

# Office of Biological and **Environmental Research**

**Biological Systems Science Division** Update

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April 20, 2017

Office U.S. DEPARTMENT OF of Science

Office of Biological and Environmental Research

# **Programmatic Activities**

# **Completed Reviews/Activities**

- Annual Genomic Sciences Program PI Meeting (Feb 6-8)
- Annual JGI User Meeting (Mar 20-23)
- Year 9 annual progress review of the Bioenergy Research Centers
- Review of applications submitted to the re-competition FOA for the Bioenergy Research Centers

# **Upcoming Reviews/Activities**

- Panel Review of Biosystems Design FOA applications (next week)
- Panel Review of Plant Feedstocks for Bioenergy FOA applications (next week)
- Committee of Visitors review of BSSD (July 11-12)
- Lawrence Livermore National Laboratory SFA Review (July 18)
- Lawrence Berkeley National Laboratory SFA Review (Aug 1-2)
- New DOE Laboratory SFA Review (mid-Aug)
- DOE Systems Biology Knowledgebase review (Sept 26-27)
- Joint Genome Institute (JGI) triennial review (Dec 6-8)

# Strategic Planning/Activities

#### <u>Workshops</u>

#### ASCR-BER Workshop on Exascale Computational Requirements

Draft completed – out brief – this meeting (Dorothy Koch CESD, Ramana Madupu BSSD)

#### Technologies for Characterizing Molecular and Cellular Systems Relevant to Bioenergy and Environment

Draft completed – out brief – this meeting (Paul Adams, Co-Chair, LBNL)

#### National Academy Studies

- "Developing a Research Agenda for Utilizing Gaseous Carbon Waste Streams." Board on Chemical Sciences and Technology and the Board on Energy and Environmental Systems (BER, BES, ARPA-E, FE, EERE)
- "Biosecurity implications of gene editing tools" Board on Life Sciences
- "Breakthroughs 2030: A Process for a 10-year Agenda for Food and Agricultural Research."

Board on Agricultural and Natural Resources

# Funding Opportunities for FY 2017

#### Plant Feedstock Genomics for Bioenergy: A Joint Research Funding Opportunity Announcement USDA, DOE (DE-FOA-0001688)

- > Continues research on plant responses to pathogens
- > Includes a focus on oil seed crops
- Panel review next week

#### Biosystems Design to Enable Next-Generation Biofuels and Bioproducts (DE-FOA-0001650)

- > Plant and microbial genome-scale design and engineering
- > Expands its focus to biofuels and bioproducts

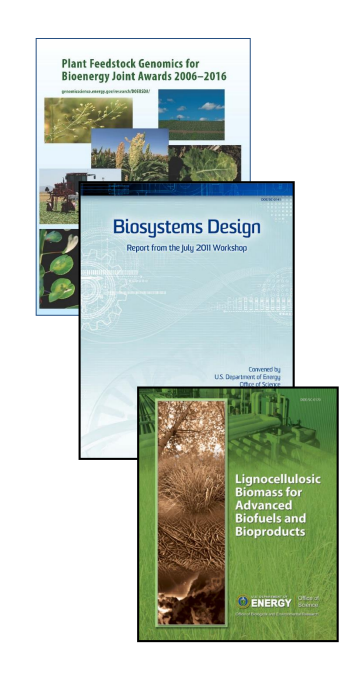
Panel review next week

#### DOE National Laboratories Science Focus Area (SFA) Opportunity

- Plant Systems Biology for Bioenergy
- Biosystems Design for Bioenergy
- Soil Microbiome Research

## Systems Biology for Bioenergy FOA

Posting TBD



# Bioenergy Research Centers - 10<sup>th</sup> Year!!

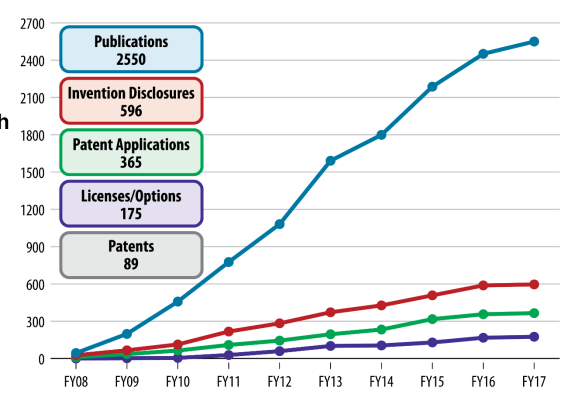
Multidisciplinary fundamental science guided by milestones & deliverables, targeted to key areas needed to improve production of biofuels from renewable biomass.







