JaponicusDB: rapid deployment of a model organism database for an emerging model species

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Abstract

The fission yeast Schizosaccharomyces japonicus has recently emerged as a powerful system for studying the evolution of essential cellular processes, drawing on similarities as well as key differences between S. japonicus and the related, well-established model Schizosaccharomyces pombe. We have deployed the open-source, modular code and tools originally developed for PomBase, the S. pombe model organism database (MOD), to create JaponicusDB (www.japonicusdb.org), a new MOD dedicated to S. japonicus. By providing a central resource with ready access to a growing body of experimental data, ontology-based curation, seamless browsing and querying, and the ability to integrate new data with existing knowledge, JaponicusDB supports fission yeast biologists to a far greater extent than any other source of S. japonicus data. JaponicusDB thus enables S. japonicus researchers to realize the full potential of studying a newly emerging model species and illustrates the widely applicable power and utility of harnessing reusable PomBase code to build a comprehensive, community-maintainable repository of species-relevant knowledge.

Keywords: model organism database; Schizosaccharomyces japonicus; genome; annotation; curation; evolution

Introduction

In recent years, the fission yeast Schizosaccharomyces japonicus has emerged as a powerful system for studying evolutionary cell biology, via comparison to the related, well-established model eukaryote Schizosaccharomyces pombe. Although these two fission yeasts carry out conserved cell-level processes using similar gene complements, substantial physiological differences in the mechanisms underpinning cell growth and division (Aoki et al. 2011; Yam et al. 2011; Gu et al. 2015; Makarova et al. 2016), physiology (Okamoto et al. 2013; Kinnaer et al. 2019), and metabolism (Builder 1963; Kaino et al. 2018; Makarova et al. 2020) provide invaluable opportunities to study how processes accomplished by universally conserved gene products may diverge. Indeed, S. pombe and S. japonicus may be regarded together as a composite model system in which conserved processes, gene products, and associated phenotypes can be compared side by side (Gu and Oliferenko 2015; Russell et al. 2017; Oliferenko 2018). Furthermore, S. japonicus can be used as a standalone model organism for studying biological processes not apparent or readily tractable in other model yeasts, such as nuclear envelope breakdown and reassembly (Yam et al. 2013; Pieper et al. 2020), cellular geometry scaling (Gu and Oliferenko 2019), and quorum sensing (Gómez-Gil et al. 2019).

The current trajectory of S. japonicus research also aptly illustrates how the demands placed on species-specific data resources grow and change as an organism becomes established as a model. To date, the S. japonicus genome sequence and associated computationally generated annotation available from existing species-neutral data aggregators and repositories, such as the INSDC databases (https://www.insdc.org/), Ensembl Genomes (Howe et al. 2020), UniProtKB/TrEMBL (The UniProt Consortium 2020), and FungiDB (Basenko et al. 2018), has sufficed for the early stages of research in this model organism. As a model system matures, however, researchers move beyond simple exploratory screens to designing more ambitious research programs that generate heterogeneous gene-specific molecular data for entire processes—a stage that S. japonicus research is now reaching. To accommodate growing bodies of literature and data, and to realize the full potential of coordinated studies investigating the divergent biology of fission yeasts, the accumulated results of S. japonicus experiments must be made readily available to the research community in the expertly curated and integrated state provided only by a model organism database (MOD; Oliver et al. 2016, Lipshultz 2021).

To address the urgent need for MOD infrastructure and services for S. japonicus, we have created JaponicusDB using the suite of open-source modular, customizable tools and code originally developed for PomBase, the S. pombe MOD (Lock et al. 2019 and Harris et al. 2021). The PomBase database system was designed to facilitate reuse for emerging model species and comprises an online curation environment (Canto; Rutherford et al. 2014), a curation database using the Chado schema...
(Mungall et al. 2007), and code to import data and generate a website that features intuitive displays, a versatile query system, a genome browser (JBrowse; Buels et al. 2016), and support for daily data updates. Here, we describe the initial configuration and population of JaponicusDB and report on its current status. We summarize a round of manual curation in which we corrected and updated gene structure predictions, named genes, improved ortholog detection, and provided a greatly improved corpus of Gene Ontology (GO; The Gene Ontology Consortium 2000, 2021) annotation. Because community curation using Canto has proven successful in enhancing S. pombe literature curation (Lock et al. 2020), we have launched an analogous community curation approach for S. japonicus as a core part of JaponicusDB.

