One of the world’s largest living organisms can be found in a forest, but its presence can only be detected by the appearance of fruiting bodies, better known as honey mushrooms, around infected trees. Armillaria fungi are among the most devastating fungal pathogens; the fungus Armillaria ostoyae was sequenced and analyzed by the JGI to better understand the evolution of Armillaria’s abilities to spread and infect, and effectively break down all components of plant cell walls.

(Armand Robichaud, Flickr CC BY-NC 2.0)

Impact Section Case Study credits (in order of appearance):
Laccaria bicolor (Francis Martin, INRA); Postia placenta (Courtesy of Tom Kuster, FPL); Scheffersomyces stipitis (Courtesy of Tom Jeffries, University of Wisconsin, Madison); and, Daldinia eschscholzii (Petra Gloy CC BY-NC-ND 2.0)

All photography, unless otherwise noted, by Marilyn Chung, Berkeley Lab.
Table of Contents
Aerial view looking toward Berkeley Lab (middle left), with JGI near the base of Mount Diablo. (Jeremiah Higgins, Unsplash)
The U.S. Department of Energy (DOE) Joint Genome Institute’s (JGI) mission is to serve the diverse scientific community as a national user facility, by making available large-scale genomics and analysis of plants, microbes, and communities of microbes. This addresses the DOE mission goal of harnessing science and technology to address energy and environmental challenges.
Director's Perspective

Nigel Mouncey, Director, DOE Joint Genome Institute
Continuing Evolution

In 2017, the JGI continued its evolution towards an Integrative Genome Science User Facility that provides a diverse scientific community with access to state-of-the-art technologies and scientific expertise, enabling biological discoveries and applications in the mission areas of bioenergy, nutrient cycling, and biogeochemistry.

The JGI offers a suite of capabilities unique in their ability and scale to advance energy and environmental science. Collectively, the JGI’s investments in infrastructure and data management, computational tool development, and cutting-edge approaches to data mining have paved the way to translate large-scale and diverse data into biological discoveries. The JGI identifies grand scientific challenges and provides the leadership, tools, expertise and resources to address them. I joined as the JGI’s new Director in March 2017, bringing with me almost 20 years’ experience in conducting and managing research programs and research and development organizations in the industrial biotechnology private sector, most recently at Dow AgroSciences. It is the tremendous science, expertise and the ability to advance scientific knowledge that attracted me to the JGI and Lawrence Berkeley National Laboratory’s (Berkeley Lab) Biosciences Area.

I would like to recognize the outstanding leadership of Axel Visel, who served as Interim Director from March 2016 until my arrival.

Over the last year, the JGI has continued to successfully grow its user and science programs, in alignment with its scientific directions. We are developing a wide range of large-scale experimental and computational capabilities to link sequence to biological insights whose targets are largely driven by systematic (phylogenetic) and problem-focused (e.g., plant-microbe interactions) approaches, as well as large-scale functional studies. A major continued thrust at the JGI is to provide integrated capabilities that, for example, allow users to mine genomic databases, design and synthesize constructs, introduce these into suitable hosts, and determine gene/pathway function through metabolomics. Seamless integration of these capabilities provides users with a powerful suite of technologies that offer value far beyond their individual components. Significant progress has been made towards the two- and five-year milestones defined in the 2016 Strategic Planning Update, with 63% of the two-year milestones now completed.

As a User Facility, in fiscal year (FY) 2017 the JGI served a worldwide community of 1,598 users from academia, government, and industry. In support of their projects, we produced nine new and 2,646 resequenced plant genomes, 194 annotated fungal genomes, and 1,826 microbial genomes (including single-cell draft genomes), as well as 1,291 metagenomic data sets. As we continue to advance our functional genomics, we have generated 1,523 plant transcriptomic and 511 metatranscriptomic datasets. In total, the JGI produced a record sequencing output of 178 trillion bases of sequence data in FY17 alone. Additionally, our DNA Synthesis Science Program produced six million bases of DNA. Data from the sequencing projects are disseminated to the broader scientific community via the respective Science Program data portals, as well as the JGI’s genome portal. These data platforms continue to attract new users: more than 11,000 new portal users accessed our data portals in FY17, and more than 100,000 visitors came for data access and analysis.
The demand for our user programs is also increasing. In 2017, we received 76 proposals submitted to the large-scale Community Science Program (CSP), as well as 153 proposals in response to the small-scale microbial/metagenome and DNA synthesis CSP calls, and 51 proposals to the joint Facilities Integrating Collaborations for User Science (FICUS) call with the Environmental Molecular Sciences Laboratory (EMSL) at Pacific Northwest National Laboratory (PNNL). FICUS is a user program collaboratively offered by the JGI with, as of 2017, two other national user facilities: EMSL and the National Energy Research Scientific Computing Center (NERSC) at Berkeley Lab. This Microbiome Data Science FICUS provides for the first time a service that is based on data rather than traditional products (e.g., sequencing). Across all of these programs, a total of 108 proposals were approved and accepted after peer review and are now in progress.

The impact of the research performed through all of these accepted proposals is reflected by the large number of scientific publications originating from the JGI and its users. In 2017, the JGI contributed to 147 peer-reviewed publications, including 17 in the most selective journals, Science and Nature. Seven of our scientists were included in the annual Highly Cited Researchers list by Clarivate Analytics (formerly Thomson Reuters), and several of our scientists serve on editorial boards of leading scientific journals. Selected scientific highlights published in 2017 included:

- **Toward More Effective Carbon Fixation.** By tapping the DNA synthesis expertise of the JGI, a team from the Max Planck Institute (MPI) for Terrestrial Microbiology has reverse engineered a biosynthetic pathway for more effective carbon fixation. This novel pathway is based on a new CO$_2$-fixing enzyme that is nearly 20 times faster than the most prevalent enzyme in nature responsible for capturing CO$_2$ in plants by using sunlight as energy. Successfully reconstituting a synthetic enzymatic network in a test tube for the conversion of CO$_2$ into organic products could lead to potential future applications, such as artificial photosynthesis (Science, November 18, 2016).

- **Antarctic Adaptations in Diatoms.** To learn more about how the Antarctic alga *Fragilariopsis cylindrus* adapted to its extremely cold environment, a team led by University of East Anglia (UEA) scientists conducted a comparative genomic analysis involving three diatoms by tapping the JGI's sequencing and annotation expertise. The paper was recently named a Fast-Breaking Paper for Biology and Biochemistry by Clarivate Analytics (Nature, January 16, 2017).

- **Aspergillus Diversity for Industrial Applications.** An international team including JGI researchers reported sequencing the genomes of 10 novel *Aspergillus* species, more than doubling the number of *Aspergillus* species sequenced to date. Comparing the newly sequenced genomes with the eight other sequenced *Aspergillus* species provided the first ever genus-wide view. The consortium found that *Aspergillus* has a greater genomic and functional diversity than previously understood, broadening the range of potential applications for the fungi considered one of the most important workhorses in biotechnology (Genome Biology, February 14, 2017).

- **Major Gene Expression Regulator in Fungi.** Just four letters—A, C, T, and G—make up an organism’s genetic code. Changing a single letter, or base, can lead to changes in protein structures and functions, affecting an organism’s traits. In addition, though, subtler changes can and do happen, involving modifications of the DNA bases themselves. A JGI-led team reported the prevalence of 6-methyladenine (6mA) modifications in the earliest branches of the fungal kingdom, and proposed the role of 6mA in gene regulation (Nature Genetics, May 8, 2017).

- **Fungal Enzyme Complexes to Break Down Cellulose.** A team led by researchers at the University of California (UC), Santa Barbara, has found for the first time that early lineages of fungi can form complexes of enzymes called “cellulosomes” capable of degrading plant biomass. Consolidating these enzymes, in effect into protein assembly lines, makes them team up to work more efficiently than they would as individuals. Through the FICUS JGI-EMSL program, the team found cellulosomes in anaerobic gut fungi that attack plant biomass as a cluster of enzymes (Nature Microbiology, May 26, 2017).
**IMG/M Database Helps Fill Previously Unknown Protein Structures.** The Pfam database contains close to 15,000 protein families: groups of proteins that share an evolutionary origin. A team led by the University of Washington’s David Baker reported that structural models have been generated for 614 (12%) of the protein families that had previously had no structural information available. This accomplishment was made possible through a collaboration in which the Baker lab’s protein structure prediction server, Rosetta, analyzed the publicly available metagenomic sequences on the JGI’s Integrated Microbial Genomes (IMG) system (Science, January 20, 2017).

**1003 Microbial Genomes Released.** Culminating nearly a decade’s worth of work, JGI’s Prokaryotic Super Program head Nikos Kyrpides and his team reported the release of 1,003 phylogenetically diverse bacterial and archaeal reference genomes — the single largest release to date. With the release of this high-quality genomic information, the JGI provides a wealth of new sequences that will be invaluable to scientists interested in experiments such as characterizing biotechnologically relevant secondary metabolites or studying enzymes that work under specific conditions (Nature Biotechnology, June 12, 2017).

**Novel Group of Giant Viruses Discovered.** Giant viruses were first discovered in 2003, and a handful of other giant virus groups have been found since. JGI scientists’ report of a novel group of giant viruses — dubbed “Klosneuviruses” — with a more complete set of translation machinery genes than any other virus known to date resolved the debate over viruses as a possible fourth domain of life by demonstrating that eukaryote translation-encoding genes had been taken up and incorporated into the viral genomes (Science, April 7, 2017).

**Minimum Metadata Standards.** As genomic data production has ramped up over the past two decades and is being generated on various platforms around the world, scientists have worked together to establish definitions for terms such as “draft assembly” and data collection standards that apply across the board. An international team led by JGI researchers has developed standards for the minimum metadata to be supplied with single amplified genomes and metagenome-assembled genomes submitted to public databases (Nature Biotechnology, August 8, 2017).

To continue enabling these types of consequential scientific discoveries into the future, we invest in an expert workforce and develop genomic technologies that expand frontiers. Over the last year, we have brought into production higher throughput and capacity short-read instrumentation and more robust single-molecule real time long read technology, and continue to develop nanopore-based sequencing technologies. For functional genomics, we have established new capabilities in experimental transcription factor binding motif detection (DAP-Seq), chromatin...
conformation analysis (ChIP-Seq), and virus-based sequencing. In single-cell genomics, we have expanded into eukaryotic de novo genome assembly and the binning of cells containing similar transcriptomes. We continue to provide users with capabilities to determine gene and pathway function through expansion of our DNA synthesis science and metabolomics platforms. These capabilities allow researchers to delve deeper into genomes and maintain the JGI’s relevance through the continuous addition of novel, pioneering approaches available to users.

The JGI Science Programs also develop novel approaches and resources relevant to our mission, showcasing these through high-visibility publications such as the ones cited earlier. These initial demonstration studies serve as an inspiration to our user base to adopt new JGI capabilities and have proven invaluable in developing the JGI user communities. For example, we continue our leadership role in microbial diversity research by expanding the tree of life through exploiting differential coverage binning of deeply sequenced metagenomes to identify new phyla. Additionally, the Plant Program, led by Jeremy Schmutz and largely executed at our partner institute, HudsonAlpha Institute for Biotechnology in Huntsville, Alabama, continues to be one of the world’s leading plant genomics programs. Through this long-standing partnership, the JGI generates extensively annotated de novo and improved genomes of JGI Flagship plants and comparator species, performs large-scale population genetics and gene expression studies to link genes to bioenergy-relevant traits, and characterizes evolutionary genome dynamics and their influence on functional adaptations through pan-genome analyses.

Today’s ability to generate data at a scale and pace that outweighs the capacity to manage, analyze, and interpret data requires scalable computational infrastructure and tools. NERSC and the JGI have developed a deeper partnership that has led to the successful implementation of several computing initiatives, including migration of pipelines to Docker for increased portability, consolidation of workflow management systems, and extensive training on emerging computing technologies. The DOE Office of Science has a major initiative, the Exascale Computing Project (ECP), dedicated to developing an exascale ecosystem to advance scientific discovery. The JGI partners with the Berkeley Lab Computational Research Division and participates in ECP through the ExaBiome project. The main aim of this work is to adapt key elements of the JGI computing workload to run at exascale (a billion billion calculations per second), so the JGI analysis can advance in step with the enormous volumes of data that will be generated. In 2017, we also completed the automated GenBank submission system’s development to release our data to NCBI.

With the wealth and breadth of data being generated, the JGI continues to develop the tools and systems for users to extract new insights. For example, the number of registered users in our IMG system doubled (surpassed 18,000 registrations from 102 countries), as has the number of metagenome datasets integrated in IMG. We have now launched an institutional effort to overhaul IMG (with the working title IMG-Next), which is making good progress towards providing users with improved pipelines and interfaces, and providing scalable database architecture to accommodate and enable the analysis of rapidly growing amounts of microbial genome and microbiome data.

The Emerging Technologies Opportunity Program (ETOP) was designed to facilitate JGI user access to the latest sequence-to-function capabilities developed by external groups collaborating with the JGI. Excellent progress has been made in our ongoing projects, some examples of which are detailed in the “Science: Year in Review” section. This year, we also launched a new ETOP call for proposals related to rapid-prototyping systems to rapidly and at scale determine gene/pathway function.

As a DOE User Facility, the JGI must engage actively with users. Our 12th Annual Genomics of Energy and Environment Meeting filled our Walnut Creek meeting space to capacity. As a result, the meeting will be held in San Francisco in 2018. We had an exciting lineup of speakers, with keynote addresses from Tobias Erb of the Max Planck Institute (MPI) for Terrestrial Ecology and Titus Brown of the University of California, Davis (see Appendix E.) The JGI also hosted the NeLLi (New Lineages of Life) and Microbial and Plant Systems Modulated by Secondary Metabolites meetings, and the first National Microbiome Data Collaborative workshop. The Microbial Genomics and Metagenomics (MGM) workshops marked their tenth year in 2017, and along with that, the milestone of training more than 1,000 scientists from 55 countries in using the IMG resources.
We are also expanding our communication and outreach efforts to attract new users from both the academic and private sectors, and in 2017 formed a Business Development Team to lead a new Industry Engagement Program. This program has already had discussions with more than 50 companies and launched its first sponsored research project with an industrial partner at JGI.

The foundations of building and sustaining high-performing teams at the JGI are extensive training programs (especially for safety), the Performance Management Process, and people development opportunities. The JGI has an exceptionally highly skilled workforce motivated to deliver science and expertise to our users. I have established a new JGI Leadership Team (JLT) comprised of the JGI Director; Deputies for User Programs, Science Programs, Genomic Technologies, and Operations; Chief Informatics Officer, Senior Communications and Outreach Manager, and Business Development and Program Management Office Leader. The JLT has continued to emphasize the safety and wellness of our staff as an institutional priority, and our outstanding safety record has had a positive impact on our overall operational efficiency and productivity.