- BioEnergy Science Center (Oak Ridge National Laboratory)
- Great Lakes Bioenergy Research Center (Univ. of Wisconsin, Michigan State Univ.)
- Joint BioEnergy Institute (Lawrence Berkeley National Laboratory)







Key traits leading to reduced recalcitrance remain stable following three years of field trials

**Objective** Evaluate retention of lignin-altered traits in field trials of lignin altered switchgrass.

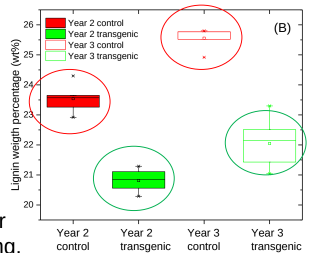
## Approach

Chemical composition, cellulose crystallinity and degree of polymerization and cellulose accessibility were measured in down-regulated COMT switchgrass following growth in the field for two and three seasons.

#### Results/Impact

- Reduced lignin content and resultant reduced biomass recalcitrance remained stable in field-grown trials for at least three seasons.
- The findings are consistent with the originally reported improvements in cellulose accessibility via down-regulation of COMT in switchgrass.
- Demonstrates the long(er) term stability of introduced traits for improved biofuel production in bioenergy crops in a field setting.





Li, M. et.al., "Study of traits and recalcitrance reduction of field-grown COMT down-regulated switchgrass" *Biotechnol Biofuels* (2017). DOI: 10.1186/s13068-016-0695-7

## <u>BRC Science Highlight</u>

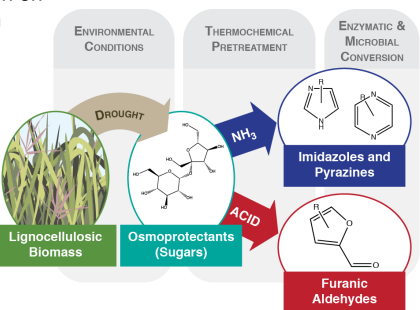


# Inhibition of microbial biofuel production in drought stressed switchgrass hydrolysate

**Objective** Examine impact of variable precipitation on biomass hydrolysate composition and fermentation

## Approach

Corn stover and switchgrass samples collected from field sites over a 3-year period
Time period included a major drought year and two years of average precipitation
Biomass samples were analyzed for hydrolysate composition, and fermentative performance.



#### Result/Impacts

- Growth of yeast was completely inhibited in hydrolysate made from drought stressed switchgrass and contained elevated levels of pyrazines and imidazoles
- Environmental variation can have significant effects on biomass hydrolysate properties
- Knowledge of how these factors impact biofuel production can guide pretreatment and biocatalyst choices to mitigate these inhibitory effects.

Ong, R. G. *et al.* Inhibition of microbial biofuel production in drought stressed switchgrass hydrolysate. *Biotechnology for Biofuels* (2016) DOI: 10.1186/s13068-016-0657-0

April 2017 BERAC Meeting

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#### BRC Science Highlight



# • On-chip integration of droplet microfluidics with nanostructure initiator mass spectrometry for enzyme assay

## **Objective**

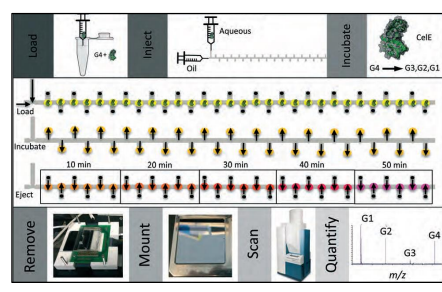
Develop new high throughput technologies for rapid screening of enzyme activity

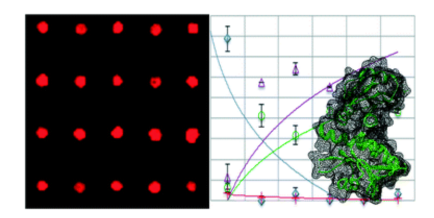
## Approach

- Designed microfluidics chip for assaying enzyme activity with 1000x fold reduction in reagent consumption.
- Integrated microfluidics with NIMS to establish a new screening platform