Now that both PomBase and JaponicusDB use the same intuitive database system and curation environment, all fission yeast researchers can easily carry out numerous activities that would not otherwise be feasible. Researchers can curate detailed data from small-scale experiments, including phenotypes, modifications, molecular functions, interactions, and processes; display data from publication-based curation rapidly and conveniently; query curated data via simple, intuitive tools; and compare lists of genes identified in specific experiments with comprehensively curated lists in an intuitive and meaningful way. The rapid deployment of JaponicusDB showcases how reusing the PomBase system brings immediate benefits to a model organism community in exchange for very reasonable input funds and effort.

**Methods**

JaponicusDB shares a codebase with PomBase, which is used to build the database, load the data and run the website. All database-specific differences are captured in configuration files. The entire codebase is available from the PomBase GitHub organization (https://github.com/pombase/), and the JaponicusDB configuration files and manual curation in a separate GitHub organization (https://github.com/japonicusdb/). Now that the initial setup is complete, JaponicusDB can be maintained almost entirely by editing files stored in GitHub repositories; anyone with a GitHub account can thus be authorized to make corrections and other changes. The database and website are automatically updated daily using the latest code, ontology versions, configuration, and data files. Problems detected during this process are recorded to publicly visible log files to allow remote users to investigate issues.

**Database initialization**

The first step in each daily update is to initialize a PostgreSQL database with an empty Chado schema (Mungall et al. 2007) and populate it with the required ontologies [GO; The Gene Ontology Consortium 2000, 2021], the Fission Yeast Phenotype Ontology (FYPO; Harris et al. 2013), the Sequence Ontology (SO; Eilbeck et al. 2005), the chemical ontology ChEBI (Hastings et al. 2005), the InterPro and Gene Ontology Annotation (GOA), all annotation files are now maintained in GitHub repositories.

**Prepping datasets**

We prepared data files for the first JaponicusDB load as described below. As noted above, with the exception of external files from InterPro and Gene Ontology Annotation (GOA), all annotation files are now maintained in GitHub repositories.

**Genome**

The S. japonicus genome contigs (EMBL: KE651166–KE651197; Rhind et al. 2011) and mitochondrial genome (EMBL: AF547983; Bullerwell et al. 2003) were downloaded from the European Nucleotide Archive (ENA; Harrison et al. 2021). Genome locus tags were repurposed to provide systematic identifiers, and identifiers (SJAGMIT_01–SJAGMIT_37) were minted for the mitochondrial genome.

**Transposons and LTRs**

All transposons, transposon-related fragments, and LTRs reported by Rhind et al. (2011) were added to the sequence contigs, assigned IDs in the range SJATN_00001–SJATN_00265, and loaded into the S. japonicus JBrowse instance.

**Orthologs**

UniProtKB (The UniProt Consortium 2020) records were downloaded using a query for taxon ID 402676 (the sequenced strain). Gene names applied by UniProtKB or via ENA submissions were imported. These were augmented by automated transfer of S. pombe gene names and products from PomBase where there is a one-to-one ortholog. Automatically transferred names are updated daily, with a configurable provision to override any unsuitable gene names or product descriptions. This configuration file also supplies manually curated names and product descriptions for paralogs, which we resolved based on synteny, and for genes with complicated orthology relationships, for which we used protein family membership to guide our decisions.

**Schizosaccharomyces pombe orthologs**

The S. japonicus–S. pombe orthologs identified by Rhind et al. (2011) for 4302 gene products were imported. These predictions were supplemented with orthologs from Ensembl Compara (Herrero et al. 2016), bringing the number of S. japonicus proteins with identified S. pombe orthologs to 4504. To find missing divergent orthologs, we identified proteins that are conserved between S. pombe and other species but did not have identified orthologs in S. japonicus, and devised gene-specific search strategies using a combination of FASTA (Pearson and Lipman 1988), JackHMMER (Johnson et al. 2010), PSI-BLAST (Altschul et al. 1997), TBLASTN (Gertz et al. 2006), and synteny. This procedure also identified missing genes and gene structures that required revision.

