Ensembl Genomes 2022: an expanding genome resource for non-vertebrates

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Received September 21, 2021; Revised October 07, 2021; Editorial Decision October 08, 2021; Accepted November 10, 2021

ABSTRACT

Ensembl Genomes (https://www.ensemblgenomes.org) provides access to non-vertebrate genomes and analysis complementing vertebrate resources developed by the Ensembl project (https://www.ensembl.org). The two resources collectively present genome annotation through a consistent set of interfaces spanning the tree of life presenting genome sequence, annotation, variation, transcriptomic data and comparative analysis. Here, we present our largest increase in plant, metazoan and fungal genomes since the project’s inception creating one of the world’s most comprehensive genomic resources and describe our efforts to reduce genome redundancy in our Bacteria portal. We detail our new efforts in gene annotation, our emerging support for pangenome analysis, our efforts to accelerate data dissemination through the Ensembl Rapid Release resource and our new AlphaFold visualization. Finally, we present details of our future plans including updates on our integration with Ensembl, and how we plan to improve our support for the microbial research community. Software and data are made avail-

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able without restriction via our website, online tools platform and programmatic interfaces (available under an Apache 2.0 license). Data updates are synchronised with Ensembl’s release cycle.

INTRODUCTION

Ensembl Genomes (https://www.ensemblgenomes.org) provides access and analysis for non-vertebrate genomes across the domain of life. It is organised around the five kingdoms of life: plants (https://plants.ensembl.org), invertebrate metazoans (https://metazoa.ensembl.org), fungi (https://fungi.ensembl.org), protists (https://protists.ensembl.org) and bacteria (https://bacteria.ensembl.org). These five resources complement the Ensembl project (1) (https://www.ensembl.org), whose focus is vertebrate metazoans and model organisms.

As previously reported, we provide high-quality annotated genome assemblies, integrate and link with other complementary genome resources, represent genomic diversity and deliver a comprehensive analysis platform (2). We provide secondary analysis platforms including whole genome pairwise and multiple sequence alignment, homology prediction and transcriptomic analysis, ontology-based gene annotations and pathway associations. Our secondary analyses are enabled by a shared data representation and infrastructure with Ensembl, meaning tools and analysis methods developed for vertebrates are compatible with the non-vertebrate genomes with minimal, or no, modification required.

All genome assemblies are imported from the International Nucleotide Sequence Database Collaboration (INSDC) (3). Only INSDC accessioned sequences are hosted as part of our joint browser agreement with NCBI (4) and UCSC(5). We also import variation data sets from the European Variation Archive (EVA) (https://www.ebi.ac.uk/eva/) and provide automated alignment of plant transcriptome data as submitted to the European Nucleotide Archive (ENA) (6) through our collaboration with Expression Atlas (7). Our resources are further enhanced by our active collaborations with other major non-vertebrate genome providers including Gramene for plant genomes of crops, models, and species of evolutionary importance (8), VEUPathDB for eukaryotic pathogens (9) and invertebrate vectors of disease-causing pathogens (10), WormBase providing for nematodes and flatworms (11) and PHI-base for manually curated pathogen-host interactions (12).

Genomes can be accessed via one of our dedicated taxonomic websites or through the Ensembl Rapid Release resource (https://rapid.ensembl.org). All Ensembl sites provide genome browsing functionality; a way to explore the spatial relationships between annotated genomic elements. Functional annotation of genes, transcripts and proteins are enabled through imports of UniProt curated functions (13), imputation from sequence analysis tools such as InterProScan (14) and imports of manual curation of host-pathogen interactions from PHI-base. We provide comparative genomic analysis including whole genome alignments and gene orthology prediction (available for all eukaryotic taxonomic divisions), a pan-taxonomic gene orthology prediction covering key species across the tree of life and PANTHER based classification of bacterial gene families (15). Search and BLAST is available for all genomes (16). A public MySQL database server, Perl and RESTful Application Programming Interfaces (APIs) (https://rest.ensembl.org), BioMart (17) and bulk access flat-files (ftp.ensemblgenomes.org) is available for all genomes hosted in our taxonomic sites. Genomes can be analysed with standard Ensembl tools such as the Ensembl Variant Effect Predictor (VEP) (18). Each taxon-specific website is archived once per year with releases 45 (e.g. https://eg45-plants.ensembl.org/) and 49 (e.g. https://eg49-plants.ensembl.org/) being nominated for archive in 2019 and 2020, respectively. Genomes provided via Rapid Release, described later, only have a genome browser, minimal functional data imports, BLAST and flat-file access via Ensembl’s FTP site (ftp.ensembl.org/pub/rapid-release/species/). All data generated by Ensembl Genomes are available for use without restriction.