#### Result/Impacts

- Device can manipulate150 nl droplets with subsequent deposition onto the NIMS surface, achieving a significant decrease in reagent use
- Disruptive technology based on a grid of 50 µm resolution that enables >100 000 pads per 5 cm2 NIMS array when scaling the current design
- Potential significant impact on enzyme engineering, DNA manipulation, microbial screening for bioenergy applications





Heinemann et al. (2017) "On-chip integration of droplet microfluidics and nanostructure-initiator mass spectrometry for enzyme screening". Lab Chip. doi, 10.1039/c6lc01182a

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Genomic Science Program-Carbon Cycling

# Microbial Community Interactions Drive Methane Consumption in Lakes

**Objective** Evaluate methane metabolism in lake sediments

## Approach

- Assembled a microbial community consisting of methanotrophs and co-occurring nonmethanotrophic bacteria found in lake sediments
- Used community-scale meta-omics analyses of shifts in gene expression to track metabolism of associated microbes during methane-driven growth



#### Results/Impact

Image courtesy of iStock

- In the co-culture system, presence of non-methanotrophs altered the gene expression and metabolism of the methanotrophs to convert a portion of the available methane to methanol
- Methanol served as an energy source for the non-methanotrophs suggesting a cross-feeding mechanism
- Beneficial partner-induced changes in gene expression has implications for understanding the microbial interactions occurring in complex environments

SMB Krause et al. 2017. "Lanthanide-dependent cross-feeding of methane-derived carbon is linked by microbial community interactions." **PNAS** [DOI: 10.1073/pnas.1619871114]

Genomic Science Program-Biosystems Design

# **Engineering Oleaginous Yeast for High Productivity of Biodiesel from Sugars**

Glucose -

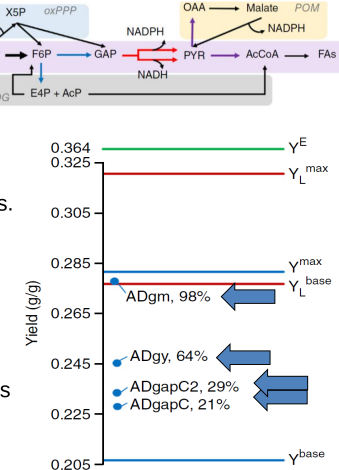
**Objective** Engineer Yarrowia lipolytica for high oil production yield

## Approach

- Model results identified a metabolic bottleneck in oil biosynthesis pathway
- Introduced heterologous yeast and bacterial genes to convert NADH into NADPH, required for lipid biosynthesis.
- Increased oil production by overexpressing a bacterial malic enzyme (MCE2)

## Results/Impact

- Lipid production in the engineered Y. lipolytica increased 25% relative to control strains.
- Approach could be optimized for converting plant biomass into biofuel precursors and other DOE-relevant bioproducts.



K. Qiao, T. Wasylenko, K.Zhou, P.Xu & G. Stephanopoulos, "Lipid production in *Yarrowia lipolytica* is maximized by engineering cytosolic redox metabolism." *Nature Biotechnology* 35, 173 (2017)

#### Genomic Science Program-Plant Feedstocks

# Phosphate Stress and Immunity Systems in Plants are Orchestrated by the Root Microbial Community

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**Objective:** Evaluate how plant immune systems coordinate microbial recognition with nutritional cues during microbiome assembly.

#### Approach

Mutants of Arabidopsis thaliana with altered phosphate starvation response (PSR) to test impact of genes controlling PSR on normal root microbiome assembly

## Results/Impact

- PSR regulation and pathogen defense are coordinated, providing insight into the coordinated interchange of plant response to nutritional stress, the plant immune system, and the root microbiome
- Provides a foundational basis for using the soil microbiome to enhance phosphate use efficiency in plants.