**Human and budding yeast orthologs**

In PomBase, orthologs between S. pombe and Saccharomyces cerevisiae (budding yeast) and between S. pombe and human have been manually curated over 20 years using multiple ortholog prediction methods and tailored search strategies to provide complete and highly accurate coverage of known orthologs, including many not identified by automated methods (Wood 2005; Hu et al. 2011). We inferred human and budding yeast orthologs by transferring orthologs manually curated by PomBase for S. pombe genes to the orthologous S. japonicus genes. These were supplemented with orthologs from Compara for proteins that have no S. pombe ortholog but are conserved in other species.

**Families and domains**

Protein domain information from InterPro version 85 (Blum et al. 2021), together with coiled-coil and low-complexity regions from Pfam version 34.0 (El-Gebali et al. 2019), was imported using existing PomBase Chado loading code. Transmembrane domains were predicted using TMHMM (Krogh et al. 2001).
Gene Ontology

GO annotations were obtained by downloading the GOA UniProt file produced by the GOA project at EBI (Huntley et al. 2015), and then filtering for NCBI taxon ID 4897 or 402676. The loading script checks the GOA UniProt file (located at ftp://ftp.ebi.ac.uk/pub/databases/GO/goa/UNIPROT/goa\_uniprot\_all.gaf.gz) for updates upon each run. For the first JaponicusDB load, we used the GOA file from July 2021, which contains 36,878 S. japonicus annotations.

The filtering protocol used by PomBase to retain only relevant and nonredundant automated annotation, in which GO annotations are filtered at two stages, was applied (Lock et al. 2019). Prior to loading into Chado, the GOAs were filtered to exclude any derived from false positive or otherwise inapplicable InterPro2GO (Finn et al. 2017, https://github.com/geneontology/go-site/blob/master/metadata/gorefs/goref-0000002.md) and keyword (https://github.com/geneontology/go-site/blob/master/metadata/gorefs/goref-0000041.md, https://github.com/geneontology/go-site/blob/master/metadata/gorefs/goref-0000043.md) mappings, or phylogeny-based transfer (Gaudet et al. 2011), and any GO terms that are flagged by GO as not usable for direct annotation, or as inapplicable due to taxon restrictions (Deegan née Clark et al. 2010; The Gene Ontology Consortium 2021); after filtering, 24,159 annotations were loaded.

To enhance the S. japonicus GO annotation set, S. pombe GO annotations were transferred to S. japonicus orthologs, and manual annotations from literature curated in Canto were added. The annotations loaded into Chado were then subjected to the second round of filtering, which removes redundant nonexperimental GO annotation.

The JaponicusDB GO pipeline also supports configurable filtering that can be used in cases where paralogs have diverged (e.g., one or both of a paralogous pair has undergone neofunctionalization or sub-functionalization), making the S. pombe annotation inappropriate for S. japonicus; with automated annotation transfer suppressed, the paralogs can be manually annotated. Annotation from PomBase is also not transferred for any gene–term combination that contradicts a manually curated negated (NOT) annotation in JaponicusDB (e.g., Mid1 GO:1902408). Finally, for known proteins conserved in other species but absent from S. pombe, GO annotations were made manually by inference from experimentally characterized orthologs. As of September 2021, JaponicusDB has 29,589 GO annotations.

Identification of missing genes and revision of gene structures protein-coding genes

During manual ortholog assignment, we observed that a number of genes at syntenic locations did not display any clear sequence similarity. Inspecting the structures of these genes revealed errors, which were corrected. All gene structure editing was done using the Artemis genome annotation tool (Rutherford et al. 2000). Gene structures annotated in the published assembly without methionines were either trimmed to the appropriate methionine or revised to include a missing N-terminal exon. Gene structure errors reported in publications were also corrected (Makarova et al. 2016, 2020).

