CAMERA A Metagenomics Resource for Microbial Ecology

> Saul A. Kravitz Rekha Seshadri J. Craig Venter Institute

DOE GTL PI Meeting February 12, 2008









Presentation Outline

- Introduction to Metagenomics
- Global Ocean Sampling (GOS) Expedition
- Challenges of Metagenomics
- The CAMERA Project
- Live Demonstration of CAMERA



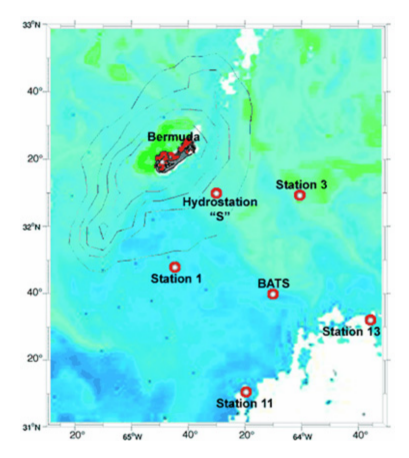
Genomics vs Metagenomics

- Genomics 'Old School'
 - Study of a single organism's genome
 - Genome sequence determined using shotgun sequencing and assembly
 - ~1300 microbes sequenced, first in 1995
 - DNA usually obtained from pure cultures (<1%)
- Metagenomics
 - Application of genome sequencing methods to environmental samples (no culturing)
 - Environmental shotgun sequencing is the most widely used approach
 - Environmental Metadata provides key context



Metadata

- Metagenomics
 - Genomics + Metadata
- Environmental Metadata
 - Time and location (lat, long, depth) of sample collection
 - Correlate w/remote sensing data
 - Physico-chemical properties (e.g. temperature, nutrients, salinity)



MODIS-Aqua satellite image of ocean chlorophyll in the Sargasso Sea grid about the BATS site from 22 February 2003





Metagenomic Questions

- Within an environment
 - What biological functions are present (absent)?
 - What organisms are present (absent)
- Compare data from (dis)similar environments
 - What are the fundamental rules of microbial ecology
- How do organisms adapt to environmental conditions?
- Search for novel proteins and protein families



Complexity of Microbial Communities

- Simple (e.g., AMD, gutless worm)
 - Few species present (<10)
 - Diverse
 - ➔ Variations on standard genomics techniques
- Complex (e.g., Soil or Marine)
 - Many species present (>10, often >1000)
 - Many closely related
 - ➔ New techniques





JCVI Global Ocean Sampling Expedition Largest Metagenomic Study to Date











Global Ocean Sampling (GOS)

178 Total Sampling Locations

Phase 1: 41 samples, 7.7M reads, >6M proteins

Diverse Environments

Open ocean, estuary, embayment, upwelling, fringing reef, atoll, warm seep, mangrove, fresh water, biofilms, sediments, soils



Challenges of Complex Metagenomic Datasets

- Most sequence reads are unique
 - Very limited assembly
 - Most sequences not taxonomically anchored
 - Annotation of annotated reads required
 - Relating shotgun data to reference genomes
- Annotation challenging
 - Which ORFs code for predicted proteins?
 - Lack of genomic context
 - Large data volume







Motivations for CAMERA

- Significant investment in sequencing
 - Only accessible to bioinformatics elite
 - Diversity of user sophistication and needs
- Bioinformatics and Computation Challenges
 - Assembly, annotation, comparative analysis, visualization
 - Dedicated compute resources
- Importance of Metadata
 - Metadata required for environmental analysis
 - Need to drive standards
- Compliance with Convention on Biodiversity









CAMERA Team

- J. Craig Venter Institute
 - Marv Frazier, co-PI
 - Saul Kravitz, Program Manager
 - Rekha Seshadri, Chief Scientist
- UCSD/Calit2
 - Larry Smarr, co-PI
 - Paul Gilna, Executive Director
- UC Davis
 - Jonathan Eisen, co-investigator
- Gordon and Betty Moore Foundation
 - David Kingsbury and Mary Maxon





J. Craig Venter

NSTIT





Demo of CAMERA 1.3

- Data and Metadata Collections
 - ALL metagenomic nt sequence and metadata
 - Coding sequences functionally annotated
- BLAST with Environmental Metadata
 - Large query and subject sizes
 - 454 FLX run or microbe VS All Metagenomic
- Fragment Recruitment Viewer
 - Based on Rusch's GOS viewer
- Text Search of Annotations











