *C. tropicalis* strains. Scatterplots show the relationship between normalized median animal length (y-axis) and three amino acid substitution scoring metrics (x-axis): BLOSUM ($R^2 = 0.04$, p -value = 0.87), Grantham ($R^2 = 0.03$, p -value = 0.89), percent protein ($R^2 = 0.77$, p -value = 0.32). Each point represents a *C. tropicalis* strain with a missense substitution in BEN-1 or TBB-2. Gray lines indicate the linear regression fit of these models.



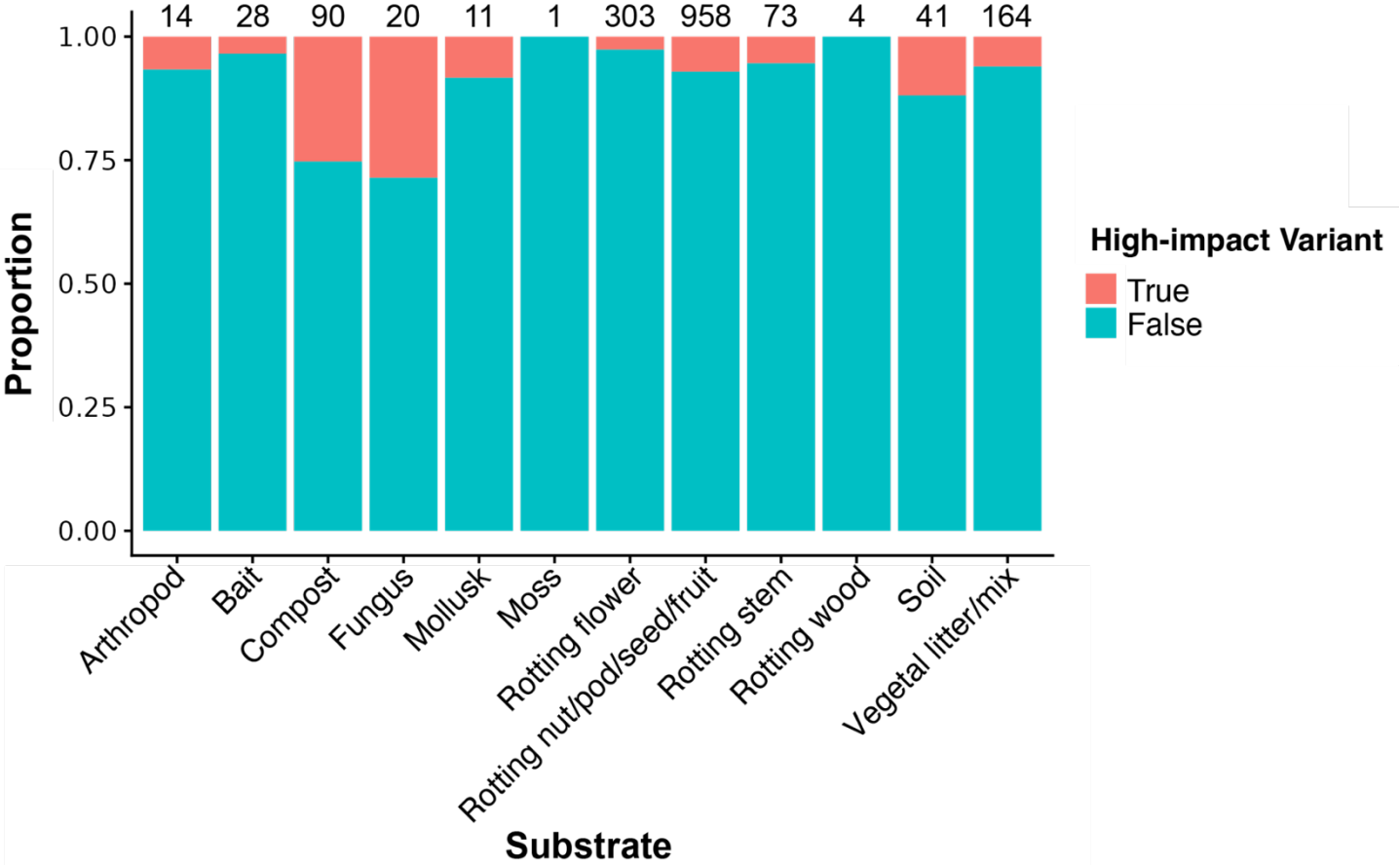
Supplemental Figure 13. The global distribution of *Caenorhabditis* strains that contain predicted high-impact variation in *tbb-2*

(A) Each point corresponds to the sampling location of an individual *C. briggsae* or *C. tropicalis* isotype reference strain with a predicted high-impact consequence in the gene *tbb-2*. A genome-wide phylogeny of (B) 641 *C. briggsae* and (C) 518 *C. tropicalis* isotype reference strains where each point denotes an isotype reference strain with a predicted high-impact consequence in *tbb-2*.



Supplemental Figure 14. The global distribution of *Caenorhabditis* strains that contain predicted high-impact variation in *tbb-1*

(A) Each point corresponds to the sampling location of an individual *C. briggsae* or *C. tropicalis* isotype reference strain with a predicted high-impact consequence in the gene *tbb-1*. (B) A genome-wide phylogeny of 641 *C. briggsae* isotype reference strains where each point denotes an isotype reference strain with a predicted high-impact consequence in *tbb-1*. One isotype, XZ1213 has a high-impact *tbb-1* variant, but sampling coordinates were not recorded. (C) A genome-wide phylogeny of 518 *C. tropicalis* isotype reference strains where each point denotes an isotype reference strain with a predicted high-impact consequence in *tbb-1*.



Supplemental Figure 15. The proportion of strains with high-impact resistant variants in beta-tubulin genes and the substrates where those strains were found
The proportion of strains (y-axis) found in a given substrate (x-axis) are displayed. Strains with a high-impact variant in a beta-tubulin gene are colored salmon. Strains with no variants in a beta-tubulin gene are colored teal. The total number of strains isolated from a given substrate is displayed above each column. Moss and rotting wood were not included in the substrate enrichment analysis due to the small sample size. No significant relationship between beta-tubulin gene variant status and substrate were identified (Fisher's Exact Test, $p=1$).