We created a list of S. pombe genes that are broadly conserved in single copy throughout eukaryotes (predominantly one-to-one to human) using the PomBase advanced search and imported the result list into JaponicusDB using the identifier mapping feature (Harris et al. in this issue). This query combination provided a list of conserved proteins present in S. pombe but absent from S. japonicus.

Since we expected most of these genes to be present in S. japonicus, we used TBLASTN to perform directed searches against the S. japonicus genome, using the S. pombe proteins as input, coupled with manual inspection and gene structure curation in Artemis to identify highly spliced genes at syntenic locations.

Noncoding RNAs

Annotated features imported with the genome sequence included 347 tRNA genes and 17 rRNA genes, but no other noncoding RNAs. We used tRNAscan-SE (Chan and Lowe 2019) to confirm the tRNA complement, identifying three additional tRNA genes (SJAG_08001–SJAG_08003), and as a source of codon information for the product descriptions. Other missing noncoding RNAs were imported from RNAcentral (The RNAcentral Consortium 2019).

Website code and documentation

The JaponicusDB website uses the PomBase website code (Lock et al. 2019), configured for S. japonicus. In brief, the Chado database is processed nightly to build a Docker container that holds a complete instance of the website, including all documentation, HTML, code, and data.

All documentation for JaponicusDB is maintained in Markdown format in a GitHub repository and the source files are shared with PomBase where possible.

Canto setup and literature import

The Canto community curation tool was designed to support different species and datatypes via simple configuration adjustments (Rutherford et al. 2014). As deployed for S. japonicus, Canto currently supports annotating phenotypes using FYPO, alleles and genotypes, protein modifications, genetic and physical interactions, qualitative gene expression, and GO, including associated annotation extensions (Huntley et al. 2014; Lock et al. 2019) and metadata. A PubMed search using a set of S. japonicus-related keywords retrieved 179 publications, which were imported into Canto along with all citation details needed to populate Canto and the website. Publications containing gene-specific data were flagged using Canto’s literature triage function, which allows users to classify papers by type (e.g., curatable, review, methods), and then assign them to authors for full curation. The triage procedure also flagged 91 false positives, most of which were publications describing a “japonicus” species in a genus other than Schizosaccharomyces. We have refined the PubMed search keywords based on these results, so future Canto publication updates will include fewer false positives.

Results

Adding value to imported data

Adding and revising gene structures

The original gene models for S. japonicus were built using aligned RNA-seq transcripts from S. pombe, Schizosaccharomyces cryophilus, and Schizosaccharomyces octosporus orthologous groups compared to protein sequence (Rhind et al. 2011). Gene models with discrepancies were then manually reviewed to provide a set of high-quality gene predictions. However, continual revisions are required to make corrections and keep gene structures in line with published knowledge.
In this work, we used previously published information (Makarova et al. 2016) and detailed orthology inspection to revise 36 gene structures, illustrated in Figure 1. We identified 18 additional conserved genes (SJAG_07000–SJAG_07017) by searching for small, universally conserved proteins absent from the predicted gene set. In eight cases, we replaced erroneous gene predictions with genes correctly identified in an alternative reading frame. Newly identified gene products include Atp19, an F-type ATPase subunit for which no gene structure previously existed (Figure 1A), and the recombination protein Rec7, for which the originally predicted structure was completely replaced by a new structure (Figure 1B). For SJAG_03830, the predicted structure was revised but not completely replaced (Figure 1C), allowing the gene product to be identified as the ubiquinol-cytochrome-c reductase complex subunit Qcr10, previously thought absent from S. japonicus.

Using tRNAscan-SE (Chan and Lowe 2019), we added three genes to the tRNA complement and added product descriptions specifying codons to the 325 nuclear-encoded tRNAs. Additional noncoding RNAs including snRNAs, snoRNAs, and telomerase RNA were imported from RNAcentral (The RNAcentral Consortium 2019).