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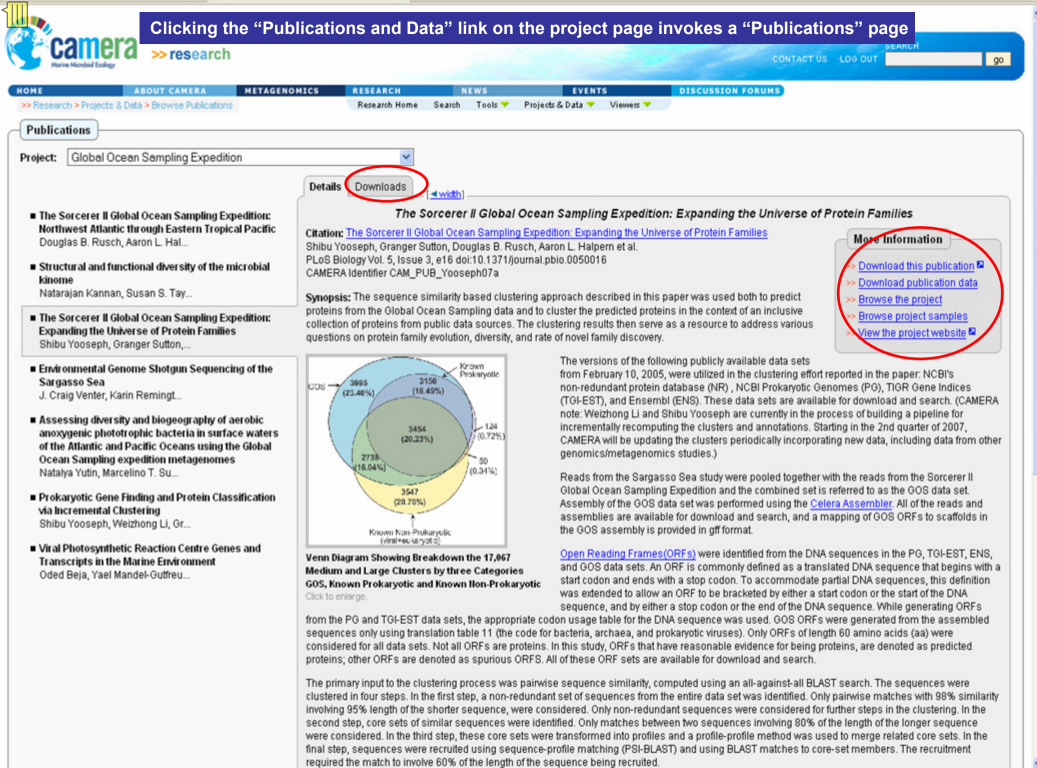
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This is the "Projects" page. A synopsis of the selected project is displayed
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rojects	Details
	Global Ocean Sampling Expedition
 Acid Mine Drainage Metagenome 	Principal Investigator: J. Craig Venter Funded By: Moore Foundation, DOE, Venter Science Foundation
 Alvinella pompejana Epibiont Metagenome 	Organization: J. Craig Venter Institute >> Publications & Data >> Website I
 Chesapeake Bay Virioplankton Metagenome (Data Coming Soon) 	The broad objective of the global ocean sampling expedition is to expand our understanding of the microbial world by studying the gene complement of marine microbial communities. Marine microbes influence the cycling of carbon (and other elements) in the world's oceans, acting as a biological conduit that transports carbon dioxide from the surface to the deep oceanic realms.
 Global Ocean Sampling Expedition 	By sequestering carbon from the atmosphere, marine microorganisms (eukaryotes, prokaryotes and viruses) may significantly affect global climate. How they do so, however, is poorly understood, and our attempts to study their activities are limited by our inability to culture the vast majority of them.
 Mediterranean Gutless Worm Metagenome 	One avenue of exploration is to sequence the genomes of marine microbes using a metagenomics approach. In Spring of 2003, the J. Craig Venter Institute conducted a whole environment shotgun sequencing project to study marine microorganisms in the nutrient-poor Sargasso Sea near Bermuda. This study
 Mediterranenan Bathypelagic Habitat Metagenome 	revealed an unforeseen breadth and depth of microbial diversity - about 1,800 different microbial species encoding over 1.2 million genes were discovered, nearly doubling the number of prokaryotic genes available in public databases.
 Metagenome of Marine NaCI-Saturated Brine 	Notably, this study expanded our knowledge of ocean photobiology and nutrient pools. Results from the pilot study were reported in Science in 2004.
 Microbial Community Genomics at the HOT/ALOHA 	This pilot study served as the springboard for launching a more comprehensive full collection of articles survey of the bacterial, archaeal and viral diversity of the world's oceans. A global circumnavigation aboard the Sorcerer II sailing yacht began in August 2003, starting in Halifax, Canada and samples were collected at sites along the U.S. east coast, Gulf of
 Moore Marine Microbial Sequencing 	Mexico, Galapagos Islands, central and south Pacific Oceans, Australia, Indian Ocean, South Africa, across the Atlantic back to the U.S., and was completed in January 2006. An initial analysis of the microbial data from the first leg of the trip - Halifax to the Galapagos Islands - was reported in PLoS Biology in January
Ocean Viruses	The Sorcereril Research Vessel Courtesy - J.Craig Venter Institute 2007. Shotgun sequencing and deep sequencing of 16S and 18S rRNA continues on the samples from the
 Phosphorous Removing (EBPR) Sludge Community 	These studies have produced the largest catalogue of genes to date from thousands of new species, with no apparent slowing of the rate of discovery (i.e., attaining saturation of data). These data have potentially far-reaching implications for biological energy production, bioremediation, and creating solutions for reduction/management of greenhouse
 Viral Metagenomes from Terrestrial Hot Springs 	gas levels in our biosphere. For more information, see the J.Craig Venter Institute's <u>Global Ocean Sampling Project</u> . JCVI's GOS data is also available through NCBI's <u>Genban k</u>
 Waseca County Farm Soil Metagenome 	JCVI has had many requests for GOS materials (clones, templates, DNA). Unfortunately, managing this large collection of materials is an expensive endeavor, and JCVI is not curently equipped to provide this service. JCVI is exploring the possibility of a 3rd party taking on this role, but there are no definitive plans as of yet.