Since our last review, we have seen one of the largest increases in eukaryotic genomes available through our platform with over 500 new species. As the number of genomes increased, we have had to adapt both our infrastructure and analyses to ensure scalability, continue to provide world-class genomic annotations and make available new data visualizations. Below, we highlight the new genomes and features that have been introduced over the last two years.

NEW AND IMPROVED GENOMES

The past two years have seen significant increases in our plant, metazoan and fungal genome collections (see Table 1), totalling 588 additional genomes. We have expanded our taxonomic breadth of plants, which now includes asterids (e.g. sesame, lettuce), grasses (barley and wheat cultivars) and Brassicaceae (false flax and alpine rock-cress). Thirteen tree genomes have been added including Pistacia vera (pistachio), Olea europaea (olive tree), Corylus avellana (common hazel), Eucalyptus grandis (eucalyptus) and Quercus lobata (Valley Oak). Many of these species have a long generation time, as in the case of Corylus avellana (hazel) which takes up to eight years to reach full productivity (19). Analysing and integrating these trees has required novel method development due to their genome size and complexity and is detailed later.

Our metazoa resource has added sets of new or improved assemblies for pathogenic disease vectors including Aedes aegypti (vector for yellow fever, zika and chikungunya), Anopheles coluzzii (vector for malaria), Phlebotomus papatasi (vector for leishmaniasis), six species of the Glossina complex (vector for sleeping sickness) (20) and the livestock pest Stomoxys calcitrans (21). Our twelve hosted Drosophila fly genomes have been refreshed to mirror those in FlyBase (22). Six strains of Bemisia tabaci, a cassava insect pest, are now available through our collaborative work with the African Cassava Whitefly Project (http://www.cassavawhitefly.org/) (23). Similarly, our collaboration with the Marine Invertebrate Models Database (MARIMBA) and CORBEL has brought two new marine metazoan genomes; Actinia equina (beadlet anemone) and Clytia hemisphaerica (a cnidarian). We also host a selec-
tion of well-studied nematode and flatworm genomes from the WormBase ParaSite project (https://parasite.wormbase.org) to enrich our comparative analysis. These include Caenorhabditis elegans; five other Caenorhabditis; parasites of humans and livestock including Brugia malayi (lymphatic filariasis) and Loa loa (African eye worm). Our fungal genomes coverage has increased significantly due to a new public archive import and 15 genomes originating from VEuPathDB’s fungal database, FungiDB, making Ensembl Fungi the most comprehensive collection of free/open access fungal genomes.

We chose to freeze our protists collection as we switched our focus towards identifying redundant genomes in our bacterial collection. We have adopted UniProt’s prokaryotic proteome redundancy definitions, which removes closely related genomes based on the protein coding content (24). UniProt’s methodology first creates a directed weighted graph of proteome similarity based on proteome content, taxonomic filtering and proteome size. It then finds the dominating set by repeatedly removing the weakest nodes until no more removals are possible. Adopting this approach has resulted in the removal of 12 716 genomes, whilst maintaining the coverage of 527 known bacterial families (Figure 1A and B). Cross referencing the removed genomes against NCBI’s family classification showed reductions in the Streptococcaceae (–5367), Enterobacteriaceae (–5278), Staphylococcaceae (–4877) and Mycobacteriaceae (–3547) families showing a previous over-representation in well studied bacterial families (Figure 1A). We also observed an increase of 957 genomes with no assigned taxonomy at the family level, raising the percentage of unclassified bacteria to ~10% (Figure 1C). All removed genomes remain accessible from our release 49 Ensembl Bacteria archive and FTP site. Further details can be found in our blog (https://www.ensembl.info/2020/09/21/ensembl-bacteria-updates/).