Identifying distant orthologs

Accurate detection of orthologs, i.e., genes in different species related by vertical descent (Fitch 1970), between two proteomes allows for comprehensive comparative study of the biological processes in two species. Orthologs provide a framework for reconstructing the evolutionary events that have given rise to observed biological differences. Thorough ortholog inventories require the targeted detection of distant orthologs and the identification of the orthologous relationship type (i.e., one-to-one, one-to-many, or many-to-many). Ortholog inventories also facilitate the accurate identification of lineage-specific gene losses, species-specific genes, and protein family expansions.

In this study, we identified 67 previously undetected distant orthologs between S. pombe and S. japonicus, 25 of which are also conserved in human (Supplementary Table S1). These newly detected orthologs include the 18 newly identified small genes noted above, and two genes for which we revised structures. Candidates for a further 17 orthologs (including the kinetochore regulator Meikin, the kinetochore protein Mis19, and the spindle component Dms1) are recorded as high confidence based on a one-to-one relationship between S. pombe and other species and...
syntenic location between \textit{S. pombe} and \textit{S. japonicus}, although sequence similarity has not yet been detected. These regions are currently being reviewed for possible gene prediction errors. Finally, some additional ortholog calls from the automated pipelines were revised to include in-paralogs and remove out-paralogs.

\textbf{Gene names and product descriptions}

\textbf{Gene names}

In consultation with \textit{S. japonicus} researchers, we have assigned standard gene names in JaponicusDB, using unified nomenclature to make conserved loci readily recognizable and to avert potential naming conflicts with other species. Only 101 gene names were imported from UniProtKB for \textit{S. japonicus}. Where a named \textit{S. pombe} gene has a one-to-one ortholog in \textit{S. japonicus}, the gene name was transferred; this method assigned 3471 \textit{S. japonicus} names. Manual curation of 119 exact or near-exact duplicate proteins (ribosomal proteins, histones, translation elongation factors) enabled us to assign names for syntenic orthologs. Finally, 184 gene names were assigned manually to \textit{S. japonicus} specific families, one-to-many, and many-to-one paralogs. Overall, JaponicusDB now provides standard names for 3875 protein-coding genes (of 4886 total). The remit of the fission yeast Gene Naming Committee (GNC; \url{https://www.pombase.org/submitted-data//gene-names}) has expanded to cover \textit{S. japonicus} as well as \textit{S. pombe} gene names; the GNC will approve all new gene names in both species.

\textbf{Gene product descriptions}

Informative gene product descriptions allow users to browse proteomes effectively, scan input and output lists from experiments, and review genome contents. Gene product descriptions manually curated by PomBase were automatically transferred to one-to-one orthologs, providing descriptions for 4251 protein-coding genes. A further 645 descriptions were manually curated based on one-to-many, many-to-one, or many-to-many orthologs or protein family descriptions. JaponicusDB now provides gene product labels that have been reviewed by fission yeast experts, and accurately reflect current knowledge for 4886 protein-coding genes.

Our analysis and curation pipeline have improved gene product descriptions dramatically compared to previously available data. UniProtKB provides descriptions for protein gene products, but over 99% of \textit{S. japonicus} proteins remain in the unreviewed TrEMBL database and therefore have only automated descriptions. The quality of TrEMBL’s inferred descriptions is generally high, but coverage is incomplete. Notably, we have provided informative product labels for 522 proteins described as “uncharacterized protein” in UniProtKB. Many of these are broadly conserved, well-studied proteins identified and annotated via our distant ortholog detection or manual curation pipelines. For example, we found five proteins involved in TOR signaling and three cytochrome oxidase assembly factors.

The JaponicusDB pipeline updates imported gene names and product descriptions upon each run. Simple configuration files maintained in GitHub repositories specify manually curated names and descriptions that take precedence over automated imports.