A core JGI value is diversity and inclusion (D&I), and our culture of inclusion helps us attract, retain, and develop outstanding talent, thereby directly promoting our scientific excellence. Our D&I Working Group is very active in promoting monthly social events, arranging talks and videos to stimulate discussion, and organizing trainings (e.g., to raise awareness of implicit bias). Additionally, the JGI is committed to ensuring that its management teams, advisory bodies, and speakers at JGI meetings and review panels represent the full demographic and scientific diversity of the JGI and its user community. We launched the OneJGI initiative in 2017 to enhance collaboration, trust, and respect at the JGI by revisiting and communicating the JGI’s core values and norms and ensuring open and transparent communication.

Finally, the first step in realizing Berkeley Lab’s vision to co-locate Biosciences activities and organizations at the main Berkeley Lab site is the JGI’s new home, the Integrative Genomics Building (IGB). A groundbreaking ceremony was held at the end of January 2017 to kick off the construction phase of this project. Excellent progress has been made on the construction, with foundations completed and the first-floor structure erected. We aim to move into this building in the first half of 2019.

We are very excited about this move, and the expected scientific and operational productivity gains for the JGI and the DOE Systems Biology Knowledgebase (KBase), as well as related research programs in the Biosciences Area. Achieving systems biology’s promise to understand and solve basic energy and environment problems requires the ability to iteratively predict the behavior of organisms and biological systems, then test those predictions and adjust models accordingly. To fulfill this promise, we have strengthened our partnership with KBase, with the aim of enabling seamless interactions to provide our users with access to KBase tools and data that go beyond what is offered through JGI resources.

In summary, 2017 was another highly productive year for the JGI, and we continue to offer our global user community a unique suite of state-of-the-art capabilities essential to enable the advancement of energy and environmental science and address scientific grand challenges. As the JGI continues its evolution into an Integrative Genome Science User Facility that is the leading center worldwide for energy and environmental genomics, our science, our capabilities, our users, and our OneJGI culture all position us for growing JGI’s future success. This will be the guiding principle as we conduct a refresh of our strategic plan in 2018 that will guide the JGI to meet the future scientific needs of its broad and diverse users.

Nigel J. Mouncey, DPhil
Director, DOE Joint Genome Institute
Achieving the DOE Mission
The Joint Genome Institute (JGI) is a national user facility funded by the DOE’s Office of Biological and Environmental Research (BER) that conducts high-throughput DNA sequencing, synthesis, and analysis aligned with BER’s bioenergy and environmental missions. These missions mirror DOE and national priorities to:

- Develop renewable and sustainable sources of plant biomass feedstocks for biofuels by exploiting genomic knowledge of plants, microbes, fungi, and microbial communities.
- Gain insights into biogeochemical processes controlling the cycling of carbon, nitrogen, and key nutrients in environments and the mobility of heavy metals and radionuclides at contaminated sites for which DOE has stewardship responsibilities.

**Bioenergy**

The United States is the world’s largest consumer of petroleum, with most used for transportation and industry. This drives the DOE’s focus on developing clean, renewable, and sustainable alternative fuel sources from lignocellulosic biomass. Such fuels would ideally offer energy content on par with gasoline while being compatible with the existing fuel distribution infrastructure. Sequencing projects at the JGI that contribute to meeting this goal focus on at least one of three categories: terrestrial plant feedstocks for biofuel production and their associated microbial communities (microbiomes); fungi, microbes, and microbial communities that break down the lignin and cellulose in plant walls; and organisms that convert lignocellulose-derived sugars or lignols into biofuels or other bioproducts currently produced from petroleum, such as plastics.

**Biogeochemistry**

Many DOE-relevant environmental processes are controlled by complex, interconnected biogeochemical reactions. The JGI engages in projects that can couple a genome-enabled understanding of biological processes in the context of the physical, chemical, and geochemical processes controlling the cycling and fate of key elements in environments affecting BER’s energy and environmental missions. Microbes and microbial communities of interest to the JGI as targets for sequencing include those involved in terrestrial carbon, nitrogen, phosphorus, sulfur, and other macronutrient cycles that affect sustainable bioenergy crop growth or global carbon cycling. Others include those involved in the iron, sulfur, and manganese cycles that mediate the transformation of DOE-relevant contaminants, such as heavy metals or radionuclides in soils, freshwater aquatic sediments, and the subsurface. As microbes constitute the largest component of Earth’s biodiversity and biomass, understanding how they metabolize these elements and how environmental changes affect these processes is crucial.

With the support of BER through the Subsurface Systems Scientific Focus Area (SFA) 2.0, Berkeley Lab’s Earth and Environmental Sciences Area (EESA) and their JGI collaborators have conducted major investigations at the field site in Rifle, Colorado to facilitate integrated, field-based subsurface biogeochemical and microbial genomics research relevant to uranium mobility to improve the predictive understanding of subsurface flow and transport relevant to metal and radionuclide contaminants. *(Roy Kaltschmidt, Berkeley Lab)*
Organizational Structure
Senior Management Team

Nigel Mouncey  
JGI Director

Kjiersten Fagnan  
Chief Informatics Officer

Tootie Tatum  
Business Development & Project Management Office Lead

Susannah Tringe  
Deputy, User Programs

Len Pennacchio  
Deputy, Genomic Technologies

Axel Visel  
Deputy, Science Programs

Ray Turner  
Deputy, Operations

David Gilbert  
Senior Manager, Communications & Outreach

Ronan O’Malley  
Sequencing Technologies

Yasuo Yoshikuni  
DNA Synthesis Science Program

Trent Northen  
Metabolomics Group

Atif Shahab  
Institutional Informatics

Tanjia Woyke  
Microbial Program

Daniel Rokhsar  
Eukaryote Super Program

Igor Grigoriev  
Fungal Program

Jeremy Schmutz  
Plant Program

Tanja Woyke  
Microbial Program

Emiley Eloe-Fadrosh  
Metagenome Program

Nikos Kyrpides  
Prokaryote Super Program
Impact 2017
Primary Users **Fiscal Year 2017**

This category captures the primary users of the JGI, which include PIs and their collaborators on all user projects that were active during FY 2017. Each user is uniquely identifiable and is counted once per year regardless of the number of active projects in which he/she may be involved. This count does not include collaborators who are employed by the JGI or funded through the JGI’s partner subcontracts.

**Spending Profile FY2017**

- **41%** Science Programs & Analysis
- **30%** Genomic Technologies
- **9%** Institutional Informatics
- **6%** National Energy Research Scientific Computing Center (NERSC)/IT
- **3%** Lease
- **3%** Project Management Office
- **3%** Management
- **3%** Operations
- **2%** Emerging Technologies Opportunity Program (ETOP)
## Users on the Map: 1,598

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Users of JGI Tools & Data

The JGI produces high-quality data that are made available to the community through our data portals. These investigators are not included in the primary user count because their projects were not conducted as part of JGI’s user programs. When users log in to the systems and download data, this activity is tracked in order to help us understand which data sets are of greatest value to our Users. Additionally, the JGI’s data management system is able to restore data from HPSS upon user request and this activity is logged. In 2017, JGI users downloaded more than 1.8 million data sets and 1 PB of data that were restored through the JAMO system.

Workshops and Meetings

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<td>MSA JGI Fungal Program/MycoCosm Workshop</td>
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<td>American Plant Biology talk and joint outreach booth with Kbase</td>
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Sequence Output

(in billions of bases or GB)

Massively Parallel Short Read

The JGI supports short and long read sequencers, where a read refers to a sequence of DNA bases. Short read sequencers produce billions of 300 base reads used for quantification such as in gene expression analysis. Long read sequencers currently average 12,000 base reads and are used for de novo genome assembly.

Single Molecular Long Read Sequencing

Combined short read and long read totals per year give JGI’s annual sequence output.
Papers published 2013-2015 were cited by researchers over 5,200 times in 2016.
Case Study: **10 Years of MycoCosm: Approaching 1,000 Fungal Genomes**

MycoCosm is the JGI’s web-based repository of fungal resources. With nearly 1,000 genomes, it is the world’s largest interactive collection of fungal genomes. MycoCosm started with the integration of six genome projects for six plant pathogenic Dothideomycetes; the JGI held jamborees in 2007–2008 that brought together the six labs responsible for the individual genome projects. This historic exercise in community building and data integration led to the first large-scale fungal comparative genomics project at JGI, and a series of joint publications.

MycoCosm became an official JGI community resource in 2010 and, due to near exponential growth, is approaching an important milestone of 1,000 fungal genomes. Through its community building framework, there were 8,500 registered users and 130,000 distinct web visitors in 2017 alone.

JGI has transformed mycology into genome-based science. MycoCosm enabled the transition from a single-genome analysis — sequencing the first mycorrhiza, the first wood decayer, and the first of many other targeted DOE mission-relevant groups — to comparative genomics.

**Plant-Mycorrhizal Associations.** Over 80% of plant species depend on mycorrhizae. The first genomes of each major type of mycorrhizae — arbuscular (*Rhizophagus irregularis*), ericoid (*Oidiodendron maius*), orchid (*Tulasnella calospora*, learn more in the “Science: Year in Review” section), and ectomycorrhizae (*Laccaria bicolor*) — were sequenced and annotated by the JGI. The sequencing and analysis of a larger group of mycorrhizal fungi identified similar but independently evolved symbiotic toolkits, demonstrating their convergent evolution in mycorrhizal fungi across the entire Fungal Tree of Life. The JGI supports studies of fungal interactions with plants in several model plant-fungal systems; for example, the genome of the dominant ectomycorrhizal *Cenococcum geophilum* suggested clues to plant drought tolerance.

**Lignocellulose Degradation by Filamentous Fungi.** Lignin is the second-most important component of plant biomass after cellulose. Together the two biopolymers support plant growth and protect plants from microbial attack. The first genomes of a white rot fungus (*Phanerochaete chrysosporium*) and a brown rot fungus (*Postia placenta*) were published in 2004 and 2009, respectively, and demonstrated dramatic differences in the fungi’s approaches to lignocellulose degradation. In 2012, the JGI published a comparative genomics analysis of 12 wood decay fungi and traced the evolution of class II peroxidases, lignin-degrading enzymes that distinguish white rot from brown rot fungi. Further analysis suggested a continuum rather than a dichotomy between the white-rot and brown-rot modes of wood decay, and identified several gene families predictive of wood decay mode.
Sugar-Fermenting Yeasts. Sugar fermentation is a critical step in converting sugars extracted from plant biomass into biofuels and bio-based products. Plant biomass is composed of a variety of sugars, but the model yeast *Saccharomyces cerevisiae* grows primarily on glucose. The genome of the xylose-fermenting yeast *Scheffersomyces stipites*, sequenced by the JGI in 2007, revealed genes later used in engineering xylose-fermenting *S. cerevisiae* strains. In 2016, the JGI sequenced and analyzed diverse Ascomycete yeasts growing on other types of sugars to produce an expanded catalog of genes, enzymes, and pathways. This yeast project also discovered a genetic code change in *Pachysolen tannophilus* where the codon CUG encodes the amino acid alanine instead of the expected leucine or serine seen in nearly all other eukaryotic nuclear genomes.

Genomics of Consolidated Bioprocessing. Most state-of-the-art platforms for the bioconversion of plant biomass into biofuels involve multiple costly steps of biomass pretreatment and saccharification, challenging the economic feasibility of these technologies. An alternative strategy, consolidated bioprocessing (CBP), aims to combine all these steps into a single unit operation. In collaboration with the Joint BioEnergy Institute (JBEI), the JGI sequenced and analyzed four endophytes (*Hypoxylon* and *Daldinia* spp.) to assess their potential to produce “mycodiesel” (volatile organic compounds with properties similar to fossil fuels) while growing on lignocellulosic plant and agricultural residues. Their ability to simultaneously deconstruct lignocellulose and convert it into mycodiesel makes them candidate CBP hosts for biofuel production.

More recent Community Science Program (CSP) projects have significantly advanced MycoCosm’s development. The production of fungal multi-omics datasets for functional genomics is pioneered by fungal miniENCODE (Encyclopedia of DNA Elements) projects. The ongoing JGI 1000 Fungal Genomes project, aimed at producing at least one reference genome for every known family of fungi, is one of the world’s largest genomics studies of fungal diversity. These and similar projects not only ensure further growth of MycoCosm, but also transformed the portal from a static data repository resource into a dynamic platform for the user community to analyze sequenced genomes and nominate new species for sequencing to fill gaps in the Fungal Tree of Life.

Growth in number of genomes in MycoCosm

![Graph showing the growth in number of genomes in MycoCosm from 2008 to 2017.](image)
Among the JGI’s major initiatives for the Plant Program is focusing computational and experimental efforts to move beyond sequence to function, ultimately depositing validated accurate data sets into KBase. To that end, the Plant Functional Genomics team uses the model grass *Brachypodium distachyon* and other plants to develop and utilize resources to understand traits such as genome organization and regulation, abiotic stress tolerance and the molecular basis of perenniality.

Science:

A Year in Review
The ability to design and manufacture synthetic DNA has opened tremendous possibilities in genomic research. In addition to providing access to samples that are hard to find in nature (as well as crafting genomic sequences not known to occur in the natural world), manufacturing DNA enables scientists to test any sequence in a wide variety of contexts and environments.

BioCAD/CAM software tools help researchers design sequences that can be critical to discovering new solutions for energy and the environment, but so far the software has not been able to automatically fix problematic sequences. This delays synthesis of the designed DNA and the subsequent transition to the manufacturing process.

To solve this problem, JGI researchers Ernst Oberortner, Jan-Fang-Cheng, Nathan Hillson, and Sam Deutsch developed a suite of build-optimization software tools (BOOST) to automate the synthetic DNA design process—and do away with the trial-and-error process scientists currently utilize to determine a sequence that can be synthesized. As reported December 6, 2016, in ACS Synthetic Biology, BOOST can automatically detect “difficult” sequences (of nucleotides) and redesign them for DNA synthesis, addressing DNA sequences with certain problematic characteristics (e.g., extreme %GC content, sequence patterns, and repeats), which decrease the success rate of DNA synthesis. By improving the design and manufacture of synthetic DNA through enhanced tools, scientists can accelerate gene discovery and gene characterization toward practical applications for energy and the environment. The BOOST suite of tools is available as a web application at https://boost.jgi.doe.gov.
Finding a New Major Gene Expression Regulator in Fungi

Just four letters — A, C, T, and G — make up an organism’s genetic code. Changing a single letter, or base, can lead to changes in protein structures and functions, affecting an organism’s traits. In addition, though, subtler changes can and do happen, involving modifications of the DNA bases themselves. The best-known example of this kind of change is a methylation of the base cytosine at the fifth position on its carbon ring (5mC). In eukaryotes, a less well-known modification involves adding a methyl group to base 6 of adenine (6mA).