Table 1. Ensembl non-vertebrate growth/update 2019–2021

| Release | Date         | Bacteria | Protists | Fungi | Plants | Metazoa |
|---------|--------------|----------|----------|-------|--------|---------|
| 45      | September 2019 | 44 048   | 237      | 1014  | 67     | 78      |
| 52      | October 2021  | 31 332   | 237      | 1505  | 119    | 123     |
| Change  |              | -12 716  | 0        | +491  | +52    | +45     |

SCALING GENOME RESOURCES

In response to the recent increases in non-vertebrate genomes, we identified a need to accelerate researcher access to emerging data sets and scale our infrastructure to meet that demand. In 2020, Ensembl provided the ‘Ensembl Rapid Release’ website to support large-scale biodiversity studies and enables annotation release every two weeks, in contrast to its three-month integrated release cycle. Clytia hemisphaerica and Actinidia equina were the first non-vertebrates to be made available via rapid release in 2020 and have been joined by Vigna unguiculata (black-eyed pea), Cajanus cajan (pigeon pea) and Digitaria exilis (fonio millet) representing crops of agricultural importance. We also redesigned our portal site (https://www.ensemblgenomes.org) to streamline user access to key genomes, switch our technology to the static site builder eleventy.js and to provide a new dynamic text search enabled by the EBI Search API (29).

SUPPORTING PANGENOMES

Pangene adoption is a growing area of interest and is considered a credible solution to reference biases and missing elements of a single reference genome. One such case is in Triticum aestivum where 12 150 genes were found to be missing from the reference assembly of the variety Chinese Spring Wheat, but were found in at least one of the 18 re-sequenced modern varieties (30, 31). To better model the wheat pangenome, we added nine new chromosome-scale wheat lines, alongside five additional scaffold-level assemblies published as part of the 10+ wheat genome consortium (http://www.10wheatgenomes.com/). Each assembly can be viewed individually via our genome browser or using our cultivar view, which reuses visualization views originally developed for mouse strains.

Generating high quality whole genome alignments is a key component in creating graph genomes. In preparation
for increasing our pangenome support, we benchmarked Ensembl’s existing whole genome aligner Enredo-Pecan-Ortheus (EPO) (32) against a set of 11 *Oryza* (rice) assemblies (Figure 2).

**LINKING GENOMES TO PREDICTED 3D STRUCTURE**

AlphaFold (33) has been a revolutionary advancement in 3D protein structure prediction and the release of AlphaFold DB in July 2021 (Varadi et al. in preparation) made available predictions across 17 non-vertebrate species providing previously unimaginable 3D proteome coverage. In the case of *Arabidopsis thaliana*, PDBe (34) contains 1661 experimental structures compared to 27434 predicted structures available from AlphaFold DB. We used *A. thaliana* as a test for integration due to the availability of high-quality variant data and shared identity between ourselves and UniProt’s reference proteome. We have successfully integrated AlphaFold models, visualized via Mol* (35), with exon and protein altering SIFT scored variants (Figure 3) (36). This view is available from our protein information page. We plan to expand our coverage to all available
Figure 2. EPO multiple genome alignment visualization of chromosome 1 in three rice genomes: Oryza sativa indica Group (top), Oryza sativa japonica Group (middle) and Oryza glaberrima (bottom). Orange discontinuous blocks represent the areas of alignment across all three genomes. Each genome displays its genes and can be used to identify regions of uniqueness in each genome and identify potential areas of mis-assembly or mis-annotation. This alignment can be browsed at http://plants.ensembl.org/Oryza_nivara/Location/Compara_Alignments/Image?align=9910;db=core;r=1:586653--632276.

AlphaFold DB proteomes where possible across the non-vertebrate domain.