\textbf{Extending GO annotation coverage and specificity}

To generate a GO (\textit{The Gene Ontology Consortium 2000, 2021}) annotation dataset for \textit{S. japonicus}, we imported computationally generated annotations from the GOA UniProt (\textit{Huntley et al. 2015}), and added new annotations from the PomBase ortholog pipeline, manual inferences for genes without \textit{S. pombe} orthologs, and experimental data from literature manually curated in Canto. We applied the filtering protocol used by PomBase (\textit{Lock et al. 2019}) to retain only relevant and nonredundant automated annotation. Annotations manually curated from the literature in Canto take precedence over any automatically transferred annotations (e.g., as noted in Methods, manual negated annotations suppress import of contradictory automated annotations). The JaponicusDB GO annotation procedures thus support robust transfer of large annotation sets, while also allowing the fine tuning of specific annotation to capture known biological differences (reviewed in \textit{Gu and Oliferenko 2015; Russell et al. 2017; Oliferenko 2018}; also see references cited in \textit{Introduction}). The resulting GO annotation dataset comprises 29,589 annotations as of September 2021.

PomBase maintains subsets of GO, known as “GO slims,” of selected biologically meaningful terms from each branch of GO—Molecular Function (MF), Biological Process (BP), and Cellular Component (CC)—that are used to summarize the functional capabilities of \textit{S. pombe} (\textit{Lock et al. 2019; Wood et al. 2019}). JaponicusDB uses the PomBase GO slims for the same purposes, illustrating the value added by our GO curation procedures. GO slim analysis shows that annotation breadth has significantly increased compared to the GO data originally available from GOA despite an overall decrease in annotation number (from 36,878 to 29,512). For each aspect of GO, more genes are annotated to terms in the GO slim (MF coverage increased from 3087 to 3652, BP from 3238 to 4187, and CC from 2999 to 4578 genes), and the number of genes that either have no annotation or are annotated but not covered by the GO slim, is correspondingly decreased (Figure 2). Notably, we have increased annotation coverage in several areas of active \textit{S. japonicus} research. For example, the annotation increases for “membrane organization” and “lipid metabolism” in BP and “endoplasmic reticulum” (ER) and “mitochondrion” in CC are highly relevant to comparative studies of respiration and membrane and lipid biology. In MF, there is a large increase in annotations to “hydrolase activity,” including 21 gene products annotated to “lipid metabolism.” Newly annotated examples in this set include the mitochondrial cardiolipin-specific phospholipase Clδ1, the acyl-coenzyme A thioesterase Thε4, and SJAG_00199, a mitochondrial DDHD family phospholipase. The relatively new GO term “molecular adaptor activity,” relevant to research in organelle organization, also has increased annotations including many newly described protein- and organelle-to-membrane adaptors.

Annotation depth, defined as increased distance from the root node in an ontology graph, provides a measure of the biological specificity of the annotations. Our work has increased annotation depth in at least one GO aspect (MF, BP, or CC) for 4312 proteins (>88% of the total), and in all three aspects for 1385 (>28%, Figure 3).

\textbf{Manual literature curation in Canto}

Using Canto’s literature triage function, we have manually classified the 179 publications found in our initial PubMed query (see Methods). After discarding spurious matches, we identified 31 papers containing gene-specific data suitable for curation, as well as 39 papers in other categories. The latter reflect the status of \textit{S. japonicus} as an emerging model and include publications describing wild-type features and cell composition, methods and
reagents, or phylogenetic studies, as well as reviews. The 31 curatable papers were assigned to the authors (or to a Canto administrator) for curation, and 23 are now fully curated in JaponicusDB (see https://www.japonicusdb.org/reference_list/community). To date, manual curation has supplied 135 experimentally supported GO annotations and 168 phenotype annotations.

**Website**
The JaponicusDB website, modeled on PomBase, provides convenient access to all molecular and cell biological data curated for S. japonicus. Like PomBase, JaponicusDB includes pages for each gene, publication, and genotype annotated, as well as for ontology terms. JaponicusDB also uses the same simple, advanced, and peptide motif search tools as PomBase. To facilitate comparison between S. japonicus and S. pombe, the “Ortholog” section of each gene page includes reciprocal links between PomBase and JaponicusDB. On both sites, orthologs can also be retrieved via the advanced search or via an ID mapping tool. The JaponicusDB front page (Figure 4) presents news, database usage hints, publications recently curated by the community, and “Research Spotlight” panels, which feature graphical abstracts from recent papers (one of the most popular features of PomBase). An instance of the genome browser JBrowse (Buels et al. 2016) currently displays the S. japonicus genome and annotated features, and will host any available high-throughput sequence-based datasets. Online documentation describes all page contents, search capabilities, and how to contribute to JaponicusDB (also see Lock et al. 2019 and Harris et al. in this issue). JaponicusDB uses Google Analytics to monitor website usage.