As reported May 8, 2017, in Nature Genetics, a team led by JGI scientists described the prevalence of 6mA modifications in the earliest branches of the fungal kingdom. Though fungi have existed for a billion years and collectively are capable of degrading nearly all naturally occurring polymers and even some human-made ones, most of the species that have been studied belong to just two phyla: the Ascomycota and Basidiomycota. The remaining six groups of fungi are classified as “early diverging lineages,” the earliest branches in fungal genealogy. They comprise a little-explored realm of fungi, providing a repertoire of important and valuable gene products for DOE bioenergy and environment missions.

“By and large, early-diverging fungi are very poorly understood compared to other lineages. However, many of these fungi turn out to be important in a variety of ways,” said study first author and JGI analyst Stephen Mondo.

Many of the fungal genomes used in the study were sequenced as part of the JGI’s 1000 Fungal Genomes initiative aimed at producing at least one reference genome for every family of fungi. For the study, the team used 16 fungal genomes sequenced at the JGI using the Pacific Biosciences sequencing platform. JGI scientists took advantage of this sequencing platform to explore epigenetic (5mC, 6mA) modifications. They discovered very high levels of 6mA in fungi, where up to 2.8% of all adenines were methylated, and confirmed these findings using multiple independent methods. The previous record holder for genomic 6mA, noted Mondo, is the alga Chlamydomonas reinhardtii (sequenced and annotated by the JGI), in which just 0.4% of adenines were methylated.
“This is one of the first direct comparisons of 6mA and 5mC in eukaryotes, and the first 6mA study across the fungal kingdom,” said JGI Fungal Program head and senior author Igor Grigoriev. “6mA has been shown to have different functions depending on the organism. For example, in animals it is involved in suppressing transposon activity, while in algae it is positively associated with gene expression. Our analysis has shown that 6mA modifications are associated with expressed genes and are preferentially deposited based on gene function and conservation, revealing 6mA as a marker of expression for important functionally-relevant genes. The discovery of DNA methylation in early diverging fungi helps the research community better understand regulation of genes that encode the parts for bio-based economy and bioenergy applications.”

The team found that, in addition to 6mA performing what seems to be the opposite role of 5mC (which suppresses expression), the presence of 5mC and 6mA are inversely correlated. Mondo described the identification and proposed role of 6mA in early diverging fungi at the 12th Annual Genomics of Energy and Environment Meeting. Watch his talk on the JGI’s YouTube channel at http://bit.ly/JGI2017Mondo.

**New Technology to Access Microbial Dark Matter**

New technologies that can be harnessed to illuminate so-called microbial “dark matter,” the majority of the planet’s microbial diversity that remains uncultivated, have many benefits. The genes and metabolic functions of these yet-undiscovered microbes enable them to live in a wide range of environments, and could have potential applications in fields ranging from bioenergy to biotechnology to environmental research.
More than 50,000 microbial genome sequences are in the JGI’s Integrated Microbial Genomes & Microbiomes (IMG/M) publicly accessible database, and many of them have been uncovered through the use of metagenomic sequencing and single-cell genomics. Despite their utility, these techniques have limitations: single-cell genome amplifications are time-consuming and often incomplete, and shotgun metagenomics sequencing generally works best if the environmental sample is not too complex.

A team of Stanford University researchers reported the development of a microfluidics-based, mini-metagenomics approach to mitigate these challenges on July 5, 2017, in *eLife*. The technique starts with reducing the environmental sample’s complexity by separating it using microfluidics into 96 subsamples of five to 10 cells. Then the genomes in the few cells in each subsample are amplified, and libraries are created for sequencing of these mini-metagenomes. The smaller subsamples can be held to single-cell resolution for statistical analyses. Co-occurrence patterns from many subsamples can also be used to perform sequence independent genome binning. The technology was developed through resources provided by the JGI’s Emerging Technologies Opportunity Program (ETOP), which was launched in 2013.

The team validated the technique using a synthetic microbial community, and then applied it to samples from the Bijah and Mound hot springs at Yellowstone National Park. Among the team’s findings was that the microbes at Mound Spring had higher potential to produce methane than the microbes from Bijah Spring. The team also identified a microbial genome from Bijah Spring that could reduce nitrite to nitrogen. Applying this new technology to additional sample sites will add to the range of hitherto uncharacterized microbial capabilities with potential DOE mission applicability.

**IMG/M Database Helps Fill Previously Unknown Protein Structures**

Proteins largely form a cell’s structures and carry out its functions. They control growth and influence mobility, serve as catalysts, and transport or store other molecules. On paper, the one-dimensional amino acid sequence may seem meaningless, but when viewed in three dimensions, a protein’s structure, and particularly the way it folds, determines its functions.

The Pfam database contains close to 15,000 protein families: groups of proteins that share an evolutionary origin. For nearly a third of these protein families, at least one protein in each family already has an experimentally determined structure. For another third of the protein families, comparative models could be built with some degree of confidence. For the final third of the protein families in the database, however, no structural information exists.

In the January 20, 2017, issue of *Science*, a team led by the University of Washington’s David Baker collaborated with JGI researchers to report that structural models have been generated for 614, or 12%, of the protein families that had previously had no structural information available. This accomplishment was made possible through a collaboration in which the Baker lab’s protein structure prediction server, Rosetta, analyzed the metagenomic sequences publicly available on the JGI’s IMG/M system. “That this could be accomplished using computational modeling methods was not at all apparent five years ago,” the team noted in its paper.

Armed with genome sequences, researchers like Baker have been able to identify sets of amino acids that evolve simultaneously, even though they are nowhere near each other on the unfolded chain. JGI Prokaryote Super Program head Nikos Kyrpides said the collaboration between the Baker lab and the JGI enabled the team to come up with a powerful way of predicting structures and structural alignments. “Such efforts were previously restricted on protein families generated from sequences found on the isolate genome only. These genomes comprise about 200 million sequences. As expected, by harnessing the 5 billion assembled metagenome sequences available on our IMG/M database, we were able to dramatically increase the coverage of many of the known protein families. Efforts like this one heavily depend on the availability of assembled metagenomics sequences, which is an advantage the JGI brings to the table with our high-quality assemblies. The application of such tools on our data provides a great example of how the larger community can utilize JGI resources for discovery.”
Provisioning high-quality, publicly accessible sequence data goes hand-in-hand with developing and maintaining the databases and tools that the research community can employ to help answer scientific questions. In the “Database” issue of the journal *Nucleic Acids Research*, released January 1, 2017, JGI researchers reported on the latest updates to several publicly accessible databases and computational tools that benefit the global community of microbial researchers. One report focused on a new database dedicated to global viral diversity called IMG/VR (https://img.jgi.doe.gov/vr/), which followed on the heels of a recent JGI viral diversity study in *Nature*.

Additional articles in the same issue describe updates to several publicly accessible, interactive databases since the last set of reports was published in 2014. For example, as of July 2016, 47,516 archaeal, bacterial, and eukaryotic genomes were in the Integrated Microbial Genomes with Microbiome Samples (IMG/M: https://img.jgi.doe.gov/m/) system, with researchers noting that number “represents an over 300% increase since September 2013.” Another paper concerns the Genomes Online Database (GOLD: https://gold.jgi.doe.gov), a manually curated data management system that catalogs sequencing projects with associated metadata from around the world. A fourth paper focuses on the Integrated Microbial Genomes Atlas of Biosynthetic gene
Clusters (IMG-ABC: https://img.jgi.doe.gov/abc/). Launched in 2015, IMG-ABC allows researchers to search for biosynthetic gene clusters and secondary metabolites. The latest update now incorporates ClusterScout, a tool for targeted identification of custom biosynthetic gene clusters across several thousand isolate microbial genomes, and a new search capability.

Microbes play key roles in maintaining the planet’s biogeochemical cycles. Viruses, thought to outnumber microbes by 10-fold, exert major influences on microbial survival and community interactions. Advances in sequencing technologies have generated vast amounts of data about viruses, requiring tools to manage and interpret the information. These updates focus on database analytical tools for microbial genomics and viruses relevant to DOE bioenergy and environment missions.

Minimum Information Standards for Uncultivated Microbial Genomes

During the Industrial Revolution, the establishment of standards allowed builders to produce supplies in bulk, maintain production quality, and fuel interstate commerce. The importance of standards is dramatically illustrated when they don’t exist or are not commonly accepted.

More than a century after the Industrial Revolution, advances in DNA sequencing technologies have caused similarly dramatic shifts in scientific research. One aspect is studying the planet’s biodiversity. An international team led by JGI researchers has developed standards for the minimum metadata to be supplied with single amplified genomes (SAGs) and metagenome-assembled genomes (MAGs) submitted to public databases. The team’s work was published August 8, 2017, in *Nature Biotechnology*.

“Over the last several years, single-cell genomics has become a popular tool to complement metagenomics,” said study senior author Tanja Woyke, head of the JGI Microbial Program. “Starting in 2007, the first single-cell genomes from environmental cells appeared in public databases, and they are draft assemblies with fluctuations in the data quality. Metagenome-assembled genomes have similar quality challenges. For researchers who want to conduct comparative analyses, it’s really important to know what goes into the analysis. Robust comparative genomics relies on extensive and correct metadata.”

Microbes play crucial roles in regulating global cycles involving carbon, nitrogen, and phosphorus among others, but many of these microorganisms remain uncultured and unknown. Learning more about this “microbial dark matter” involves sequencing metagenomes or the amplified DNA of single cells, then bioinformatically...
identifying genomes. As genomic data production has ramped up over the past two decades and is being generated on various platforms around the world, scientists have worked together to establish definitions for terms such as “draft assembly” and data collection standards that apply across the board.

In their paper, Woyke and her colleagues proposed four categories of genome quality. Low-Quality Drafts would be less than 50% complete, with minimal review of the assembled fragments and less than 10% contaminated with non-target sequence. Medium-Quality Drafts would be at least 50% complete, with minimal review of the assembled fragments and less than 10% contamination. High-Quality Drafts would be more than 90% complete with the presence of the 23S, 16S, and 5S rRNA genes, as well as at least 18 tRNAs, and with less than 5% contamination. The Finished Quality category is reserved for single contiguous sequences without gaps and less than 1 error per 100,000 base pairs.

Moving from a proposal in print to implementation requires community buy-in. Woyke and her colleagues conceived of the minimum metadata requirements for SAGs and MAGs as extensions to existing metadata standards for sequence data, referred to as “MIxS,” developed and implemented by the Genomic Standards Consortium (GSC) in 2011. The GSC is an open-membership working body that ensures the research community and the data repositories are engaged in the standards development process. By working directly with these groups, the GSC can assist both large-scale data submitters and databases to align with the MIxS standard and submit compliant data.

“I think it helps that the people developing standards are the people conducting the studies,” said GSC President and study co-author Lynn Schriml of the Institute of Genome Sciences at the University of Maryland School of Medicine. “We have a vested interest in the data. Developing these novel metadata standards enables researchers to consistently report the most critical metadata for analysis. Capturing data using controlled vocabularies facilitates data consistency, thus making the databases richer and reusable.”

As the number of single-cell genomes and metagenome-assembled genomes in public databases continues to rise, developing minimum metadata standards enables researchers to consistently perform robust comparative analyses. (Zosia Rostomian, Berkeley Lab Creative Services)
Mining Metagenomics Data for New CRISPR-Cas Systems

In microbes, CRISPR-Cas systems grant adaptive immunity, and these gene-editing tools are the bases of versatile technologies revolutionizing research. Thus far, CRISPR-Cas technology has been based only on systems from isolated bacteria. In a study led by longtime JGI collaborator Jill Banfield of UC Berkeley and Berkeley Lab’s Earth & Environmental Sciences Area (EESA) published in *Nature* on December 16, 2016, researchers reported discovering a CRISPR-Cas9 system in archaea, as well as two previously unknown simple CRISPR-Cas systems in uncultivable bacteria. To identify these new systems, Banfield and her colleagues harnessed more than a decade’s worth of metagenomic data from samples sequenced and analyzed by the JGI.

Microbes play key roles in the planet’s cycles, and characterizing them helps researchers work towards solutions for energy and environmental challenges. Examining environmental microbial communities has provided access to an unprecedented diversity of genomes and CRISPR-Cas systems that have many applications, including biological research. By using the combined computational-experimental approach that was successful in this study, nearly all environments where life exists can be investigated. The researchers analyzed 155 million protein coding genes from uncultivated microbial communities, leading to the novel discoveries. The CasX and CasY proteins were found in bacteria from groundwater and sediment samples. The archaeal Cas9 were identified in samples taken from the Iron Mountain Mine Superfund site as part of Banfield’s pioneering metagenomics work with the JGI. Both CasX and CasY are among some of the most compact systems ever identified. Banfield’s team noted that this application of metagenomics gives a significant validation to studies of CRISPR-Cas proteins using living organisms.

The newly discovered CRISPR-CasY system was found in bacteria from deep underground at Utah’s Crystal Geyser.
Novel Group of Giant Viruses Discovered

Viruses are everywhere; their population is estimated to surpass the number of stars in the Milky Way. Giant viruses are characterized by disproportionately large genomes and virions that house the viruses’ genetic material. They can encode several genes potentially involved in protein biosynthesis, a unique feature which has led to diverging hypotheses about the origins of these viruses.

As reported April 7, 2017, in *Science* by a JGI-led team that included collaborators at the National Institutes of Health, the University of Vienna, and CalTech, the discovery of a novel group of giant viruses (dubbed “Klosneuviruses”) with a more complete set of translation machinery genes than any other virus known to date lends insights into viral evolution.

The predicted hosts for the Klosneuviruses are protists: single-celled microorganisms. And while their direct impacts on protists are not yet worked out, these giant viruses are thought to have a significant impact on these protists that help regulate the planet’s biogeochemical cycles.

“The discovery presents virus evolution for us in new ways, vastly expanding our understanding of how many essential host genes viruses can capture during their evolution,” said National Institutes of Health evolutionary and computational biologist Eugene Koonin, a study co-author whose lab collaborated with the JGI on analyzing the Klosneuvirus genome. “Since protein synthesis is one of the most prominent hallmarks of cellular life, it shows that these new viruses are more ‘cell-like’ than any virus anyone has ever seen before.”

Scientists have been fascinated by giant viruses since 2003, when a group of French biologists discovered the Mimiviruses. Since then, a handful of other giant virus groups have been found. Giant viruses have the unique ability to encode proteins involved in translation (typically DNA to RNA to protein), piquing researchers’ interests as to the giant viruses’ origin. Two evolutionary hypotheses have emerged; one posits that giant viruses evolved from an ancient cell, perhaps one from an extinct fourth domain of cellular life. Another — a scenario championed by Koonin — presents the idea that giant viruses arose from smaller viruses.