PUBLIC ENGAGEMENT, OUTREACH AND TRAINING

We continue to offer training on our tools, interfaces, and APIs conducted virtually during the COVID-19 pandemic via video conferencing platforms. These platforms are used alongside participant interaction tools such as living documents (an open Google Doc where participants can type questions and answers), real-time messaging services, e.g. Slack and interactive polling software, e.g. Slido ensuring participants have multiple methods to communicate with trainers. There was also the return of the annual Wellcome Advanced course on fungal pathogen genomes co-developed with Wellcome, FungiDB, JGI and SGD (Saccharomyces Genome Database) and conducted virtually this year, after a break in 2020. A key teaching point of this course is the effective piecing together of features from multiple fungal resources to find the best answer to a biological question.

Finally, the global pandemic has also made public engagement increasingly difficult as social distancing requirements have challenged in-person interaction. Working in conjunction with the Cambridge University Botanic Garden, we co-developed ‘DNA in the Garden Trail’; a unique COVID-safe self-guided tour around the botanic gardens with a focus on plants hosted in Ensembl Plants (https://www.botanic.cam.ac.uk/education-learning/trails/dnatrail/). A companion application was built using Guidemap and offered a plant genomics quiz for families to complete as they used the trail.

SOFTWARE ANALYSIS RESOURCES

Over the past two years, we have released two new non-vertebrate software resources; a collection of Ensembl Genomes data production workflows (https://github.com/Ensembl/ensembl-production-imported) and a set of analysis scripts for Ensembl Plant genomes (https://github.com/Ensembl/plant-scripts). These scripts are provided in a number of programming languages (Python, R, Perl) and detail common tasks using our programmatic interfaces and databases. We also released a de novo repeat analysis method for plant genomes, which uses a combination of repeat finders, repeat libraries, such as RepBase (37) or REDat (38), Red (39) and a curated set of transposable elements from a well characterized set of plants enabling fast and accurate annotation of new genomes (40). These new methods help to maintain sustainable genome analysis through accurate annotation of repetitive sequence.
**Figure 3.** An AlphaFold 3D prediction for the Arabidopsis thaliana protein Q00958 (LFY: AT5G61850.1) displayed as a Richardson model using Mol*. The central panel annotates the model with regions of high confidence (blue) to low confidence (orange) with its protein sequence displayed above. The right hand panel enables highlighting of one or more exons, variants and protein features which are controlled by clicking on the eye icon. Variants can be turned on/off according to how deleterious or tolerated they are or individually. Only variants resulting in protein changes with SIFT scores are made available for display.

**FUTURE PLANS**

Whilst most of our annotation comes from third party imports, we have grown our ability to annotate a diverse range of non-vertebrate genomes in-house. Ensembl Rapid Release has been a vital component of this strategy enabling fast dissemination and is becoming our preferred method of distribution for newly annotated genomes. We plan to continue annotating non-vertebrate genomes in-house, expand data types available via Rapid Release and release a new scalable homology prediction method in collaboration with Ensembl. Genomes will still flow into our taxonomic sites based on their importance, scientific interest, broadening of our comparative analyses and when in-line with Ensembl’s strategy for genome inclusion.

Continued growth in bacteria genomes necessitates a different strategy to handle duplication, inconsistencies of annotation and prioritize the needs of microbial researchers (41). As mentioned previously, 10% of bacteria lack a taxonomy, and this coupled with the continued growth in bacterial genomes derived from both isolate and environmental source will require further deployment of dereplication methods such as those used by Genome Taxonomy Database (42) to ensure our resources represent the breadth of bacterial diversity, yet continue to scale. Many newly submitted genomes lack gene annotation, and those that do have annotation can be of varying quality and/or show other issues e.g. inconsistent gene naming. To overcome these issues, we plan to re-annotate our hosted bacterial genomes and enrich them with functional annotations such as pathways and secondary metabolite gene clusters. We will also maintain existing annotations on key community reference genomes, e.g. *Escherichia coli K12* (U00096.3). Consistent high-quality annotation is key to enabling better downstream analysis such as developing new methods for deeper/broader functional annotation using machine learning. Our collaboration with MGnify (43) – EMBL-EBI’s metagenomics resource - will continue to expand. Briefly, we will focus on harmonizing the bacteria in Ensembl Genomes with the metagenome assembled genomes (MAGs) available in MGnify, through the adoption of common annotation pipelines, utilization of similar methods for the removal of genomic redundancy, i.e. GTDB (44, 45), and application of the same web presentation layers in both resources, making it easier for users to transition between the two resources. We will use the collections of Ensembl (isolate) genomes and MAGs as reference databases for determining their presence in metagenomes, to better
understand the biological environments these genomes are found in. As the range of microbes presented in MGnify expands beyond prokaryotic microbes, we anticipate further synergies. Part of this effort will involve improving our coverage of protists in Ensembl Genomes.