**Conclusions**
For several decades, fission yeast research has been a mainstay of progress in cell and molecular biology, due to pioneering work

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**Figure 2** Effect of curation on S. japonicus GO annotation coverage. Comparison of the distribution of annotations for all S. japonicus protein-coding genes (4896 total) to selected high-level terms in each aspect of GO between the UniProt GOA file (“before”, left-hand columns) and the JaponicusDB GO annotation set (“after”, right-hand columns). The number of annotated proteins has increased for most MF, BP, and CC, with associated decreases in the proportions of proteins annotated as “unknown.” Highlighted blocks represent GO terms relevant to processes that are intensively studied in S. japonicus: “gene expression,” “membrane organization,” and “lipid metabolism” in BP; “endoplasmic reticulum” and “mitochondrion” in CC, and “hydrolase activity” and “molecular adaptor activity” in MF. Images were generated using QuiLT (Harris et al., in this issue), in which only one term per gene can be included for display. If a gene is annotated to more than one GO term, one is selected according to a set order of precedence. Here, terms are arranged by order of precedence in the charts and key for each GO aspect.

**Figure 3** Schizosaccharomyces japonicus proteins with increased GO annotation specificity. GO annotation specificity, defined as distance from the ontology root, was measured for each aspect of GO and for each of the 4896 proteins in S. japonicus. The Venn diagram shows the number of proteins that have more specific annotations in the JaponicusDB GO annotation set than in the UniProt GOA file for one or more GO aspect(s). MF, Molecular Function; BP, Biological Process; CC, Cellular Component.

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on cell cycle regulation, as well as recent innovative experimentation in additional areas such as chromosome segregation, epigenetics, and cytokinesis, using *S. pombe*. The growing amount and sophistication of research on *S. pombe*’s “sister” species *S. japonicus* provide a prime opportunity to add evolutionary biology to the roster of topics that can be fruitfully investigated using fission yeast.

We have launched JaponicusDB to enable *S. japonicus* researchers—and the wider scientific community—to realize the full promise of experiments in an emerging model system. JaponicusDB gathers and interconnects diverse types of information from many sources to create a single resource that provides coherent, intuitive access to rich, interconnected data sets. As a fully functional MOD, JaponicusDB supports robust genome-wide data interrogation and mining at any level of detail, experimental planning, hypothesis generation, and interpretation of results, thereby meeting research needs not addressed by any other system.

**PomBase code reuse**

We have reused the modular, configurable open-source code underlying PomBase, enabling us to deploy a curation database, a Canto instance, and a functioning, intuitive website for JaponicusDB with modest developer and curator input. (We estimate that essential development took approximately one full-time equivalent (FTE) for 6 weeks, and PomBase curators devoted a similar amount of time to curation, i.e., 12 FTE-weeks total for all tasks except the literature curation contributed by the community.) For updates, the PomBase system relies primarily on data and configuration files stored (with version control) in GitHub repositories, making future maintenance straightforward. Because JaponicusDB uses PomBase code, both databases will simultaneously deploy any new features developed by the PomBase team.

**Community literature curation**

The Canto community literature curation system enables bench biologists to contribute directly to the knowledge integrated into JaponicusDB. Although Canto is also being used for phenotype curation in FlyBase (Larkin et al. 2021) and pathogen–host interaction phenotypes in PHI-Base (Urban et al. 2020), JaponicusDB has embraced Canto for community curation at a uniquely early stage in its development. The curatable literature corpus is small at present, but over two-thirds of suitable papers have already been curated. *S. japonicus* researchers can use community curation to meet data dissemination objectives, helping ensure timely FAIR (Findable, Accessible, Interoperable, and Reusable; Wilkinson et al. 2016) sharing of new information. To enhance engagement with JaponicusDB and interactions among community...
Community MOD management

To ensure that JaponicusDB meets its users’ needs in the future, we will invite experts in relevant areas, including S. japonicus biology and database curation and maintenance, to form a Scientific Advisory Board (SAB). The SAB will collaborate with PomBase staff to mobilize volunteers from the S. japonicus community to carry out routine updates and maintenance, and will guide future fission yeast MOD developments that arise from S. japonicus research. PomBase staff will provide training and advice to community volunteers in configuring the website and JBrowse track metadata, Canto administration, editing gene structures, and other maintenance tasks. This group will also arrange to disseminate data regularly from JaponicusDB to external resources, e.g., GO annotations to the GO Consortium repository.