The discovery of Klosneuvirus supports the latter idea, according to Tanja Woyke, JGI Microbial Program lead and senior author of the paper. “In this scenario, a smaller virus infected different eukaryote hosts and picked up genes encoding translational machinery components from independent sources over long periods of time through piecemeal acquisition,” she said.

Sorghum is of interest to the DOE as a bioenergy crop because of its drought resistance and ability to thrive on marginal lands. JGI sequenced the sorghum reference genome (*Sorghum bicolor*), publishing it in 2009. The Epigenetic Control of Drought Response in Sorghum (EPICON), a five-year project funded by the DOE and involving JGI researchers, focuses on understanding how sorghum, a grass related to corn, thrives on limited resources, including water.
A Gene that Influences Grain Yields in Grasses

Gene discovery in maize, an important food and bioenergy crop, has been limited due to its large and complex genome. *Setaria viridis* has been proposed as a model crop for maize; *Setaria* species, among them green foxtail (*S. viridis*) and foxtail millet (*S. italica*), are related to several candidate bioenergy grasses including switchgrass and Miscanthus. They serve as model systems to study grasses that photosynthetically fix carbon from CO\(_2\) through a water-conserving (C\(_4\)) pathway. The genomes of both green foxtail and foxtail millet were sequenced and annotated through the JGI’s CSP.

A team led by the Donald Danforth Plant Science Center’s Tom Brutnell and including JGI researchers reported that it had identified genes that may play a role in flower development on the panicle of green foxtail. The panicle is the spear-shaped flowering cluster at the tip of each branch necessary for reproduction. The team’s work was published April 18, 2017, in *Nature Plants*.

The team identified four sparse panicle recessive mutants, tagged *spp1* through *spp4*, that lead to panicles with reduced and uneven flower clusters. Focusing on the *spp1* mutation, they performed deep sequencing to specifically locate the genes that cause the mutation, narrowing their search down to a 1-million base sequence. They ultimately identified the *SvAUX1* gene in green foxtail as one critical for flower cluster development in green foxtail. Panicle development is critical for determining grain yield that is crucial to food crops as well as candidate crops for producing renewable and sustainable fuels. A homologous gene in maize was identified as playing a similar role, illustrating the value of model systems in finding genes involved in important properties in potential bioenergy-relevant plants.
Mutualism and Lipid Metabolism in Fungi

To answer the challenge of producing renewable, sustainable alternative fuels, researchers aren’t just looking at developing candidate bioenergy crops but are also reviewing other natural sources of energy-dense oils, such as fungi. To learn more about how bacteria interact with fungi in a symbiotic relationship to support the biochemical processes that could contribute to the development of alternate fuel sources, Cornell and JGI researchers used a model bacterial-fungal system to reveal the mechanism for lipid production in oil-producing, or oleaginous, fungi.

The widespread use of antimicrobials has led to the perception that fungi and bacteria are antagonists, but this is not always true. Researchers at Cornell University and the JGI reported on the mechanisms influencing the mutualistic relationship between the fungal plant pathogen *Rhizopus microsporus* and a *Burkholderia* endosymbiont in the *Proceedings of the National Academy of Sciences* on December 12, 2016. The work highlighted the complicated nature of fungal-bacterial relationships that can oscillate between mutualistic and antagonistic.

*Rhizopus* is a plant pathogen of crops including rice, sunflower, and maize, and it relies on the toxin produced by the endosymbiont bacteria. This symbiotic relationship was discovered in 2005, and researchers expect more examples of fungal-bacterial mutualisms to be found now that they are being actively sought out. The Cornell team cultivated the fungi and isolated the endobacteria while the JGI team sequenced, assembled, and annotated the host and non-host genomes as part of the JGI’s 1000 Fungal Genomes project and other JGI Community Science Projects. The results indicate that the transition between symbiosis and antagonism between the bacteria and their fungal host is driven by a novel mechanism involving fungal lipid metabolism. “When lipid metabolism in the fungus is triggered and phosphatidic acid (PA)-producing enzymes are activated, the fungus and bacteria act as mutualists. When the PA-producing enzymes are inhibited, the bacteria and fungi act as antagonists,” wrote Cornell’s Teresa Pawlowska and her graduate student Olga Lastovetsky.
A Genome-Wide Map of Bacterial Genes

A plant’s health and development are influenced by microbes in the soil and in the rhizosphere where the plant roots interact with the soil, and the endophytes residing within the plant. By identifying the bacterial genes that alter how well microbes colonize a plant, researchers can develop targeted approaches to improve plant health and growth for a number of applications, including increased biomass yield for biofuel production. To better understand how microbes colonize the root environment, researchers from JGI and a team led by Jeff Dangl of the Howard Hughes Medical Institute at the University of North Carolina at Chapel Hill applied a genome-wide transposon mutagenesis approach on the model plant growth-promoting bacterium *Pseudomonas simiae*. The team used the model plant *Arabidopsis thaliana* as a host to generate a genome-wide map of bacterial genes that affect the efficacy of microbial colonization.

One of the principal challenges arising from rapid sequencing is the assignment of functions to new genes. The randomly barcoded transposon sequencing (RB TnSeq) approach used here can accelerate the association of new genes with characteristics and behaviors of importance to DOE missions, such as understanding how microbes help (or hinder) the growth of crops that could serve as bioenergy feedstocks. As reported September 22, 2017, in *Plos Biology*, they used RB Tn-Seq to identify 115 genes that, when mutated, cause reduced root colonization capabilities. These genes are involved in functions such as sugar metabolism, cell wall synthesis, and motility. The team also identified 243 genes that, when mutated, positively alter root colonization capabilities, many of which are likely involved in amino acid transport and metabolism. From these nearly 360 genes, the team identified a subset of 43 genes to which very little or no functional information could be assigned. The researchers suggested that these genes may represent novel functions or pathways yet to be characterized. The work shows that RB-TnSeq can be applied to assess *in vivo* bacterial plant root colonization.

Among the contributors to this project was Sabah Ul-Hasan, a 2015 intern through the JGI/University of California (UC), Merced Genomics Distinguished Graduate Internship Program. The program offers UC Merced graduate students hands-on experience in cutting-edge genome research as part of the JGI’s commitment to training the next generation of scientific talent.
Mutant Rice Database for Bioenergy Research

For more than half of the world’s population, rice is the primary staple crop. As a grass, it is a close relative of the candidate bioenergy feedstock switchgrass. Boosting yields of bioenergy feedstock crops such as grasses requires a better understanding of how enzymes and proteins synthesize plant cell walls to modify the processes and the composition. Until now, mutant collections for grass models have lagged behind those available for the Arabidopsis model system. In fast-neutron irradiation, exposure to high-energy neutrons induces a wide variety of mutations by making changes in DNA. Using this approach, rice researchers created the first major, large-scale collection of mutations for grass models.

As reported June 2017 in The Plant Cell, a team led by Pam Ronald of UC Davis, and including researchers at the JGI and the Joint BioEnergy Institute (JBEI), a DOE Bioenergy Research Center, have assembled the first major large-scale collection of mutations for grass models. Their goal is to have a functional genomics resource for grass models involved in plant cell wall biosynthesis studies.

The team used the model rice cultivar Kitaake (Oryza sativa L. ssp. japonica), and compared the genes against the reference rice genome of another japonica subspecies called Nipponbare available on the JGI Plant Portal Phytozome. Through fast-neutron irradiation, the time-consuming procedures involving plant transformation or tissue culture were bypassed, allowing for faster development of rice mutant collections. Resequencing the 1,504 mutants has allowed researchers to identify structural variants and mutations, providing an invaluable resource for grass models being used to improve candidate bioenergy feedstock crops such as switchgrass.

Information on this new, large-scale collection of more than 90,000 mutations affecting nearly 60% of all rice genes is available on the publicly accessible database KitBase.

Fungal Enzyme Clusters Better at Attacking Biomass

One of the biggest barriers to commercial production of sustainable biofuels is cost-effectively breaking down the bioenergy crops into sugars that can then be converted into fuel. To reduce this barrier, bioenergy researchers are looking to nature and the estimated 1.5 million species of fungi that, collectively, can break down almost any substance on earth, including plant biomass.

As reported May 26, 2017, in Nature Microbiology, a team led by researchers at UC Santa Barbara, found for the first time that early lineages of fungi can form complexes of enzymes capable of degrading plant biomass. Consolidating these enzymes, in effect into protein assembly lines, makes them team up to work more efficiently than they would as individuals.
“There are protein complexes in bacteria called cellulosomes that pack together the enzymes to break down plant biomass,” said study senior author Michelle O’Malley of UC Santa Barbara. “The idea is that these clusters are better at attacking biomass because they are keeping the different enzymes in place with plugs called dockerins, so they work more efficiently. This has been detailed in bacteria for more than 20 years, but now seen for the first time in fungi.”

The work was enabled by harnessing the capabilities of two DOE Office of Science User Facilities: the JGI at Berkeley Lab, and the Environmental Molecular Sciences Laboratory (EMSL) at Pacific Northwest National Laboratory (PNNL). With help from both user facilities through the Facilities Integrating Collaborations for User Science (FICUS) initiative, the team has now found protein complexes in anaerobic gut fungi that O’Malley said in principle do the same thing: attack plant biomass as a cluster of enzymes. While the team found that many of the enzymes in these complexes resulted from horizontal gene transfers with gut bacteria, it also noted differences in the composition compared with the bacterial cellulosomes. For one thing, dockerins and scaffoldin are not similar between fungi and bacteria. Also, the bacterial cellulosomes are species-specific.

The study involved a comparative genomics analyses of five fungi that belong to the Neocallimastigomycetes, a clade of the early-diverging lineages that are not well-studied. Three of the fungi were isolated from animal gut samples collected by the UC Santa Barbara team and sequenced and annotated by the JGI. Study co-senior author and JGI Fungal Program head Igor Grigoriev noted that a multi-omics approach that harnessed the genomics and molecular characterization capabilities through the FICUS collaborative science initiative was critical for the research.

“It’s the first time we’ve seen parts of the fungal cellulosome,” Grigoriev said. “Through the JGI-EMSL FICUS initiative, proteomics allowed us to find the first of these really large ~700 kiloDaltons (kDa) fungal proteins that hold all enzymes together, compared to the molecular weight of 34 kDa of an average protein. Then the high quality of genome assemblies enabled identification of multiple copies of this protein in each of the gut fungi genomes. Just having proteomics or sequencing tools isn’t enough, since these proteins are not similar to anything else outside of Neocallimastigomycetes. Though the fungal cellulosome was discovered through proteomics, we needed genomics and transcriptomics to decode all its parts.”

The work is an extension of O’Malley’s studies of anaerobic gut fungi, which appeared in Science. “It’s a lot of the same players, but we’re digging deeper now because we have high-resolution genomes, and we didn’t have them then,” she said. “We’re able to conduct more comparative genetics and now we’re trying to figure out the ecological roles in their microbiome.” Watch her talk on this topic from the 2016 Genomics of Energy and Environment Meeting at bit.ly/JGI2016OMalley.
A Genus-Wide View of Aspergillus Diversity

Aspergillus niger has been used for decades to produce citric acid — a compound frequently added to foods and pharmaceuticals — through fermentation at an industrial scale. Other Aspergillus species play critical roles in biofuel production and plant and human health. Since the majority of Aspergillus fungi’s 350 species have yet to be sequenced and analyzed, researchers are still at the tip of the iceberg when it comes to understanding the full spectrum of useful compounds the fungi may generate.

As reported February 14, 2017, in Genome Biology, an international team including JGI researchers sequenced the genomes of 10 novel Aspergillus species, more than doubling the number of Aspergillus species sequenced to date. The newly sequenced genomes were compared with the eight other sequenced Aspergillus species. With this first ever genus-wide view, the international consortium found that Aspergillus has a greater genomic and functional diversity than previously understood, broadening the range of potential applications for the fungi considered one of the most important workhorses in biotechnology.

Comparative growth of Aspergillus spp. (Ad Wiebenga & Ronald de Vries, Westerdijk Fungal Biodiversity Institute, Utrecht, The Netherlands)
Conducted through the JGI’s CSP, the study demonstrates the importance of evaluating biodiversity within a genus to understand how fungi can be harnessed to solve a variety of problems. Sequencing a diverse set of *Aspergillus* genomes allows researchers to build a more comprehensive catalog of enzymes for biotechnological applications, added JGI Fungal Program Head Igor Grigoriev, senior author of the paper. Those applications include harnessing *Aspergillus* to help protect crops and ward off agents that can cause diseases in plants. Comparing the newly sequenced genomes to those already available, researchers found a huge variety of carbohydrate-active enzymes (CAZymes) among the *Aspergillus* species, suggesting distinct strategies to break down plant biomass. CAZymes are responsible for breaking down plant cell walls, which is useful for industrially processing plants the DOE considers candidate bioenergy crops. The sugars that are part of these cell walls can’t be accessed and fermented to make biofuels unless the walls are broken down by agents like CAZymes.

“Several *Aspergillus* species have already established status as cell factories for enzymes and metabolites,” said study lead author Ronald de Vries of the Westerdijk Fungal Biodiversity Institute in the Netherlands. But despite the new insights gained, he emphasized that much still remains unknown about the full spectrum of what *Aspergillus* can do. “There is [still] much to learn and get from a better study [of *Aspergillus*],” he said. “The potential for applications within the genus has barely been touched.”

### Nitrogen Uptake Between Fungi and Orchids

Orchids, like the majority of terrestrial plants, form symbiotic relationships between their plant roots and soil fungi, known as mycorrhizal association. However, unlike other terrestrial plants, orchids rely on their mycorrhizal fungal partners for nutrient supply during the feed germination and development stages. Following this stage, most orchid species develop leaves and are capable of self-nourishment, whereas some species continue to rely on their fungal partners for an organic carbon supply. As reported in the January 2017 issue of *New Phytologist*, a team led by University of Turin researchers investigated the orchid mycorrhizal fungus *Tulasnella calospora* as both a free-living mycelium and in symbiosis with the photosynthetic orchid long-lipped serapias, or *Serapias vomeracea*.