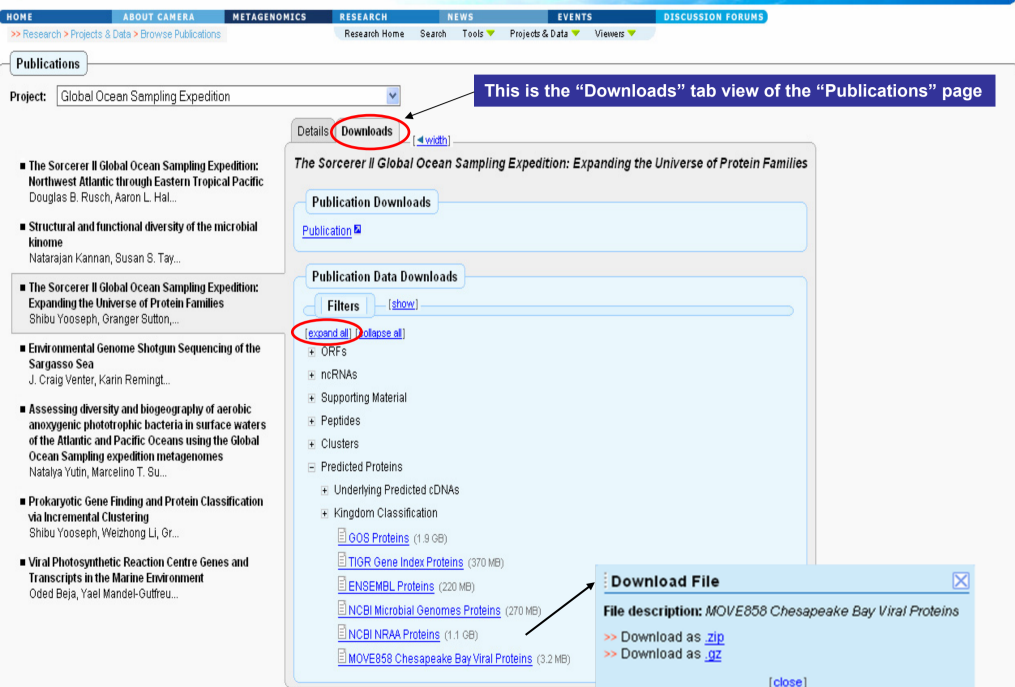
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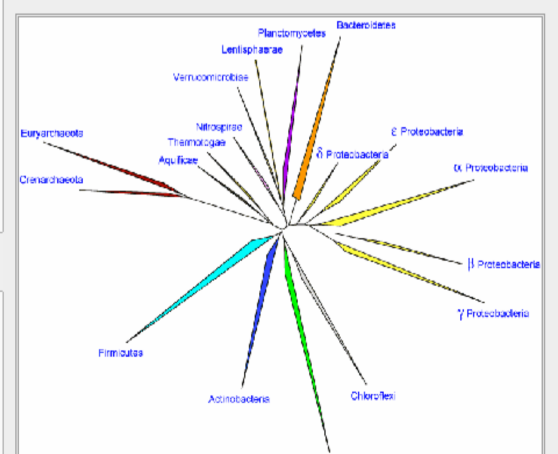
Moore Marine Microbial Sequencing