Our efforts to merge Ensembl and Ensembl Genomes resources continues within the context of Ensembl’s new infrastructure and website project (https://2020.ensembl.org). Of the seven genomes available through the new infrastructure, five are non-vertebrates; *Triticum aestivum*, *Caenorhabditis elegans*, *Saccharomyces cerevisiae*, *Plasmodium falciparum* and *Escherichia coli* K12. This puts non-vertebrate genomes at the centre of our future strategy, reflecting the increasing popularity of these genomes. Our efforts to reuse the rapid release infrastructure underlines that this strategy is not only possible but will create a better experience for researchers. We encourage those interested in shaping the future of this site to give feedback via our helpdesk and to sign up to our user experience sessions.

Finally, we expect significant progress in our support for pangenomes both in data processing and visualization. We plan to utilize our multiple sequence alignment methodology to construct genome graphs of rice and wheat cultivars and to utilize our multiple sequence alignment methodology to visualize pangenome analysis and visualization.

ACKNOWLEDGEMENTS

We thank the following Ensembl project members for their work, which underpins our own: Jamie Allen, Jorge Alvarez-Jarreta, Irina Armean, Olaranwaju Austine-Orimoloye, Konstantinos Billis, Sanjay Boddu, Lucy Brooks, Mehrnaz Charkhchi, Carla Cummins, Kamal Kumar Dodiya, Bilal El Houdaigui, Carlos Garcia Giron, Thiago Genez, Arthur Gymer, Thibaut Houflier, Thomas Juettemann, Ilias Lavidas, Diana Lemos, José Carlos Marugán, Shamika Mohanan, Tamara El Naboulsi, Marc Naven, Denye N. Ogeh, Anne Parker, Andrew Parton, Ivana Pilzota, Mira Prosovetkskaia, Helen Schuilenburg, William Stark, Kyösti Sutinen, Anja Thormann, Francesca Tricomi, David Urbina-Gómez, Andres Veidenberg, Thomas Walsh, Brandon Walts, Natalie Willholt, Andrea Winterbottom, Bethany Flint, Stefano Giorgetti, Leanne Haggerty, Sarah Hunt, Garth Ilsley, Fergal Martin, Magali Ruffier, David Thybert, Peter W. Harrison and Daniel Zerbino. We also thank Mandar Deshpande, Mihaly Varadi and Sameer Velankar for their help in enabling AlphaFold visualisation. Finally, we thank Chantal Helm, Angela Cano, Mark Danson, James Blackshaw, Alexandra Canet and Susan Wallace for their contributions to the 'DNA in the Garden Trail'. Ensembl and Ensembl VEP are trademarks of EMBL. For the purpose of open access, the author has applied a CC BY public copyright licence to any Author Accepted Manuscript version arising from this submission.

FUNDING

UK Biosciences and Biotechnology Research Council [BB/P024602/1 to F.H.R., T.L., BB/P016855/1, BB/S02011X/1, BB/M028372/1, BB/P027849/1, BB/S020020/1 to A.C., K.E.H-K, J.S. M.U.; BB/T015691/1; Ensembl-4-Breeders]; Wellcome Trust [108749/Z/15/Z, 201535/Z/16/Z, 222155/Z/20/Z]; UK Medical Research Council [MR/S000453/1]; National Science Foundation [IOS-1127112 to K.C., J.E., P.G., P.J., V.K., S.K., S.N., A.O., J.P., M.K.T-R, D.W., S.W.]; United States Department of Agriculture [8062-21000-041-00D to D.W.]; Bill and Melinda Gates Foundation [B03436X13]; ELIXIR [FONDUE, ‘Apple as a Model for Genome Information Exchange’]; European Molecular Biology Laboratory. National Human Genome Research Institute of the National Institutes of Health [U24HG002223]; National Institute of Allergy and Infectious Diseases, National Institutes of Health, Department of Health and Human Services [5N593019C00077]; the content is solely the responsibility of the authors and does not represent the views of the National Institutes of Health; European Union’s Horizon 2020 Research and Innovation Programme [731060, 654248]; the DNA in the garden trail was funded by the Wellcome Connecting Science Enabling Fund. Funding for open access charge: European Molecular Biology Laboratory.