Castrillo *et a*l "Root microbiota drive direct integration of phosphate stress and immunity." *Nature* 543, 513-518 (2017). [10.1038/nature21417]



#### Plant-Microbe Interfaces Scientific Focus Area

# **Populus trichocarpa encodes small, effector-like secreted** proteins that are highly induced during mutualistic symbiosis

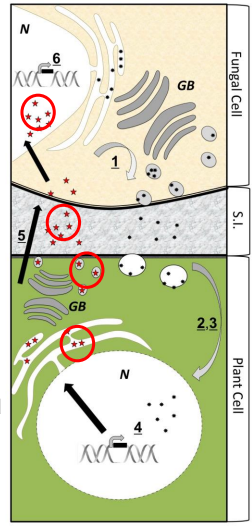
**Objective** Evaluate plant-microbe interactions in a *Populus-Laccaria* system

## Approach

- Computational analysis of the *Populus* transcriptome identified 2,819 protein-encoding genes that exhibited differential transcript abundance during mycorrhizal root tip development during symbiosis with *L. bicolor*.
- Of these, 417 were predicted to be small secreted proteins or SSPs (<250 aa in length).</p>
- A subset of SSPs were able to enter *L. bicolor* hyphae and accumulate in the nucleus altering development of multiple ectomycorrhizal and pathogenic fungi.

## Results/Impact

- This study demonstrates that SSPs in *Populus* can be secreted and function as effector proteins during symbiotic interactions.
- This study highlights a novel avenue by which plants communicate with (or control) their mutualistic microbial partners.



Plett J.M. *et al.* (2017). *Populus trichocarpa* encodes small, effector-like secreted proteins that are highly induced during mutualistic symbiosis. *Nature Scientific Reports* 

#### ENIGMA Scientific Focus Area



**Objective** Improve the annotation of genes of unknown function in bacterial genomes.

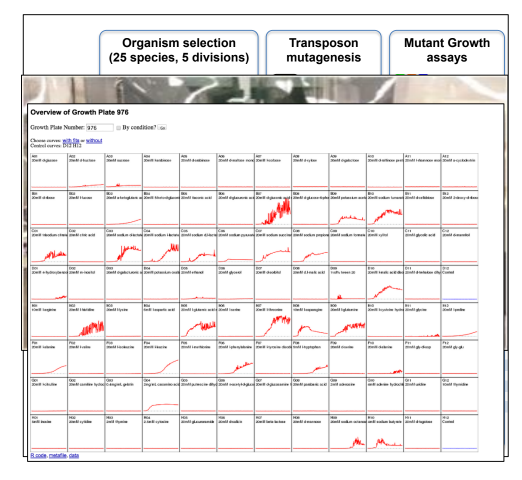
## Approach

Assay transposon mutant libraries from 25 diverse bacteria across hundreds of distinct conditions

#### Results/Impact

- From 3,903 mutant fitness assays, 14.9 million gene phenotype measurements were collected & the mutant phenotypes for 8,487 proteins with previous unknown functions were identified
- Demonstrates utility of high-throughput genetics for large-scale annotation of bacterial proteins
- Provides a large collection of experimentally-determined protein functions across diverse bacteria

## Deep Annotation of Protein Function across Diverse Bacteria from Mutant Phenotypes

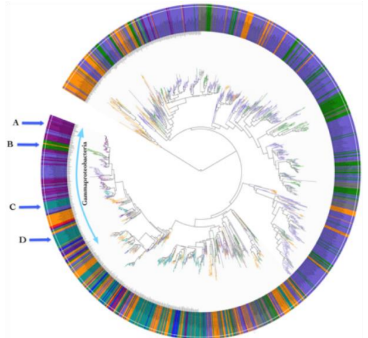


Price et al. "Deep Annotation of Protein Function across Diverse Bacteria from Mutant Phenoypes" (in review)

#### KBase PREDICTIVE BIOLOGY

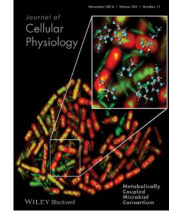
#### What's new In KBase?

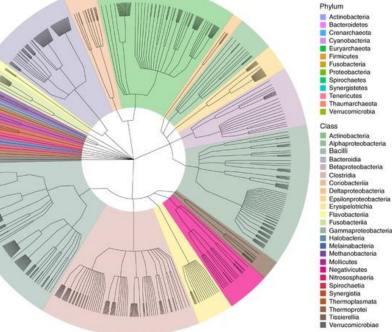
- Over 60 apps spanning assembly, annotation, comparative genomics, metabolic modeling, expression analysis, RNA-seq, and more
- Enhanced RNA-seq and metabolic modeling
- Upload large data objects from the web
- Upload and download select data in bulk
- Greatly enhanced tools supporting 3rd-party software tool development



 Constructed, compared and analyzed core metabolic models of over 8000 microbes (Edirisinghe et al., *BMC Genomics*)  Modeled and predicted interactions between heterotrophic and autotrophic species in a simple community (Henry et al., *J. of Cellular Physiology*)