Principal Investigator:	Various
Funded By:	The Gordon and Betty Moore Foundation
Organization:	The Gordon and Betty Moore Foundation
Affiliation:	The J. Craig Venter Institute

More Information >> <u>Samples</u> >> <u>Publications & Data</u> >> <u>Website</u> >> <u>Contact</u>

The overarching goal of the marine microbial sequencing project is to explore the distribution, function and ecological role of marine microbes, and their role in the overall health of oceans and climates. The field of marine microbiology has made amazing progress in recent years through advances in genomic and metagenomic

technologies, and is causing us to reevaluate many of the basic precepts of microbial ecology and evolution. Nonetheless, out of the 300+ prokaryotic genomes that have been completely sequenced to date, only about 18 are from the marine environment. In an attempt to bridge this gap, we are sequencing the draft genomes of over 155 diverse, ecologically relevant marine microorganisms that were selected for sequencing based on the following criteria:



1.Open ocean and coastal bacterioplankton isolates most closely related to uncultured marine clades abundant in ribosomal RNA surveys.

2.Open ocean and coastal bacterioplankton isolates for .cluster sequencing, from ecologically relevant marine clades.

3.Bacterioplankton isolates representing broad physiological diversity (photoautotrophs, photoheterotrophs, nitrifiers, methanotrophs, etc.).

4.Isolates from diverse environments/habitats (greater depths, more extreme temperatures, specialized habitats).

These genome data not only hold the promise of delivering tremendous biological insight through individual and broad comparative genome analyses, but also provide a key reference collection for culture-independent metagenomic surveys like the global ocean sampling expedition and many others.

The project was initiated in Fall 2004 with a grant to the J. Craig Venter Institute. Auto-annotated genome sequences have been deposited in GenBank. For further information, see the J.Craig Venter Institute's <u>Marine Microbial Genome Sequencing Project</u>.

Publications and Data

Project: Moore Marine Microbial Sequencing

This project has no papers.

Marine Organisms

1 - 20 of 168 Advanced Sort 🚽 prev 20 📔 goto 🍸 | next 20 🕨 Release Date Organism Investigator Status Acaryochloris sp. CCMEE 5410 🗖 Michelle Wood Not Yet Received Aciduliprofundum boonei T469 🗖 Anna-Louise Reysenbach 03/07/06 Released to Genbank AEGEAN 169 clade (SAR11-like) HIMB37 🗖 Michael Rappé Not Yet Received Christopher Bolch 04/07/06 Released to Genbank Alcanivorax sp. DG881 🗖 Algoriphagus sp. PR1 🗖 Nicole King 12/14/05 Released to Genbank alpha proteobacterium BAL199 🗖 Åke Hagström 04/06/06 Released to Genbank alpha proteobacterium HTCC2255 🗖 Stephen J. Giovannoni, Jang-Cheon Cho Released to Genbank 08/09/05 Alteromonadales TW-7 🗖 Faroog Azam 11/22/05 Released to Genbank Francisco Rodriguez-Valera Alteromonas macleodii 'Deep ecotype' 🗖 11/19/04 Released to Genbank Aurantimonas sp. SI85-9A1 🗖 Bradley Tebo 08/17/04 Released to Genbank Bacillus sp. B14905 🗖 11/15/05 Rob Edwards Released to Genbank Bacillus sp. NRRL B-14911 🗖 Janet Siefert 08/24/04 Released to Genbank Bacillus sp. SG-1 🗖 Bradley Tebo 11/29/05 Released to Genbank Beggiatoa sp. 'Orange Guaymas' 🗖 Andreas Teske 10/10/07 16S QC (PCR -- Analysis) beta proteobacterium KB13 🗖 Michael Rappe 02/20/07 Delivered to Researcher Blastopirellula marina DSM 3645 🗖 Rudolf Amann 09/01/04 Released to Genbank Brevundimonas sp. BAL3 🗖 Åke Hagström 04/06/06 Released to Genbank Caldanaerobacter DSM 12653 🗖 Juan Gonzalez, Tatyana Sokolova Delivered to Researcher 05/17/06