For the first time, researchers looked at the fungal genes that may have been involved in both the uptake and transfer of nitrogen to the host plant. RNA sequencing for the project was performed at the JGI as part of the 2013 CSP portfolio. The team also used the JGI fungal genome database MycoCosm to identify fungal genes coding for proteins that were involved in nitrogen uptake and transfer. They found that the *T. calospora* genome has two genes coding for ammonium transporters and several genes coding for amino acid transporters, proteins that play roles in the nitrogen nutrient pathway.
This study provided a model system amenable to experimental manipulation for plant-fungi nutrient exchanges on a symbiotic level and offers insights into how host plants benefit from the mutualistic relationships formed with soil fungi, which can expand their habitat range. Understanding these vital relationships may shed light on microbial symbioses applicable to growing bioenergy feedstock plants. Overall, the orchid mycorrhizal fungi’s use of nitrogen may broaden the habitat ranges of orchids, allowing them to grow in a variety of soil types.

Determining a Driving Force in Speciation

In a hybrid zone on the Northern Rocky Mountains, the flowering mustard plant *Boechera stricta* is evolving into a fitter species better adapted to its local environment. As reported April 3, 2017, in *Nature Ecology & Evolution*, a team including JGI researchers analyzed the mechanisms by which the *B. stricta* plants experienced positive directional selection.

As part of both the JGI’s CSP and Emerging Technologies Opportunity Program (ETOP), the researchers sequenced and analyzed the genome of *B. stricta*, a relative of the model plant *Arabidopsis*. With the genome in hand, they used techniques including gene mapping and chromosome painting methods to identify a major chromosomal inversion that controls ecologically important traits in the plant. They tested for QTLs, genome regions on chromosomes to which genetic traits can be mapped, in *B. stricta’s* chromosomal inversion. The researchers found several linked QTLs that changed ecologically important characteristics of the plant, such as flowering time and plant size, enabling it to adapt to its local environment, which in turn increased its fitness. The study shows that inversions that capture beneficial alleles such as QTLs can accelerate through the population and produce speciation.

This study offers the first direct evidence showing that QTLs are a driving force behind speciation. The knowledge gained from this study, said senior author and JGI collaborator Thomas Mitchell-Olds of Duke University, “gives evolutionary biologists experimental evidence showing how chromosomal changes contribute to adaptation and speciation. Furthermore, the genome sequence will help us understand how *Boechera* species are able to reproduce asexually by seeds, a process that can be used by farmers to speed up crop improvement practices.”
Sequenced and annotated as part of the JGI’s CSP, *Brachypodium distachyon* is a model grass for candidate bioenergy grasses such as *Miscanthus* and switchgrass. To further accelerate research in the development of biofuel feedstocks, a project to sequence thousands of *B. distachyon* mutants was selected for the 2015 CSP portfolio. This library of sequenced mutants will aid researchers to study and rapidly identify and order plants with mutations in any gene in their genomes.

Plants have evolved two kidney-shaped guard cells that swell to create the stomate. In grasses, however, the stomate have further evolved with the addition of two subsidiary cells flanking the guard cells, which may be linked to improved stomatal physiology. Unique to grasses, subsidiary cells have been associated with an improved physiological performance by allowing a greater range of pore size and quicker stomatal responsiveness. A plant’s ability to better control water loss and increase carbon assimilation could affect how it handles stressors such as drought, and play a role in the health and yields of candidate bioenergy feedstocks. Understanding the bases for water management could aid the identification and selection of individuals better suited for growing in otherwise marginal soils.

Using a genetic screen, a Stanford University team led by Dominique Bergmann identified a *B. distachyon* mutant unable to produce subsidiary cells. As reported March 17, 2017, in *Science*, by comparing the whole genome sequence of *B. distachyon* with this *sid* mutant, a 5-base pair deletion that encodes for the transcription factor BdMUTE was discovered. Further, BdMUTE was identified as the mobile transcription factor responsible for coordinating the development of subsidiary and guard cell complexes. The unique subsidiary cells in grasses may allow for an enhanced performance when stressors such as increased temperature or drought are placed on the plant. Though his contribution to the work predates his time at the JGI (he was then at the U.S. Department of Agriculture) Plant Functional Genomics lead and study co-author John Vogel provided the team with the mutant population and showed them how to manipulate the plant for their studies.
JGI researchers sequenced and analyzed the microbial communities in restored wetlands in the Sacramento-San Joaquin Delta (learn more on page 48).
Toward More Effective Carbon Fixation

Unlike DNA sequencing, where the language of life is read from the genome of an organism, DNA synthesis entails first the identification of a particular genetic element — such as an enzyme for fixing carbon from the atmosphere — and writing and expressing that code in a new system.

By tapping the DNA synthesis expertise of the JGI, a team from the Max Planck Institute (MPI) for Terrestrial Microbiology has reverse engineered a biosynthetic pathway for more effective carbon fixation. As reported November 18, 2016, in Science, this novel pathway is based on a new CO\textsubscript{2}-fixing enzyme that is nearly 20 times faster than the most prevalent enzyme in nature responsible for capturing CO\textsubscript{2} in plants by using sunlight as energy.

Photosynthesis harnesses sunlight to turn carbon dioxide into sugars that cells can use as energy along with other natural processes on the planet. It accounts for the transformation of some 350 billion tons of CO\textsubscript{2} annually. Researchers are concerned about how to capture the excess CO\textsubscript{2}, remove it from the atmosphere, and render it into energy and natural products for the economy. In support of the MPI team’s efforts, the JGI synthesized hundreds of Enoyl-CoA Carboxylase/Reductase (ECR) enzyme variants through its CSP. This enabled the MPI team to zero in on the ECR with the highest CO\textsubscript{2}-fixation activity to successfully build a more efficient artificial CO\textsubscript{2} fixation pathway in a test tube. Through sequencing and synthesis, they sourced 17 different enzymes from nine different organisms across the three kingdoms of life, and orchestrated these parts to achieve a proof of principle CO\textsubscript{2}-fixation pathway performance exceeding that found in nature. Erb dubbed it the “CETCH cycle,” because it “cetches” CO\textsubscript{2} more efficiently from the atmosphere.

Emboldened by the successful reconstitution of a synthetic enzymatic network in a test tube for the conversion of CO\textsubscript{2} into organic products that is superior to chemical processes and competes favorably with those in nature, Erb said this important discovery opens the door for other future applications. “These could include the introduction of synthetic CO\textsubscript{2}-fixation cycles into organisms to bolster natural photosynthesis, or say, in combination with photovoltaics, lead the way to artificial photosynthesis. This might at the end jumpstart the design of self-sustaining, completely synthetic carbon metabolism in bacterial and algal systems.”

Yasuo Yoshikuni, the head of the JGI’s DNA Synthesis Science group, looks to a future where DNA sequencing and biological functions further converge and leverage DNA synthesis. “Through JGI’s high-throughput sequencing capabilities coupled with the rapidly decreasing price of DNA synthesis, we continue to enable our user community in bringing to light the physiological potential of microorganisms and microbial communities. In the longer term, we hope to see these test-tube results yield a new generation of real bioproducts delivered to address critical energy and environmental challenges.”
Antarctic Adaptations in Diatoms

In the Antarctic Ocean, large populations of the diatom *Fragilariopsis cylindrus* dominate phytoplankton communities. Diatoms are a common type of photosynthetic microorganism, found in many environments from marine to soil. In the oceans, they are responsible for more than a third of the global ocean carbon captured during photosynthesis. This leads to a significant amount of sequestered carbon ending up in the sediments at the bottom of the ocean.

To learn more about how *F. cylindrus* adapted to its extremely cold environment, a team led by University of East Anglia (UEA) scientists conducted a comparative genomic analysis involving three diatoms by tapping the JGI’s sequencing and annotation expertise. Said UEA’s Thomas Mock, senior author of the work reported January 16, 2017, in *Nature*, “Our data provide first insights into how these key organisms underpinning one of the largest and unique marine ecosystems on Earth have evolved.”

The genome of *F. cylindrus* was sequenced as part of the JGI’s 2007 CSP portfolio. For the comparative analysis, its genome was compared against those of the diatoms *Thalassiosira pseudonana* and *Phaeodactylum tricornutum*, both found in temperate oceans with higher concentrations of dissolved iron, and both genomes previously reported by the JGI. *F. cylindrus* is diploid (has two copies of each chromosome, and thus two versions of each gene) and can selectively express the variant best suited to helping it deal with its environment. This provides additional genome-rooted resilience to the organism as its environment changes.

“Finding that the *F. cylindrus* population maintains and supports extensive variation to provide the adaptive ability of the population under harsh environmental conditions has broad implications for our understanding of how natural populations respond to changing environmental conditions,” said Jeremy Schmutz, head of the JGI’s Plant Program and a study co-author. “This will likely change the way the genomic techniques and assays are applied by the community to ocean-dwelling eukaryotic species.”

Algal Secrets to Thriving in the Intertidal Zone

The intertidal zone is the area between land and sea sometimes concealed by high tide or revealed by low tide. As this ecosystem is in constant flux, the organisms that inhabit the area have adapted to thrive under a range of constantly changing environmental conditions. *Porphyra* and other genera of bangiophyte red algae thrive in the intertidal zones of the northern and southern hemispheres.

As reported July 17, 2017, in the *Proceedings of the National Academy of Sciences*, the JGI sequenced, assembled, and annotated the genome of the red alga *Porphyra umbilicalis* to better understand how it harvests light and nutrients, and how warming oceans might affect its ability to fix carbon. Though red algae are one of the oldest multicellular lineages, only a few have had their genomes sequenced. *P. umbilicalis* is found in the ocean's intertidal zone, and is subject to constantly changing environmental conditions including temperature, light, and desiccation levels. As diatoms and other photosynthesizing microorganisms evolved from red algae, red algae metabolism had a significant impact on the planet's carbon cycle.

Green algae and red algae are both groups of plants that carry out photosynthesis using light-harnessing organelles called chloroplasts, which evolved from cyanobacteria that were engulfed by the ancestral eukaryotic algae. Later, other environmentally important algae, such as diatoms, dinoflagellates, and haptophytes, evolved when other non-photosynthetic eukaryotes captured red algae and integrated the red algal chloroplast and red algal nuclear genes into their genomes. These processes greatly diversified the organisms capable of conducting photosynthesis, and the red algal imprint on global productivity, aquatic food webs, and oxygen production is significant.

The team led by University of Maine researchers found that *P. umbilicalis* has previously unrecognized means of tolerating its physically stressful intertidal habitat. For example, the red alga has multiple strategies to protect cells from being damaged by high light levels, including expanded families of proteins that protect the photosynthetic apparatus from high light and unusual genomic arrangements of the genes that synthesize the mycosporine-like amino acids protecting against ultraviolet light. The team also found strong cytoskeletal limitations in *Porphyra* and most other red algae with sequenced genomes, offering a possible explanation for why red algae tend to be small compared with other multicellular eukaryotes.
Tiny Algae, Large Genomic Variation

As their name suggests, picophytoplankton such as Ostreococcus are invisible to the naked eye. Despite their size, their global abundance means they are a widespread primary producer and form the bases of several marine food webs. In coastal areas, they account for as much as 80% of the available biomass.

Ostreococcus is considered a model species to study marine picophytoplankton. A decade ago, JGI sequenced one of the Ostreococcus strains. That genome, along with other genome sequences from three Ostreococcus groups, revealed the tiny algae's diversity and adaptation to different ecological niches around the world. As reported July 5, 2017, in Science Advances, a team led by researchers at the Oceanological Observatory of Banyuls, France, and including JGI scientists, has resequenced and analyzed 13 members of a natural population of Ostreococcus tauri from the northwest Mediterranean Sea.

The analysis, enabled in part by the JGI's CSP, revealed that the O. tauri population is larger than anticipated, with high genetic and phenotypic diversity influenced by the algae's natural resistance to ocean viruses. The team identified two large candidate mating type loci, consistent with the pervasive evidence of recombination and thus sexual reproduction within the population. Understanding the genetic variability of various Ostreococcus strains will help researchers understand how environmental changes affect their abundance and ability to photosynthesize.
Nutrient Availability in Model Wetlands

Adjacent to the restored wetlands of Twitchell Island in California’s Sacramento-San Joaquin River Delta are rice fields with soil carbon contents that can vary between 2.5% and 25%, covering much of the global range of carbon found in soils. Conducting studies across a nutrient gradient allows researchers to better understand the relationship between carbon cycling, nutrient availability, and microbial communities in soil.

JGI researchers studied the ecosystems of Twitchell Island in the Sacramento-San Joaquin Delta, where the U.S. Geological Survey had a pilot study on restored wetlands. Rice fields are model wetland systems that allow researchers to focus on chosen biogeochemical variables, while factors such as water and vegetation are controlled. The DOE is interested in understanding the roles of microbial communities in long-term impacts on carbon emissions and carbon sequestration. Wetland ecosystems can trap as much as 30% of global soil carbon but contribute nearly 40% of global methane emissions, providing an opportunity to understand their roles as both carbon sinks and carbon sources. As reported July 21, 2017, in The ISME Journal, the team’s findings suggest that the microbial metabolic rates align with biological stoichiometry theory, a metabolic theory of ecology that suggests organisms with faster growth rates require more phosphorus to increase nitrogen-rich protein synthesis. Until now, this theory had not been applied to soil microbes in situ due to methodological limitations, which the scientists addressed using a novel genomic approach.

By studying the microbial communities in these soils, the researchers found that the rate at which microbes break down organic matter is coupled to the availability of carbon, nitrogen, and phosphorus in the soils. Specifically, the availability of phosphorus is a key factor in determining soil carbon cycling rates. An abundance of phosphorus increases microbial activity and metabolic rates, which in turn means higher carbon turnover. Lower phosphorus in high-carbon soils may help stabilize accumulated carbon, while high-phosphorus soils may more rapidly lose carbon stores. These associations at the ecosystem scale were also reflected in genomic data from the soil microbes that drive soil element cycling. In showing how microbial metabolism is regulated by coupled nutrient cycling and soil carbon availability, the researchers demonstrate how genomics studies of microbial communities can be scaled up to the ecosystems level, which will contribute to a deeper understanding of ecological processes and aid the development of better global carbon cycling models.
**Microbial Responses to Drilling for Oil**

Despite a shift toward more sustainable, alternative energy sources, people still rely on fossil fuels for energy and transportation. Owned by Denmark, the Halfdan North Sea oil field is an offshore petroleum reservoir managed by a joint venture that includes Shell, one of the world's largest oil companies. Over the past 15 years, 32 oil wells have been drilled, reaching depths over 2 kilometers below the seafloor to extract petroleum from this deep subsurface reservoir. Recovering gas and oil for energy and transportation fuels affects the previously isolated microbial communities in the deep subsurface. For example, water injections introduce new microbes to the existing indigenous populations, which can lead to the production of the highly toxic and corrosive hydrogen sulfide, altering oil well productivity and oil quality.