Conflict of interest statement. Paul Flicek is a member of the Scientific Advisory Boards of Fabric Genomics, Inc. and Eagle Genomics, Ltd.

REFERENCES

1. Howe,K.L., Achuthan,P., Allen,J., Allen,J., Alvarez-Jarreta,J., Amode,M.R., Armean,I.M., Azov,A.G., Bennett,R., Bhai,J. et al. (2021) Ensembl 2021. Nucleic Acids Res., 49, D884–D891.
2. Howe,K.L., Contreras-Moreira,B., De Silva,N., Maslen,G., Akanni,W., Allen,J., Alvarez-Jarreta,J., Barba,M., Bolser,D.M., Cambell,L. et al. (2020) Ensembl Genomes 2020—enabling non-vertebrate genomic research. Nucleic Acids Res., 48, D689–D695.
3. Arita,M., Karsch-Mizrachi,I. and Cochrane,G. (2021) The international nucleotide sequence database collaboration. Nucleic Acids Res., 49, D212–D214.
4. Resource Coordinators,NCBI, Agarwala,R., Barrett,T., Beck,J., Benson,D.A., Bollin,C., Bolton,E., Bourexis,D., Brister,J.R., Bryant,S.H. et al. (2018) Database resources of the National Center for Biotechnology Information. Nucleic Acids Res., 46, D8–D13.
5. Fujita,P.A., Rhead,B., Zweig,A.S., Hinrichs,A.S., Karolchik,D., Cline,M.S., Goldman,M., Barber,G.P., Clawson,H., Coelho,A. et al. (2011) The UCSC Genome Browser database: update 2011. Nucleic Acids Res., 39, D676–D682.
6. Harrison,P.W., Ahamed,A., Aslam,R., Akalo,B.T.F., Burgin,J., Buso,N., Courtot,M., Fan,J., Gupta,D., Haseeb,M. et al. (2021) The European Nucleotide Archive in 2020. Nucleic Acids Res., 49, D82–D85.
7. Papadopedalou,I., Moreno,P., Manning,J., Fuentes,A.M.-P., George,N., Feoxva,S., Fonseca,N.A., Fulligrabe,A., Green,M., Huang,N. et al. (2019) Expression Atlas update: from tissues to single cells. Nucleic Acids Res., 48, D77–D83.
8. Tello-Ruiz,M.K., Naithani,S., Gupta,P., Olson,A., Wei,S., Preece,J., Jiao,Y., Wang,B., Chougule,K., Garg,P. et al. (2021) Gramene 2021: harnessing the power of comparative genomics and pathways for plant research. Nucleic Acids Res., 49, D1452–D1463.
9. Aurrecoechea,C., Barreto,A., Basenko,E.Y., Brustelli,J., Brunk,B.P., Cadé,S., Crouch,K., Doherty,R., Falke,D., Fischer,S. et al. (2017) EuPathDB: the eukaryotic pathogen genomics database resource. Nucleic Acids Res., 45, D581–D591.
10. Giraldo-Calderón,G.I., Emrich,S.J., MacCallum,R.M., Maslen,G., Dalyanas,E., Topalis,P., Ho,N., Gesing,S. and the VectorBase Consortiumthe VectorBase Consortium and Madey,G. et al. (2015) VectorBase: an updated bioinformatics resource for invertebrate vectors and other organisms related with human diseases. Nucleic Acids Res., 43, D707–D713.

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