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Calothrix sp. SC01 🗖

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Jonathan Zehr

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Not Yet Received

12/13/05 Released to Genbank

Camera >> research



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All Metagenomic ORF Peptides (P)	4,439,529,911	peptide
All Metagenomic ORFs (N)	13,318,589,733	nucleotide
All Metagenomic Sequence Reads (N)	8,393,828,709	nucleotide
FLAS: All Metagenomic Sequence Reads (N)	2,380,900	nucleotide
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GOS: All ORFs (N)	13,186,260,663	nucleotide
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GOS: GS000b Reads from 0.22-0.8 Sargasso Sea Stations 11 and 13 (N)	321,026,307	nucleotide
GOS: GS000c Reads from 0.22-0.8 Sargasso Sea Station 3 (N)	371,688,861	nucleotide
GOS: GS000d Reads from 0.22-0.8 Sargasso Sea Station 13 (N)	335,939,509	nucleotide
GOS: GS001 a Reads from 3.0-20.0 Sargasso Sea Hydrostation S (N)	143,316,448	nucleotide
GOS: GS001b Reads from 0.8-3.0 Sargasso Sea Hydrostation S (N)	90,955,161	nucleotide
GOS: GS001c Reads from 0.1-0.8 Sargasso Sea Hydrostation S (N)	92,688,958	nucleotide
GOS: GS002 Reads from 0.1-0.8 Gulf of Maine USA (N)	128,761,768	nucleotide

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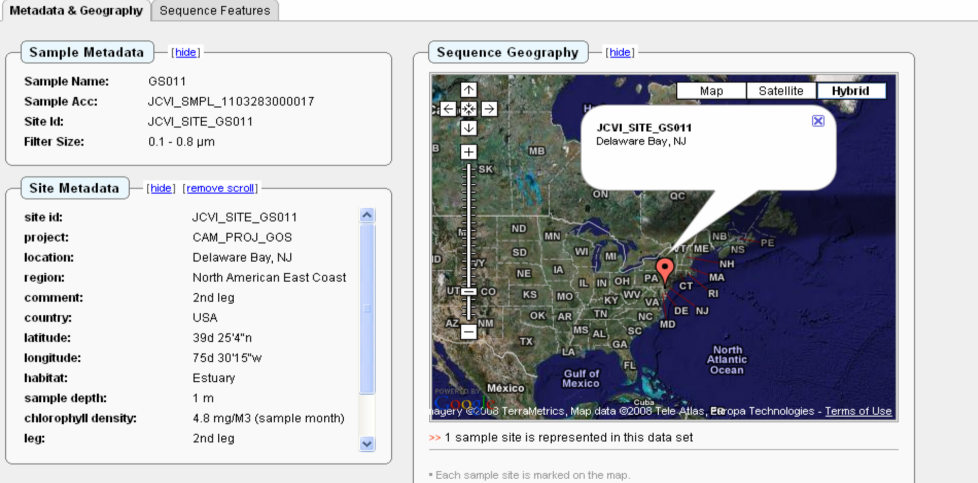
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x	NasAvsGOS [edit]	02/07/08 11:51 AM	completed	blastp	<u>1,019</u>	GOS: All ORF Peptides (P)	Results / Job 🍸 / Export 🍸
x	NasAvsGOS [edit]	02/07/08 11:36 AM	completed	blastp	<u>4,060</u>	GOS: All ORF Peptides (P)	Results / Job Y / Export Y
x	sdfs [<u>edit]</u>	01/11/08 03:14 PM	completed	blastn	<u>25</u>	GOS: All ncRNAs (N)	Results / Job 🌱 / Export 🌱
x	testg (<u>edit</u>)	01/07/08 02:15 PM	completed	blastp	<u>1,000</u>	All Metagenomic ORF Peptides (P)	Results / Job 🔨 / Export 🍸
×	testf [<u>edit]</u>	01/07/08 02:08 PM	completed	blastp	<u>1,000</u>	GOS: Combined Assembly ORF Peptides (P)	Results / Job 🌱 / Export 🌱
×	TESTaaa (<u>edit</u>)	12/10/07 11:36 AM	completed	tblastn	<u>2,615</u>	All Metagenomic ORFs (N)	Results / Job Y / Export Y
×	test [<u>edit]</u>	11/06/07 07:41 AM	completed	blastp	<u>318</u>	GOS: All ORF Peptides (P)	Results / Job 🍸 / Export 🍸
x	ntsars02vsnraa (<u>edit</u>)	10/18/07 11:17 AM	completed	blastp	<u>3,180</u>	CAMERA's Non-Identical Peptide Sequences (P)	Results / Job 🔽 / Export 🍸
×	ntsars1vsnraa (<u>edit</u>)	10/18/07 10:46 AM	completed	blastp	<u>2,447</u>	CAMERA's Non-Identical Peptide Seque Re-run this Show para	iob w/ new parameters / Export V
×	ttt [<u>edit]</u>	08/01/07 01:25 PM	completed	blastp	<u>25</u>	Microbial draft genome ORF peptide sequences (F	
×	ess (<u>edit</u>)	07/20/07 09:58 AM	completed	tblastn	<u>230</u>	GOS: All Metagenomic Sequence Reads (N)	Results / Job V / Export V
×	atz (<u>edit</u>)	07/17/07 05:05 PM	completed	blastp	<u>318</u>	GOS: All ORF Peptides (P)	Matching Query Sequences as FASTA Matching Subject Sequences as FASTA
×	atzAA (<u>edit</u>)	07/17/07 04:57 PM	completed	blastp	<u>409</u>	COO: All ODE Doutidoo (D)	BLAST Results with Metadata as CSV BLAST Results as NCBI XML
×	atzD (<u>edit</u>)	07/17/07 04:47 PM	completed	blastp	<u>11</u>	GOS: All ORF Peptides (P)	Results / Job Y / Export Y
×	atzC (<u>edit</u>)	07/17/07 04:45 PM	completed	blastp	<u>200</u>	GOS: All ORF Peptides (P)	Results / Job 🍸 / Export 🍸
×	atzB (<u>edit</u>)	07/17/07 04:44 PM	completed	blastp	<u>200</u>	GOS: All ORF Peptides (P)	Results / Job 🔨 / Export 🍸