Understanding how microbial communities in subsurface petroleum reservoirs are affected by human activity, and how their responses can sour oil production, can influence oil industry practices. To learn more about how some of these populations respond to disruptions in their environment, researchers at Shell collaborated with the JGI and Newcastle University to analyze samples from the field's 32 oil wells. As reported May 19, 2017, in *The ISME Journal*, they conducted a comparative genetic analysis of the microbial communities in multiple oil wells within an offshore oil field.

They tracked the succession of microbial populations in oil wells drilled at different times, community compositions, and how these compositions change over time the longer an oil well stays in production mode. Tools developed at the JGI and made available through the IMG/M system were utilized during the analyses. The data generated provide insights into just how deep subsurface microbial communities are perturbed by active oil wells injecting foreign substances into these previously isolated populations, and help industry researchers develop new techniques for managing microbiological problems.

“For the first time, we are able to shed light into this remote and largely unexplored ecosystem,” said study senior author Nicolas Tsesmetzis of Shell International Exploration and Production Inc. “The largely heterogeneous and highly diverse microbial communities recovered from the different oil wells of the same oil field were rather unexpected and bring about a paradigm shift in our standard microbial monitoring practices of petroleum systems.”

Shell researchers collected samples from oil wells in a North Sea oil field like this one. *(Berardo62, Flickr, CC BY-SA 2.0)*
Lessons from Simulating a Deep Ocean Oil Spill

In the three months between the time the Deepwater Horizon Macondo well exploded on April 22, 2010, until it was capped in mid-July, thousands of barrels of oil and large quantities of natural gases flowed into the Gulf of Mexico daily. It was the first major release of oil and natural gases into the deep ocean (1,500 meters). Due to the depth of the spill, vast plumes of small oil droplets remained trapped deep in the ocean (900–1,300 meters) where they underwent biodegradation by the local microbial community.

A team led by Berkeley Lab researchers reproduced the dispersal to replicate the successive populations of diverse microbes over 64 days and recover high-quality draft genomes to determine metabolic factors driving microbial community shifts. They discovered that the microbial community transformed with chemical changes in residual oil. Additionally, their lab-based method allowed for the first time the successful resolution of high-quality genomes and the characterization of functional capabilities for all the key microbes. The expertise and resources used to reconstruct the microbial genomes demonstrates how new technology development at the JGI is enabling energy and environment research. Identifying the microbes involved in degrading hydrocarbons (the chief components of petroleum and natural gas), as well as the drivers that trigger successive waves of microbial responders, allows researchers to better understand how the microbial community adapted to the events of seven years ago.

The researchers presented the first complete picture of how successive waves of microbial populations degraded the released oil on June 26, 2017, in *Proceedings of the National Academy of Sciences*. Samples from three time points (days six, 18, and 64) were chosen for metagenomic sequencing and analysis. Assembled genome fragments were binned or assigned to draft genomes of origin using resources provided by the JGI’s Emerging Technologies Opportunity Program (ETOP). The researchers were also able to recover high-quality genomes of the key microbial players, and determine the metabolic factors driving the shifts between microbial communities. This research demonstrates the rapid, specialized biodegradation by the deep ocean microbial community was possibly driven by crude oil alone.
JGI-owned Compute Resources
170+ million NERSC hours

JGI BER Allocation
25 million NERSC hours

JAMO
5.5 million records

JAMO Archived Data Footprint
5.9 Petabytes

JGI File Systems
7.1 Petabytes of data
Computational Infrastructure

The Joint Genome Institute’s (JGI) capacity as a next-generation genomics user facility has generated petabytes of high-quality sequence data and analysis. To support this workload, the JGI has invested significant resources in its high-performance compute cluster, Genepool, as well as its storage and web infrastructure, adding 192 nodes of capacity in the Cori supercomputer.

The success of these computing projects is in part due to the JGI’s ongoing partnership with the National Energy Research Scientific Computing Center (NERSC), one of the nation’s foremost centers for high-performance computing. Both sides have learned a great deal since 2010, when all of the JGI’s computational resources were moved to NERSC. The infrastructure advancements to Genepool and other JGI portals mean rapid and smooth access for users across the globe. The JGI’s partnership with NERSC enables its researchers and users to devote more of their time to cutting-edge genomics research.

In 2016, NERSC acquired Cori, a $70-million investment in data-intensive and high-performance computing infrastructure. The JGI used more than 10 million central processing unit hours on this petascale supercomputer. Many of these calculations could not have been completed on the Genepool cluster because of the computing scale required. One of the core features of Cori is hardware called the Burst Buffer, an array of non-volatile random-access memory (NVRAM) nodes that will enable input/output-intensive workloads to run at scale on the new system. NERSC also deployed software called Shifter to allow Docker containers to run on the Cray supercomputers. This work grew out of collaboration with the JGI staff and has reduced the hurdles required to move bioinformatics, light source, and astronomy workloads to Cori. In 2017, in collaboration with NERSC consultants Daniel Udvary and Anthony Wildish, JGI staff undertook a major effort to move pipelines and software to Docker containers that enable JGI software to run on Cori, Genepool, the DOE Knowledgebase (KBase), and the cloud. The JGI also leverages Bioboxes where possible, to standardize the Docker container interfaces for bioinformatics tools. Bioboxes is an effort in the community that has been led in part by Michael Barton of the Research and Development Group.

The KBase and JGI released a new feature for users that enables scientists on the KBase platform to search and retrieve any public JGI data set. This search interface relies on an API to an Elastic Search instance built on top of the JGI Archive and Metadata Organizer (JAMO) developed by Harika Tandra and Chris Beecroft of the Sequence Data Management Group. In 2017, BBtools developed by Brian Bushnell and some of the DOE QC pipelines were made available on the KBase platform. The JGI is currently working with KBase to ensure JGI software and tools are available on the KBase platform via Docker and the KBase SDK.

The JGI and NERSC co-organized a workshop at the 12th Annual Genomics of Energy and Environment Meeting. In this workshop, users learned about Docker technology and got hands-on experience running containers in the cloud. In 2018, the computational workshops at the Genomics of Energy and Environment Meeting will focus on the KBase platform, and users can learn how to access NERSC resources via the KBase graphical user interface. In 2017, the JGI and NERSC accepted six proposals (see Appendix C) from the Microbiome Data Science call under the aegis of the Facilities Integrating Collaborations for User Science (FICUS) program. This joint call, led by Emiley Eloe-Fadrosh and Nikos Kyrpides, was issued by the two DOE user facilities and helps scientists perform large-scale computational analyses of sequence data.
Appendices
Appendix A
Acronyms at a Glance

<table>
<thead>
<tr>
<th>Acronym</th>
<th>Full Form</th>
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<tbody>
<tr>
<td>AF</td>
<td>Alignment Fraction</td>
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<tr>
<td>ANI</td>
<td>Average Nucleotide Identity</td>
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<tr>
<td>BER</td>
<td>DOE Office of Biological and Environmental Research</td>
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<td>BERAC</td>
<td>Biological and Environmental Research Advisory Committee</td>
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<td>BESC</td>
<td>BioEnergy Science Center (at ORNL)</td>
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<tr>
<td>BLAST</td>
<td>Basic Local Alignment Search Tool</td>
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<td>BOOST</td>
<td>Build Optimization Software Tools for DNA Synthesis</td>
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<tr>
<td>BRC</td>
<td>Bioenergy Research Center (i.e., BESC, CABBI, CBI, GLBRC, JBEI)</td>
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<tr>
<td>CABBI</td>
<td>Center for Advanced Bioenergy and Bioproducts Innovation (at the University of Illinois)</td>
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<td>CAMI</td>
<td>Critical Assessment of Metagenome Interpretation</td>
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<td>CBI</td>
<td>Center for Bioenergy Innovation (at ORNL)</td>
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<td>CBP</td>
<td>Consolidated Bioprocessing</td>
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<tr>
<td>CRISPR</td>
<td>Clustered Regularly Interspaced Short Palindromic Repeats</td>
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<td>CRADA</td>
<td>Cooperative Research &amp; Development Agreement</td>
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<td>CRT</td>
<td>Computational Research and Theory Building (at Berkeley Lab)</td>
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<td>CSP</td>
<td>Community Science Program</td>
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<td>DOE</td>
<td>Department of Energy</td>
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<td>ECP</td>
<td>Exascale Computing Project</td>
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<td>EMSL</td>
<td>Environmental Molecular Sciences Laboratory (at PNNL)</td>
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<td>ETOP</td>
<td>Emerging Technologies Opportunity Program</td>
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<td>FACS</td>
<td>Fluorescence Activated Cell Sorting</td>
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<td>FICUS</td>
<td>Facilities Integrating Collaborations for User Science</td>
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<td>FISH</td>
<td>Fluorescent In Situ Hybridization</td>
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<td>FY</td>
<td>Fiscal Year</td>
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<td>GEBA</td>
<td>Genomic Encyclopedia of Bacteria and Archaea</td>
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<td>GLBRC</td>
<td>Great Lakes Bioenergy Research Center (at the University of Wisconsin-Madison)</td>
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<td>GOLD</td>
<td>Genomes OnLine Database</td>
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<td>GSC</td>
<td>Genomic Standards Consortium</td>
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<td>HPC</td>
<td>High-Performance Computing</td>
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<td>HPPS</td>
<td>High-Performance Storage System</td>
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<td>IEP</td>
<td>Industry Engagement Program</td>
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<tr>
<td>IGB</td>
<td>Integrative Genomics Building</td>
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<tr>
<td>IMG</td>
<td>IMG/M Integrated Microbial Genomes &amp; Microbiomes System</td>
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<td>ISM</td>
<td>Integrated Safety Management</td>
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<td>ITS</td>
<td>Integrated Tracking System</td>
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<td>JAMO</td>
<td>JGI Archive and Metadata Organizer</td>
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<td>JBEI</td>
<td>Joint BioEnergy Institute (at Berkeley Lab)</td>
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<td>JLT</td>
<td>JGI Leadership Team</td>
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<td>KBase</td>
<td>DOE Systems Biology Knowledgebase</td>
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<td>LANL</td>
<td>Los Alamos National Laboratory</td>
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<td>LBNL</td>
<td>Lawrence Berkeley National Laboratory (Berkeley Lab)</td>
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<td>Acronym</td>
<td>Full Form</td>
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<tr>
<td>LLNL</td>
<td>Lawrence Livermore National Laboratory</td>
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<tr>
<td>MAG</td>
<td>Metagenome-Assembled Genome</td>
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<td>MIMAG</td>
<td>Minimum Information about a Metagenome-Assembled Genome</td>
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<tr>
<td>MISAG</td>
<td>Minimum Information About a Single Amplified Genome</td>
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<tr>
<td>MGM</td>
<td>Microbial Genomics and Metagenomics</td>
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<tr>
<td>NERSC</td>
<td>National Energy Research Scientific Computing Center</td>
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<td>NGEE</td>
<td>Next-Generation Ecosystem Experiments</td>
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<td>NREL</td>
<td>National Renewable Energy Laboratory</td>
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<tr>
<td>ORNL</td>
<td>Oak Ridge National Laboratory</td>
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<tr>
<td>PMO</td>
<td>Project Management Office</td>
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<td>PNN</td>
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<td>QTL</td>
<td>Quantitative Trait Loci</td>
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<td>SAC</td>
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<td>SAG</td>
<td>Single Amplified Genome</td>
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<td>SFA</td>
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<td>USDA-ARS</td>
<td>United States Department of Agriculture-Agricultural Research Service</td>
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<tr>
<td>WIP</td>
<td>Work Initiation Process</td>
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Appendix B

Glossary

**Annotation:** The process of identifying the locations of genes in a genome and determining what those genes do to improve accuracy of genetic information collected.

**Archaea:** One of the three domains of life (Eukarya and Bacteria being the others) that include primitive microorganisms that can tolerate extreme environmental conditions (temperature, acid, etc.).

**Assembly:** Aligning and merging fragments of a much longer DNA sequence to reconstruct the original sequence. This is required, as DNA sequencing technology cannot read whole genomes at once, but rather reads small pieces of between 20 and 1,000 bases, depending on the technology used.

**Barcoding:** The practice of appending known unique synthetic DNA sequences to sequencing libraries to allow pooling of libraries for next-generation sequencing, after which sequence data can be assigned to particular libraries or samples based on the barcode sequence.

**Base:** A unit of DNA. There are four bases: adenine (A), guanine (G), thymine (T), and cytosine (C). The sequence of bases constitutes the blueprint of life.

**Base pair:** Two DNA bases complementary to one another (A and T or G and C) located on the complementary strands of the DNA double helix.

**Biogeochemistry:** The field of study of the biosphere's interactions with the Earth's chemical environment.

**Bioinformatics:** The use of computers to collect, store, and analyze biological information.

**Biomass:** Material derived from living or recently living organisms, usually referring to plants or plant-derived material (lignocellulosic biomass). Biomass can serve as an energy source directly by burning or indirectly, after conversion into biofuels.

**BioproSpecting:** Searching nature for genes and proteins that can be applied to help scientists solve energy and environment challenges.

**Bioremediation:** The use of microorganisms to break down contaminants and other unwanted substances in waste and other substances.

**Bridge amplification:** A proprietary technique used by Illumina sequencing platforms to generate single-stranded clusters of template DNA.

**Carbon cycle:** The biogeochemical process by which carbon is exchanged among the planet's atmosphere, land, and oceans.

**CAZymes:** Carbohydrate-active enzymes that can break down plant polysaccharides, such as cellulose, into small sugars.

**Cellulose:** An organic compound made of a long chain of several hundred to over 10,000 glucose units. It is a critical part of the cell wall of plants and many algae.

**Cellulosic biofuel:** A type of liquid transportation fuel produced from lignocellulose, a structural material that makes up much of the mass of plants. Lignocellulose is composed mainly of cellulose, hemicellulose, and lignin.

**ChIP-Seq:** A method of analyzing protein interactions with DNA.

**Contig:** A contiguous sequence resulting from the assembly of smaller sequence fragments.

**Coverage:** The number of times a region of the genome has been sequenced during whole-genome shotgun sequencing.

**Curation:** Analysis of genome annotations to improve and maintain data presentation.

**Cyanobacteria:** A phylum of bacteria that obtain their energy through photosynthesis, and named for the color of the bacteria. Although often called blue-green algae, that name is a misnomer as cyanobacteria are prokaryotic and algae are eukaryotic.

**DAP-seq:** A high-throughput method for identifying protein binding sites in DNA.
**Draft genome (also called a draft assembly):** The term for an incomplete genome sequence. It can be applied to a wide range of sequences, from those that have the minimum amount of information needed for submission to a public database, to assembled genomes that have undergone manual and automatic review but still have sequence errors to be corrected.

**Enzyme:** A protein used to induce or speed up a chemical reaction.

**Eukaryotes:** The domain of life containing organisms that consist of one or more cells, each with a nucleus and other well-developed intracellular compartments. Eukaryotes include all animals, plants, and fungi.