Curation to support S. japonicus research

Our review of the S. japonicus genome highlights the value that manual curation adds to sequence feature annotation, ortholog identification, and functional annotation, and therefore to all subsequent usage of these data in experimental and predictive studies. As a result, JaponicusDB can provide more comprehensive, experimentally informed complements of several types of genome-scale data than were previously available via data aggregators. We note that finding genes previously thought to be absent from S. japonicus has significant consequences for accurately representing molecular and cellular processes within the species as well as comparative studies of how these processes have evolved.

Gene structures for most genomes are derived by automated prediction pipelines, often incorporating information from homology and synteny. Although multiple methods are frequently used, many errors will persist even for intensively curated model species in which all gene predictions were manually inspected prior to publication. For example, since publication of the S. pombe genome, PomBase has revised >200 gene structures; similarly, WormBase has updated several thousand for Caenorhabditis elegans (P. Davis, personal communication, WormBase). Every genome-scale analysis depends upon comprehensive, manually refined gene structures (and often other sequence features) for accuracy, reproducibility, and relevance.

Our work also illustrates how biologists may benefit directly from manual ortholog curation. For example, we have identified S. japonicus orthologs of cytochrome c oxidase subunits Cox7 and Cox9, ATP synthase subunit Atp19, mitochondrial alphaketo glutarate dehydrogenase Ymr1, and ubiquinol-cytochrome-c reductase complex subunit Qcr10, all of which are important for fission yeast researchers studying the differences in central carbon metabolism between S. pombe and S. japonicus.

Future updates to ortholog predictions, gene structures, names, and product descriptions, are bound to reveal more examples where comprehensive, curated knowledge is indispensable for understanding biological processes, avoiding misinterpretation, and making reliable cross-species comparisons.

Rapid MOD deployment to support emerging model species

The rapid establishment of JaponicusDB promises to have beneficial repercussions well beyond the fission yeast community. First, simply having a full S. japonicus MOD available to facilitate and support research makes reliable information available to underpin comparative studies and data integration, not only between S. japonicus and S. pombe but also throughout all eukaryotes.

Perhaps of even greater consequence, JaponicusDB represents a proof of concept, demonstrating that a small group of people can easily deploy PomBase code to produce a comparable system for any other species. We and others have noted (Oliver et al. 2016; Alliance of Genome Resources Consortium 2019; Lipshutz 2021) that complete genome sequences continue to accumulate, but annotation even of sequence features, let alone any other associated data, cannot keep pace. A growing number of increasingly diverse research communities require creative approaches to data infrastructure to take advantage of the exploratory opportunities that a genome sequence offers, and to integrate genome-scale molecular data and methods with small-scale results from new or extant published literature. Our experience with JaponicusDB exemplifies one way forward: the PomBase system can be redeployed by a small, motivated community to create a resource that can be maintained within the scope of time and skills available to researchers, using funding allocated to data dissemination.

Data availability

As described in detail above, JaponicusDB uses the open-source PomBase code base available from the PomBase GitHub organization (https://github.com/pombase). JaponicusDB configuration and data files are available from the JaponicusDB GitHub organization (https://github.com/japonicusdb).

JaponicusDB data can be viewed directly at https://www.japonicusdb.org, query results provide data download options, and data can be downloaded in bulk from the website (see https://github.com/japonicusdb.org/datasets and https://www.github.com/japonicusdb.org/data).

Supplementary material is available at GENETICS online.

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Conflicts of interest

The authors declare that there is no conflict of interest.
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