< HC M	E	ABO	DUT CA	MERA METAGENOMICS	RESEARCH	NEWS		EVENTS	DISCUSSION FORUMS
<mark> ∳</mark> ≱ F	Research > Job	s > Job Result	ts > Job	Details	Research Home	Clusters 🥆	Jobs 🔻	Sets 🔻 🛛 Viewers 🔻	Data 🔻
Jo	b Summar	- [<u>show</u>]] [< bac	k to job results]					
_ Ma	atching Sec	uences –	- [<u>hide</u>]						Sequence Geography – [hide] –
									TX CA
1 - 1	10 of 82		1					ev 10 goto 🔻 next 10)	↑ Map Satellite Hybrid
	<u>Eval</u>	Score ▼		Query	<u>Subject</u>	Sample(Location(s)	← ☆ → Gulf of Ocean
	5.142E-3	-			JCVI_PEP_11051327			are Bay, NJ	México México Cuba
	2.097E-2	-		USER_PEP_1202414661201	JCVI_PEP_11051569			are Bay, NJ	+ PR
	2.482E-1	-		USER_PEP_1202414661201	JCVI_PEP_11051313			ort Harbor, RI	un 🕂
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	6.119E-1	-		USER_PEP_1202414661201	1			'Panama	Venezuela
				AA <mark>NL</mark> SRYRLFGN <mark>I</mark> QNSIQ <mark>E</mark> FII		7 <u>8257</u> GS006		Fundy, Nova Scotia	
] Sbjct:	77 <mark>S</mark> GDI	DWIAV	AQ <mark>SM</mark> RAYLRFSL <mark>L</mark> GGVIA <mark>K</mark> QII		05957 GS031		ling, Fernandina Island	Colombia Suri
	Query	301 SPF9	SG <mark>IM</mark> P	AIASAQLTGITEG <mark>NR</mark> NNGFCT <mark>S</mark>	DEFIGEVIEVE 340	34839 GS004		le Halifax, Nova Scotia	Ecuador
		117 STD	GD <mark>VL</mark> Q	P-PLLL <mark>LSGL</mark> GEV <mark>SR</mark> IGEVIL <mark>)</mark>		67667 GS033		Cormorant, Hypersalin	
			TONDT	EDTUDID & CTUDECOTCINC &		3 <u>9069</u> GS033	Punta	Cormorant, Hypersalin	
Se				EPTNPIDAG <mark>I</mark> WRFC <mark>QT</mark> CTKCAI LFDKPIDFG <mark>L</mark> QAFC <mark>ES</mark> CNKCAI		Advanced Sort		Export 💙	Perú Ro Br
1 - 1							a pr	goto -	ery Sequences as FASTA
= Ho				NIYGKADIVHSPGR <mark>K</mark> Q <mark>Y</mark> WTNA					bject Sequences as FASTA
= BL	L Shact :	196 P <mark>K</mark> LI	MFN	GY <mark>E</mark> I <mark>W</mark> KSDS(2KCTT <mark>Y</mark> RLTNE 221				ults as NCBI XML Paraget
	Query:	420 NG	-CGAC	MATCT <mark>F</mark> NTNTAPIH <mark>E</mark> LVRATL	ATTPTLNTFLW 457			DENOTITO	Paragua
S				MKTCP <mark>WNLEGLFKE</mark> KPFRWLA <mark>S</mark>					Chile 4
–	·		VEROU	of WERVER					
Sec				-GLNEDK <mark>E</mark> EWWDLSLPVYGFDS GGLNEIK <mark>K</mark> WWWDLEL		99/276(35%)		ap data @2008 Tele Atlas, MapLink/Tele Atlas, Europa Technologies - <u>Terms of Use</u>
Sec	-	202 ALD	DICLON	OOLMEIN <mark>N</mark> WWEELLE	KDD <u>001</u> K 271	46/276(16%)		>> 29 sample sites are represented in this data set
Alig		497 <u>K</u> 4'				ps: 8			
Cle] Sbjct:	292 <u>K</u> 2'	92			aps: 12			 Each sample site is marked on the map.
)				 Click a site marker for more information
Que	ery: 218	GKAYVFEDV	V <mark>D</mark> VGYE	GATTFV <mark>L</mark> PANK <mark>Q</mark> L <mark>Y</mark> EFCFTVP	MSKDMFRTANESO	IMYAANL <mark>S</mark> RYRI	LFGNIQ <mark>N</mark> S	SIQE <mark>FI</mark> RSLGY 294	Drag the map with your mouse, or double-click to recenter
	·	GK FEDV	7 <mark>+</mark> YE	E T V+P N + + FTV	MS+++ + +	AA <mark>+</mark> Y	I +	Q F+ LGY	
Sb	jct: 28	GKRLDFEDV	EKPYE	EETRRV <mark>I</mark> PNNA <mark>R</mark> - <mark>W</mark> VIVFTVQ	MSEELLKRRVGKTPTP	FSAAAAG <mark>N</mark> AYAI	RSRVIF <mark>D</mark> I	DTQIF <mark>L</mark> HCLGY 106	
0114	=rv 295 '	ICVGV-ASP	FSGIN	IPAIASAQ <mark>LTGI</mark> TEG <mark>NR</mark> NNGFC	TSPRVGP <mark>VT</mark> GVRS <mark>T V</mark> T	DMPIFPTNPID	AGIMEEC	TCTKCADACP 373	
Que	JIY. 200			A A + G+ E +R N					
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0		MATONDUE	DTUD	DULYCVADIU		CCLCWL TCTT	TNTADT	UCT UDATTATT AAO	
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Sb	jct: 184 !			KGPWNNPGHKTWYED					
~		TT NTET IT	ADVET	CVCT NEDVEENNDI <mark>ST</mark> DVVCE	D 405				