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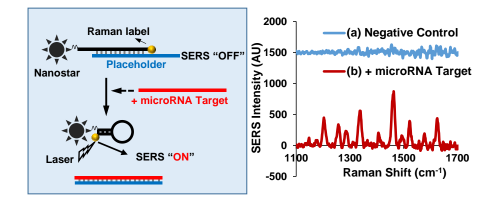
 Reconstructed 773 metabolic models of microbiome isolates (Magnúsdóttir et al., *Nature Biotech*) BER M2M Bioimaging Technologies

DUKE UNIVERSITY Fitzpatrick Institute for Photonics Multiplexed Detection of MicroRNA Biomarkers Using SERS-Based Inverse Molecular Sentinel (iMS) Nanoprobes

**Objective** Probe development for microRNAs detection in plants

## Approach

MicroRNAs play an important role in plant development such as plant floral transition



- SERS-based Inverse Molecular Sentinel (iMS) nanoprobes were developed with plasmonicactive nanostars as the sensing platform to detect microRNAs
- Differently-labeled iMS nanoprobes were designed and developed for multiplexed detection of miR-21 and miR-34a microRNAs in extracted RNA samples.

#### Results/Impact

The iMS nanoprobe as a homogeneous assay for multiplexed detection of microRNAs in extracted RNA samples without target amplification steps. The iMS biosensing platform could provide a useful tool to investigate microRNA expression.



# Seeking Structures With Metagenome Sequences

**Objective** Use metagenomic sequences to fill gaps in structural information for known protein families.

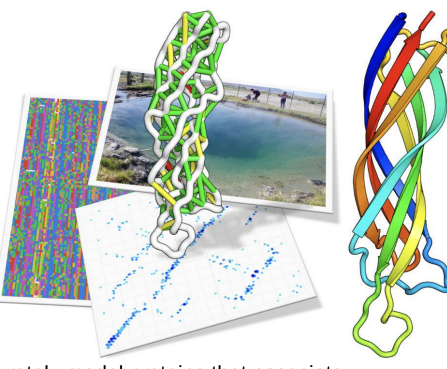
## Approach

Integrate metagenome data, contact-based structure matching, and Rosetta structure calculations to model unknown proteins.

#### Results/Impact

- Rosetta structure predictions are able to accurately model proteins that associate to large families
- Metagenome sequence data more than tripled the number of protein families with sufficient sequences for accurate modeling
- Generated models for 614 protein families with currently unknown structures; 206 putative membrane proteins and 137 with folds not represented in the Protein DataBase

Ovchinnikov et. al. "Protein Structure Determination Using Metagenome Sequence Data." *Science* 355. 6322 (2017) [DOI:10.1126/science/aah4043]





- Novel group of giant viruses discovered, Science
- Engineering a more efficient system for harnessing carbon dioxide, Science
- First CRISPR-Cas9 system in Archaea discovered, Nature
- Tracking Antarctic adaptations in Diatoms, Nature
- Metagenomics database helps fill in 10 percent of previously unknown protein structures, Science
- Harnessing Aspergillus diversity for industrial applications, Genome Biology
- Speciation driven by alleles adapted to local conditions, Nature Ecology
- IDing grass gene for stomatal morphology underscores importance of developing mutant gene index, **Science**
- Novel mechanism in bacterial-fungal symbiosis could have biodiesel production applications, **PNAS**
- 77 papers published in Q1/Q2 (through April 7, 2017)



## New Structural Biology Portal

- "One stop shop" for information about BER structural biology resources
- Highlights of the latest science developments
- Descriptions of capabilities
- Information for new users

# NEW BER Structural Biology Portal BERStructuralBioPortal.org

Bringing together information about BER's structural biology resources





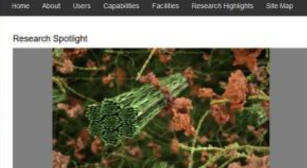






#### BER Structural Biology Resources at Synchrotron and Neutron Facilities

Enabling innovative and discovery science in support of the DOE-BER mission



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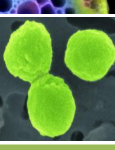
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Systems science to meet DOE mission needs in bioenergy, climate and the environment.

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# Thank you!



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ACCESS MEETING ABSTRACT!