**Finished genome:** In accordance with the 1996 Bermuda standard, a gapless sequence with a nucleotide error rate of one or less in 10,000 bases.

**Flow cell:** Resembles a microscopic slide, only with eight channels, on which DNA samples are loaded for analysis on Illumina sequencing platforms.

**Fluorescence-activated cell sorting:** A specialized type of flow cytometry used to study and purify cells. A heterogeneous mixture of cells passes through laser beams and is sorted into two or more containers, one cell at a time, based upon the specific light-scattering and fluorescent characteristics of each cell.

**Fosmid:** A vector suitable for cloning genomic inserts approximately 40 kilobases in size.

**GenBank:** Open-access, publicly available collection of annotated sequences submitted by individual laboratories and large-scale sequencing centers that is overseen by the National Center for Biotechnology Information.

**Hemicellulose:** An organic compound that is part of most plant cell walls and is made of 5-carbon sugars. Unlike cellulose, which is crystalline, strong, and resistant to being broken down, hemicellulose is much more fragile, and has a random structure.

**Informatics:** The science of information and computer information systems. At JGI, it is the science of managing and interpreting genomic information with computational tools.

**Library:** A collection of DNA fragments.

**Lignin:** A complex polymer of aromatic alcohols known as monolignols, usually derived from wood. It is a critical part of the cell wall of plants and many algae.

**Lignocellulosic biomass:** Biomass derived from plants, the most abundant raw material for the production of biofuels.

**Locus (plural loci):** The specific location of a gene or DNA sequence or position on a chromosome.

**Mapping:** Charting the location of genes on chromosomes.

**Mass spectrometry:** An analytical technique that can identify unknown compounds through their molecular weight. It can also be used to determine a molecule’s structure and chemical properties.

**Metabolomics:** A comparison of biological samples based on their metabolite profiles.

**Metagenomics (also environmental genomics or community genomics):** The study of genomes recovered from environmental samples through direct methods that do not require isolating or growing the organisms present. This field of research allows the genomic study of organisms that are not easily cultured in a laboratory.

**Metatranscriptomics:** The study of the region of the complete genetic code that is transcribed into RNA molecules and provides information on gene expression and gene function.

**Microbe:** Another name for a microorganism.

**Microbiome:** A defined environment within which a community of microbes exists and interacts.

**Molecular cloning:** The use of specialized DNA technology to produce multiple exact copies of a single gene or other segment of DNA to obtain enough material for further study.

**Multiple displacement amplification (MDA):** Method of amplifying tiny amounts of DNA in a cell so that it can be used for sequencing through single-cell genomics.
**Nitrogen cycle**: The biogeochemical process by which nitrogen is exchanged among the planet’s atmosphere, land, and oceans.

**Paired-end reads**: DNA library preparation technique that lets researchers look at both the forward and reverse template strands of a large DNA fragment and that provides positional information.

**Peptide**: Short chain of amino acids, the same compounds that make up proteins. Peptide chains are much shorter than the chains of amino acids that make up proteins.

**Phylogeny**: The evolutionary history of a molecule, such as a gene or protein, or a species.

**Polymerase chain reaction (PCR)**: A method of DNA amplification.

**Prokaryotes**: Unlike eukaryotes, these organisms (e.g., bacteria) are characterized by the absence of a nuclear membrane and by DNA that is not organized into chromosomes.

**Promoter**: A region of DNA that sends signals to a cell to tell it where a gene begins and when the gene is read. An inducible promoter signals the cell only under certain conditions while a constitutive promoter is always signaling the cell.

**Proteomics**: The large-scale study of proteins, as well as their structures and functions.

**Quantitative trait loci (QTL)**: Genome regions on chromosomes to which genetic traits can be mapped.

**RB-TnSeq**: Randomly-Barcoded Transposon Sequencing is a technique to generate sequence-tagged insertion mutant strains of a single-celled organism (typically bacteria) to simultaneously assess the functions of every gene in the genome in a variety of conditions by sequencing and counting the abundance of each tag.

**Read length**: The number of nucleotides tabulated by the DNA analyzer per DNA reaction well.

**Rhizosphere**: Microecosystem defined by a thin layer of soil where plant roots interact with microorganisms in the soil.

**Sequence**: Order of nucleotides (base sequence) in a nucleic acid molecule. In the case of DNA sequence, it is the precise ordering of the bases (A, T, G, and C) from which the DNA is composed. Also used as a verb to describe the process of determining the nucleotide order.

**Sequencing by synthesis**: Proprietary sequencing technique used by Illumina systems in which four fluorescently labeled nucleotides determine the sequence of a DNA fragment, one base at a time.

**Single-cell genomics**: Method for sequencing a genome using DNA derived from a single cell that is used to study uncultured or nonculturable organisms.

**Single-molecule real-time (SMRT) sequencing**: Single-molecule DNA sequencing performed in zero-mode waveguide (ZMW) chambers on a chip.

**Subcloning**: The process of transferring a cloned DNA fragment from one vector to another.

**Sulfur cycle**: The biogeochemical process by which sulfur is exchanged between the planet’s atmosphere, land, and oceans.

**Synthetic biology**: A field of research concerned with purposeful editing of biological systems. For JGI’s objectives, this process refers to assembling DNA sequence fragments with the goal of synthesizing sequences to experimentally validate their functions and applications.

**Transcriptome**: A collection of all the RNA transcripts in a given cell that serves as a snapshot of global gene expression.
Appendix C
2017 User Program Supported Proposals

**Community Science Program (CSP)**

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<td>Understanding polyploidy through the generation of the first sugarcane genome sequence</td>
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<td>Attwood, Graeme</td>
<td>AgResearch Ltd (New Zealand)</td>
<td>Defining gene function in rumen microbes</td>
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<td>Bohlmann, Joerg</td>
<td>University of British Columbia (Canada)</td>
<td>Exploring the G3 “Gymnosperm Giga-Genomes” for carbon sequestration, biofuels, and bioproducts</td>
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<td>Bonito, Gregory</td>
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<td>Bruns, Thomas</td>
<td>University of California, Berkeley</td>
<td>Functional genomics of pyrophilous fungi determining the fate of pyrolyzed carbon in post-fire soils</td>
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<td>Buchan, Alison</td>
<td>University of Tennessee Knoxville</td>
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<td>Coleman-Derr, Devin</td>
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<td>Cox, Michael</td>
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<td>Cullen, Daniel</td>
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<td>Metatranscriptome analysis of fungal decay of <em>Pinus contorta</em></td>
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<td>Cushman, John</td>
<td>University of Nevada</td>
<td>Ice plant gene atlas resource development for <em>Mesembryanthemum crystallinum</em>, a facultative crassulacean acid metabolism (CAM) model for improved water-use efficiency of bioenergy feedstocks</td>
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<td>Des Marais, David</td>
<td>Harvard University</td>
<td>Perenniality, abiotic stress tolerance, and biomass allocation in <em>Brachypodium</em>, a model grass genus for bioenergy</td>
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<td>Dijkstra, Paul</td>
<td>Northern Arizona University</td>
<td>Stress in microbial communities in response to changes in carbon and nitrogen availability</td>
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<td>Dopson, Mark</td>
<td>Linnaeus University (Sweden)</td>
<td>Exploring deep biosphere microbial communities by single-cell DNA sequencing</td>
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<td>Doty, Sharon</td>
<td>University of Washington</td>
<td>Functional genomics of poplar endophytes for elucidation of mechanisms of improved plant growth under challenging conditions</td>
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<td>Dudycha, Jeffry</td>
<td>University of South Carolina</td>
<td>Unlocking the photosynthetic diversity of cryptophyte algae through whole-genome sequencing</td>
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<td>Francis, Christopher</td>
<td>Stanford University</td>
<td>Spatiotemporal characterization of microbial communities controlling estuarine nitrogen and carbon cycling in the San Francisco Bay-Delta</td>
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<tr>
<td>Göker, Markus</td>
<td>DSMZ (Germany)</td>
<td>The One Thousand Microbial Genomes Phase 4 Project (KMG-4) – sequencing the most valuable type-strain genomes for metagenomic binning, comparative biology and taxonomic classification</td>
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<td>Harris, Steven</td>
<td>University of Nebraska – Lincoln</td>
<td>Fungal interaction networks in biological soil crusts</td>
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<td>Heyduk, Karolina</td>
<td>University of Georgia</td>
<td>Genome sequencing of C3 and CAM Yucca species</td>
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<td>Johnson, Matthew</td>
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<td>The role of acquired phototrophy in phytoplankton blooms: Insights from the <em>Mesodinium rubrum</em> genome</td>
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<td>Kellogg, Elizabeth</td>
<td>Donald Danforth Plant Science Center</td>
<td>Pan-genomics of big bluestem, a broadly adapted dominant grass</td>
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<td>LeBoldus, Jared</td>
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<td>RNAseq enabled metabolic modeling of disease resistance to <em>Septoria</em> canker in the DOE flagship <em>P. trichocarpa</em></td>
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<td>Lopez Peredo, Elena</td>
<td>Marine Biological Laboratory</td>
<td>Protecting photosynthesis during desiccation: do the genomes of desert-derived and aquatic <em>Scenedesmus</em> species hold the key to understanding extreme desiccation tolerance among green algae?</td>
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<td>MacGregor, Barbara</td>
<td>University of North Carolina</td>
<td>Single-cell (meta-)genomics of uncultivable large sulfur bacteria and their epibionts: Investigating host-microbe mediation of biogeochemical cycling</td>
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<td>Mockler, Todd</td>
<td>Danforth Center</td>
<td>A complete-sequence population for pan-genome analysis of sorghum</td>
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<td>Moran, Mary Ann</td>
<td>University of Georgia</td>
<td>Dynamics of bacterial carbon and sulfur cycling in a coastal environment</td>
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<td>Nguyen, Nhu</td>
<td>University of Hawai‘i at Manoa</td>
<td>A genome atlas of the ectomycorrhizal genus <em>Suillus</em>: Phylogenetic diversity and population genomics of a keystone guild of symbiotic forest fungi</td>
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<td>Nicholson, Wayne</td>
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<td>Transcriptomic and methylomic responses of <em>Carnobacterium</em> species to extreme low pressure</td>
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<td>Pett-Ridge, Jennifer</td>
<td>Lawrence Livermore National Laboratory</td>
<td>Microbial carbon transformations in wet tropical soils: Effects of redox fluctuation</td>
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<td>Raff, Jonathan</td>
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<td>Combined flux chamber and genomics approach to understanding soil emissions of reactive nitrogen oxides in a forested environment</td>
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<td>Shade, Ashley</td>
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<td>Greater than the sum of its parts? A synthetic microbial community approach to untangle member interactions and exometabolite production</td>
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<td>Smart, Christine</td>
<td>Cornell University</td>
<td>Genetic diversity of shrub willow pathogen <em>Melampsora americana</em> aided by genome sequence</td>
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<td>Stepanauskas, Ramunas</td>
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<td>Expanding the dark matter reference catalog by targeting taxonomic blind spots</td>
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<td>Whitman, William</td>
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<td>Core and pangenomes of soil and plant-associated prokaryotes</td>
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<td>Yang, Xiaohan</td>
<td>Oak Ridge National Laboratory</td>
<td>Gene atlas for <em>Kalanchoe laxiflora</em>, a obligate crassulacean acid metabolism (CAM) model for genetic improvement of water-use efficiency in bioenergy feedstocks</td>
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<td>Zhang, Chi</td>
<td>University of Nebraska – Lincoln</td>
<td>Genome sequencing of <em>Zygnematales</em>, the closest algal lineage to land plants, as a foundation for comparative genomic, transcriptomic, epigenetic, evolutionary, and biochemical studies</td>
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<td>Alvarez-Cohen, Lisa</td>
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<td>Unraveling functional dynamics and regulation crucial for the stability of an anaerobic ammonium oxidizing (anammox) community via community metatranscriptomics and 16S rRNA sequencing</td>
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<td>Averill, Colin</td>
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<td>Molecular mechanisms of ectomycorrhizal interactions that stabilize soil carbon</td>
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<td>Andras, Jason</td>
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<td>The effect of ecological restoration on the structure and function of soil microbial communities in coastal wetlands</td>
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<td>Bell, Terrence</td>
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<td>How interactions with soil microbiomes impact the survival and activity of a nitrogen-fixing soil surface consortium</td>
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<td>Bowen, Jennifer</td>
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<td>Building a foundation for understanding carbon and nitrogen cycling in microbially diverse salt marsh sediments</td>
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<td>Chistoserdova, Ludmila</td>
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<td>Probing the role of electron transfer redox mediators in methane-oxidizing communities</td>
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<td>Coleman, Maureen</td>
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<td>Cotner, James</td>
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<td>Translating stoichiometric diversity into genomic diversity: What elements are responsible for variability in bacterial biomass stoichiometry?</td>
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<td>Craft, Chris</td>
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<td>Microbial activity and community composition in a tidal freshwater marsh in response to sea level rise and saltwater intrusion: A field manipulation</td>
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<td>D'Agostino, Paul</td>
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<td>Dove, Nicholas</td>
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<td>Dynarski, Katherine</td>
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<td>Freedman, Zachary</td>
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<td>Assessing the recovery of microbial traits in bioenergy crop agroecosystems on reclaimed surface mines</td>
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<td>Gutierrez, Tony</td>
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<td>Hatzenpichler, Roland</td>
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<td>Genomic characterization of cosmopolitan sediment-dwelling archaea hypothesized to be involved in anaerobic carbon cycling</td>
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<td>Samuel, Buck</td>
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<td>Veley, Kira</td>
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<td>Venturi, Vittorio</td>
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<td>Sequencing of a set of identified and characterized rice bacterial endophytes</td>
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<td>Beisel, Chase</td>
<td>North Carolina State University</td>
<td>Profiling the functional diversity of CRISPR-Cas systems using cell-free transcription-translation systems</td>
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<tr>
<td>Bokinsky, Gregory</td>
<td>Delft University of Technology (Netherlands)</td>
<td>Plug adapters for biology: Activating heterologous iron-sulfur enzymes to fully exploit Nature’s catalytic potential</td>
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<tr>
<td>Chistoserdova, Ludmila</td>
<td>University of Washington</td>
<td>Establishing lanthanides as new life metals and understanding redox properties of lanthanide enzymes in metabolism of methane</td>
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<tr>
<td>Dueber, John</td>
<td>University of California, Berkeley</td>
<td>Identification of novel D-altronate dehydratases in the enolase superfamily enabling pectin utilization in <em>S. cerevisiae</em></td>
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<td>Huang, Possu</td>
<td>Stanford University</td>
<td>New protein platform for secondary metabolite detection</td>
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<td>Jewett, Michael</td>
<td>Northwestern University</td>
<td>Reframing combinatorial assembly and rapid prototyping of biosynthetic pathways with cell-free systems</td>
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<tr>
<td>Juminaga, Alex</td>
<td>LanzaTech, Inc.</td>
<td>Advancing understanding of acetogenic CO₂ fixation by generating a gene knock-out library to accelerate design of commercial strains for autotrophic production of fuels and bioproducts</td>
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<td>Mead, David</td>
<td>Varigen Biosciences</td>
<td>Metagenomic mining for next generation DNA polymerases</td>
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<td>Philimus, Benjamin</td>
<td>Oregon State University</td>
<td>Linking cyanobacterial orphan biosynthetic gene clusters to secondary metabolites</td>
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<td>Redding, Kevin</td>
<td>Arizona State University</td>
<td>Chloroplastic CO₂ reduction to formate supporting synthetic carbon fixation</td>
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<tr>
<td>Schmeing, Thomas</td>
<td>McGill University</td>
<td>Synthetic DNA to facilitate structural and functional understanding of nonribosomal peptide synthetase production of secondary metabolites</td>
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<tr>
<td>Scott, Kathleen</td>
<td>University of South Florida</td>
<td>Creation of constructs for functional expression of recently described inorganic carbon transporters widespread among sulfur- and iron-oxidizing chemolithoautotrophic microorganisms</td>
</tr>
<tr>
<td>Subramanian, Venkataramanan</td>
<td>National Renewable Energy Laboratory</td>
<td>Molecular engineering of <em>Trichoderma reesei</em> for improved cellulase production</td>
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<tr>
<td>Trudeau, Devin</td>
<td>Weizmann Institute of Science (Israel)</td>
<td>Genome mining to find a novel ribulose-1-phosphate kinase</td>
</tr>
<tr>
<td>Welander, Paula</td>
<td>Stanford University</td>
<td>Expression of novel triterpenoid biosynthesis proteins from environmental metagenomes</td>
</tr>
</tbody>
</table>
Facilities Integrating Collaboration for User Science (FICUS) Proposals