Query: 450 PTLNTFLWQADKFFGYGLNEDKEEWWDLSLPVYGFD 485

Peptide Det	ail — [hide] [< back to job details] —	
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Def Line:	JCVI_PEP_1105132741961 /read_id=JCVI_READ_1092351470431 /begin=0 /end=900 /orientation=1 /5_prime_stop=0 /3_prime_stop=TAG /orf_id=JCVI_ORF_1105132741960 /ttable=11 /length=300 /ergatis_id=7720 /read_defline="/library_id=JCVI_LIB_GS-11-01-01-1P3-1P8KB /template_id=JCVI_TMPL_1061001659921 /sequencing_direction=forward /sample_id=JCVI_SMPL_1103283000017 /clr_range_begin=194 /clr_range_end=1073 /full_length=1124"	Intellectual Property Notice
Seq Length:	300	
Seq Type:	AA	
Sequence	ERSGTSHVAFLELDEVNKKFIYSVDNDGKRLDFEDVEKPYEEETRRVIPNNARWVIVFTVQMSEELLKRRVGKTPTPFSAAAA YYGLMAYWFNGLGIAPALGIMAGLGELSRLNRMITPEYGPMVRVFRMITNLPLAPNKPIDAGIMRFCKTCKKCATECPSGTLS EDSPKCMAYWLKATASCATCFSVCPFSKKDKSFMHHFVEATIAKTPIFNSLFTKMDTLLGYDIPRDPESWWGTNMPPYGIDGI	GLDTEPSWEVKGPWNNPGHKTWY