Through the joint Facilities Integrating Collaborations for User Science (FICUS) initiative, the JGI has partnered with other national user facilities and called for Collaborative Science Initiative proposals. The accepted proposals began on October 1, 2017, providing the researchers with access to the capabilities of both user facilities and datasets beyond what could be generated by either facility alone.

JGI-EMSL Collaborative FICUS proposals:

The FICUS JGI-EMSL call represents a unique opportunity for researchers to harness the combined power of genomics and molecular characterization in one research project to help advance the missions of the DOE’s Office of Biological and Environmental Research (BER).

<table>
<thead>
<tr>
<th>INVESTIGATOR</th>
<th>AFFILIATION</th>
<th>DESCRIPTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Baldrian, Petr</td>
<td>Institute of Microbiology, ASCR (Czech Republic)</td>
<td>The impacts of nitrogen availability and seasonal dynamics on plant-microbial interactions affecting C and N cycling in coniferous forest soils</td>
</tr>
<tr>
<td>Bartley, Laura</td>
<td>University of Oklahoma</td>
<td>Systems analysis of grass secondary cell wall development and regulation for biofuel production</td>
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<tr>
<td>Bianchi, Thomas</td>
<td>University of Florida</td>
<td>The role of priming effects on the conversion of blue carbon to CO₂ in the coastal zone</td>
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<tr>
<td>Blanchard, Jeffrey</td>
<td>University of Massachusetts Amherst</td>
<td>Molecular mechanisms underlying changes in the temperature sensitive respiration response of forest soils to long-term experimental warming</td>
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<tr>
<td>Cattolico, Rose Ann</td>
<td>University of Washington</td>
<td>Global warming induced salinity shifts: metabolic responses by algal-bacterial consortia</td>
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<tr>
<td>Fendorf, Scott</td>
<td>Stanford University</td>
<td>Metabolic constraints on organic matter decomposition and metal cycling in sediment deposits</td>
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<tr>
<td>Liao, Hui-Ling</td>
<td>Duke University</td>
<td>Combined ‘omics approaches for the study of ectomycorrhizal symbiosis between Suillus and Pinaceae, with emphasis on their role in nutrient cycling</td>
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<tr>
<td>Rich, Virginia</td>
<td>The Ohio State University</td>
<td>Something old, something new: systems-level insights into plant-microbial-permafrost carbon dynamics by parallel high-resolution organic matter and microbial meta-omics</td>
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<tr>
<td>Skerker, Jeffrey</td>
<td>University of California, Berkeley</td>
<td>Understanding conversion of biomass-derived carbon into lipids and terpenoids in the oleaginous yeast Rhodosporidium toruloides</td>
</tr>
<tr>
<td>Wrighton, Kelly</td>
<td>The Ohio State University</td>
<td>Deciphering controls on plant decomposition in Arctic ecosystems: Identifying unknown microbial condensed tannin degradation pathways</td>
</tr>
</tbody>
</table>
JGI-NERSC Collaborative FICUS proposals:

The expertise and capabilities available at the JGI and the National Energy Research Scientific Computing Center (NERSC) were made available through the FICUS JGI-NERSC call. Researchers can explore the wealth of genomic and metagenomic data generated worldwide through access to supercomputing resources and computational science experts to accelerate discoveries.

<table>
<thead>
<tr>
<th>INVESTIGATOR</th>
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<th>DESCRIPTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Babbitt, Patricia</td>
<td>University of California, San Francisco</td>
<td>Environmental profiling of enzyme superfamilies for function prediction</td>
</tr>
<tr>
<td>Baker, David</td>
<td>University of Washington</td>
<td>Eukaryotic protein structure determination using metagenome and metatranscriptome sequence data</td>
</tr>
<tr>
<td>Brooks, Phillip</td>
<td>University of California, Davis</td>
<td>Advancing metagenome classification and comparison by MinHash fingerprinting of IMG/M data sets</td>
</tr>
<tr>
<td>DeLong, Edward</td>
<td>University of Hawaii at Manoa</td>
<td>Quantitative analyses of naturally occurring small RNAs in global metagenomic and metatranscriptomic datasets</td>
</tr>
<tr>
<td>Hallam, Steven</td>
<td>University of British Columbia (Canada)</td>
<td>Charting global biogeochemical cycles using fast phylogenetic mapping of functional anchor genes</td>
</tr>
<tr>
<td>Konstantinidis, Konstantinos</td>
<td>Georgia Institute of Technology</td>
<td>Assessing microbiomes at the individual population level: Tool development and applications to soil carbon cycling</td>
</tr>
</tbody>
</table>
Appendix D
Advisory and Review Committee Members

The Scientific Advisory Committee (SAC)
The SAC is a board convened by the JGI Director to provide a scientific and technical overview of the JGI. Responsibilities of this board include providing technical input on large-scale production sequencing, new sequencing technologies, and related informatics; an overview of the scientific programs at the JGI; and an overview of the Community Science Program (CSP). A crucial job of the committee is to take the input from the CSP Proposal Study Panel on prioritization of CSP projects and, with DOE BER concurrence, set the final sequence allocation for this program.

Members
Mark Adams, J. Craig Venter Institute
Carol Bult, The Jackson Laboratory
Steve Briggs, University of California, San Diego
Jeff Dangl, University of North Carolina
Paul Flicek, European Bioinformatics Institute (ad hoc)
Claire M. Fraser, University of Maryland
N. Louise Glass, Lawrence Berkeley National Laboratory
Glenn Kubiak, Lawrence Berkeley National Laboratory
Mary Ann Moran, University of Georgia
Trina McMahon, Arizona State University
Deirdre Meldrum, University of Wisconsin-Madison
Juan Meza, University of California, Merced
Sue Wessler, University of California, Riverside (ad hoc)

Informatics Advisory Committee

Members
Adam Arkin, Lawrence Berkeley National Laboratory; University of California, Berkeley
Judith Blake, The Jackson Laboratory
David Dooling, Monsanto
Paul Flicek, European Molecular Biology Laboratory (EMBL)-European Bioinformatics Institute (EBI)
Saul Kravitz, The Howard Hughes Medical Institute (HHMI)
Jill Mesirov, University of California, San Diego (Chair)
Granger Sutton, J. Craig Venter Institute
Cathy Wu, Georgetown University
Kathy Yelick, Lawrence Berkeley National Laboratory

Fungal Program User Advisory Committee

Members
Scott Baker, Pacific Northwest National Laboratory
Randy Berka, ADM
Ronald de Vries, Westerdijk Fungal Biodiversity Institute (Netherlands)
Audrey Gasch, University of Wisconsin-Madison; Great Lakes Bioenergy Research Center
N. Louise Glass, University of California, Berkeley; Lawrence Berkeley National Laboratory
Stephen Goodwin, Purdue University
David Hibbett, Clark University
Francis Martin, INRA (France)
Michelle O’Malley, University of California, Santa Barbara
Joseph Spatafora, Oregon State University
Kathleen Treseder, University of California, Irvine
Adrian Tsang, Concordia University (Canada)

Plant Program User Advisory Committee

Members
Siobhan Brady, University of California, Davis
Gloria Coruzzi, New York University
Jeff Dangl, University of North Carolina, Chapel Hill
Joe Ecker, The Salk Institute for Biological Studies
Samuel Hazen, University of Massachusetts-Amherst
Tom Juenger, University of Texas, Austin
Toby Kellogg, Donald Danforth Plant Science Center
Sabeeha Merchant, University of California, Los Angeles
Stephen Moose, University of Illinois
Sue Rhee, Carnegie Institution for Science, Stanford
Bob Schmitz, University of Georgia
Gary Stacey, University of Missouri

Prokaryotic Super Program Advisory Committee

Members
Jill Banfield, University of California, Berkeley
Ed DeLong, University of Hawai‘i at Manoa
Jonathan Eisen, University of California, Davis
George Garrity, Michigan State University (Chair)

Steve Hallam, University of British Columbia
Phil Hugenholtz, University of Queensland (Australia)
Janet Jansson, Pacific Northwest National Laboratory
Kostas Konstantinidis, Georgia Institute of Technology
Monica Medina, Penn State University
Trina McMahon, University of Wisconsin-Madison (Vice-Chair)
Mary Ann Moran, University of Georgia
Nancy Moran, University of Texas at Austin
Jennifer Pett-Ridge, Lawrence Livermore National Laboratory
Rich Roberts, New England Biolabs
Ramunas Stepanauskas, Bigelow Laboratory for Ocean Sciences
Matt Sullivan, The Ohio State University

DNA Synthesis Science User Advisory Committee

Members
Richard Bailey, Independent Consultant
Doug Cameron, Firstgreen Partners
Sunil Chandran, Amyris, Inc.
Elizabeth Shank, University of North Carolina, Chapel Hill

James Flatt, Synthetic Genomics
Jay Keasling, Lawrence Berkeley National Laboratory
Megan Palmer, University of California, Berkeley
Elizabeth Sattely, Stanford University

David Weller, USDA-ARS
Appendix E
2017 Genomics of Energy and Environment Meeting

With more than attendees at the Walnut Creek Marriott, the 12th Annual Genomics of Energy and Environment Meeting took place March 20–23, 2017.

Keynote Speakers

“Our lab has started to ask different questions. What if we were able to find or design new enzymes for CO₂ fixation? And what if we were able to actually design new pathways that would give us faster access to biomass productivity and convert CO₂ to useful compounds?”

— Tobias Erb

Tobias Erb of the Max Planck Institute for Terrestrial Microbiology in Marburg, Germany opened the meeting. A microbial synthetic biologist, he spoke about the importance of understanding how biochemical pathways are made in nature to construct more efficient synthetic approaches. Case in point, his research team has been working to engineer an approach to fixing carbon that would be as good or better than the ones that have evolved from nature (see page 41).

Watch his talk on the JGI YouTube channel at bit.ly/JGI2017Erb

Read about the meeting at http://bit.ly/JGIPrimerSpring17
Learn more about the meeting talks at https://usermeeting.jgi.doe.gov/past-meetings/2017-agenda/
Videos of the talks are available on JGI’s YouTube channel at http://bit.ly/JGI2017_Videos
In his closing keynote, C. Titus Brown (2nd from the left above) from the University of California, Davis, spoke about the issues he’s faced wrestling with the unique challenges posed by genomic sequence analysis. As the datasets continue to pile on, he said, researchers must think about what they need to get out of the data, analyze that, and discard the rest. He also suggested that researchers take both past and future analyses of large-scale datasets to a higher level, and closed with a few recommendations: integrate multiple data types, develop better metadata exploration tools, and boost computational training.

Watch his talk on the JGI YouTube channel at bit.ly/JGI2017Brown.

Other Featured Speakers (in order of appearance):

Jan Leach, Colorado State University
Cat Adams, UC Berkeley
Colleen Hansel, WHOI
Sunny Liao, University of Florida
Jonathan Lynch, Penn State University
Renee Wegrzyn, Defense Advanced Research Projects Agency
Lauren Alteio, University of Massachusetts, Amherst
Patsy Babbitt, UC San Francisco
Philipp Zerbe, UC Davis
Mikael Andersen, Technical University of Denmark
Stephen Mondo, JGI
Gloria Coruzzi, New York University
Jay Chen, Oak Ridge National Laboratory
Frederik Schulz, JGI
Katherine (Trina) McMahon, University of Wisconsin – Madison
J. Chris Pires, University of Missouri
Thomas Mock, University of East Anglia
Alex Greenspan, University of California, Davis
Andrew Roger, Dalhousie University
Teresa Pawlowska, Cornell University
Abhishek Biswas, University of Tennessee
Mary Lipton, Environmental Molecular Sciences Laboratory
Harry Beller, JBEI
Appendix F

2017 Publications


Butterfield CN et al. Proteogenomic analyses indicate bacterial methylotrophy and archaeal heterotrophy are prevalent below the grass root zone. *PeerJ*. 2016 Nov 8;4:e2687.


NASA astronaut and microbiologist Kate Rubins, the first woman to sequence DNA in space, visited Berkeley Lab on May 11, 2017 and gave a talk about “Science in Extreme Environments.” As her talk was hosted by the Molecular Foundry and the JGI, she came out to Walnut Creek later that day. From left to right: CASIS associate program scientist Liz Warren, JGI User Programs Deputy Susannah Tringe, NASA astronaut Kate Rubins, JGI Director Nigel Mouncey, JGI Science Programs Deputy Axel Visel, and JGI Genomic Technologies Deputy Len Pennacchio.
Ongoing construction of the Integrative Genomics Building (IGB) as of January 2018, future home of the JGI and KBase at Berkeley Lab.

The work conducted by the U.S. Department of Energy Joint Genome Institute is supported by the Office of Science of the U.S. Department of Energy under Contract No. DE-AC02-05CH11231.