Click a site marker for more information on Site Metadata panel

411	С	D	E		F	G		Н	1	J	K	L	М	N	0	Р	Q
	ilter Size (Lat/Long	Smpl Po		Geo Pos	Nation	Time	Date	Habitat	Smpl Dept				Chlorophyll Densi A
	.1 - 0.8							North American East			11/16/2003		1 m	12 m	9.4 C	26.5 ppt	2.2 mg/M3 (sample1
	.1 - 0.8					79 Gulf of Pa		Eastern Tropical Pac		4:48 PM			1.6 m	76 m	27.6 C	30.7 ppt	0.50 mg/M3 (sampl0
	.1 - 0.8			-				North American East		5:25 AM			2 m	142 m	17.3 C	28.3 ppt	0.4 mg/M3 (sample0
	.1 - 0.8							North American East		11:50 AM			1 m	119 m	11.7 C	29.9 ppt	1.4 mg/M3 (sample1
_	.1 - 0.8			_		79 Gulf of Pa		Eastern Tropical Pac		4:48 PM			1.6 m	76 m	27.6 C	30.7 ppt	0.50 mg/M3 (sampl0
_	.1 - 0.8							North American East			11/16/2003		1 m	12 m	9.4 C	26.5 ppt	2.2 mg/M3 (sample 1
-	.1 - 0.8			_				North American East			11/16/2003		1 m	12 m	9.4 C	26.5 ppt	2.2 mg/M3 (sample 1
_	.1 - 0.8					n ; Cape Mag		North American East			11/18/2003		1 m	10 m	12 C	31 ppt	2.0 mg/M3 (sample 2
_	.1 - 0.8							North American East			12/19/2003		2.1 m	20 m	9.3 C		3.0 mg/M3 (sample 2
_	.1 - 0.8			_				North American East			12/20/2003		1 m	31 m	18.6 C	-	1.70 mg/M3 (sampl1
and the second division of the second divisio	.1 - 0.8								Ecuador	5:06 PM			2.1 m	35 m	27.5 C	20.7	0.36 mg/M3 (sampl0
and the second se	.1 - 0.8							Eastern Tropical Pac		4:48 PM			1.6 m	76 m	27.6 C	30.7 ppt	0.50 mg/M3 (sampl0
_	.1 - 0.8					79 Gulf of Pa		Eastern Tropical Pac		4:48 PM			1.6 m	76 m	27.6 C	30.7 ppt	0.50 mg/M3 (sampl0
	1-0.8							North American East			11/16/2003		1 m	12 m	9.4 C	26.5 ppt	2.2 mg/M3 (sample 1
	.1 - 0.8							North American East	-		11/16/2003		1 m	12 m	9.4 C	26.5 ppt	2.2 mg/M3 (sample 1
	.1 - 0.8							Caribbean Sea	Panama	9:03 AM			1.7 m	3336 m	27.7 C	35.4 ppt	0.23 mg/M3 (sampl0
_	.1 - 0.8							Galapagos Islands	Ecuador	11:41 AM			2.2 m	2.3 m	25.5 C	34.9 ppt	0.40 mg/M3 (sampl0
	.1 - 0.8							North American East			12/19/2003		2.1 m	20 m	9.3 C	-	3.0 mg/M3 (sample 2
_	.1 - 0.8			-				Galapagos Islands	Ecuador	3:47 PM			2 m	156 m		045	0.35 mg/M3 (sampl0
	.1 - 0.8			_		; 9 Wolf Islar		Galapagos Islands	Ecuador	4:44 PM			1.7 m	71 m	21.8 C	34.5 ppt	0.28 mg/M3 (sampl0
_	.1 - 0.8					n; Gulf of M		Caribbean Sea	USA	2:15 PM		Coastal Sea	2 m	3333 m	26.4 C	35.8 ppt	0.16 mg/M3 (sampl0
_	.1 - 0.8					n; Gulf of M		Caribbean Sea	USA	2:15 PM		Coastal Sea	2 m	3333 m	26.4 C	35.8 ppt	0.16 mg/M3 (sampl0
	.1 - 0.8							Galapagos Islands	Ecuador	2:43 PM		Coastal upwe		19.6 m	18.6 C		0.35 mg/M3 (sampl0
	.1 - 0.8							Galapagos Islands	Ecuador	2:43 PM		Coastal upwe		19.6 m	18.6 C		0.35 mg/M3 (sampl0
	.1 - 0.8							Galapagos Islands	Ecuador	2:43 PM		Coastal upwe		19.6 m	18.6 C		0.35 mg/M3 (sampl0
	.1 - 0.8					91 Upwelling	, Fernar	Galapagos Islands	Ecuador	2:43 PM		Coastal upwe	lli 12 m	19.6 m	18.6 C	-	0.35 mg/M3 (sampl0
the second se	.1 - 0.8			_GS031										100	18.6 C	-	0.35 mg/M3 (sampl0
	.1 - 0.8			_GS031										-	18.6 C		0.35 mg/M3 (sampl0
Concession in the local division in the loca	.1 - 0.8			_GS031										-	18.6 C		0.35 mg/M3 (sampl0
	.1 - 0.8			_GS031										-	18.6 C		0.35 mg/M3 (sampl0
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_	.1 - 0.8			_GS031											18.6 C	<u></u>	0.35 mg/M3 (sampl0
	.1 - 0.8			_GS031				Warm See	р						18.6 C	<u></u>	0.35 mg/M3 (sampl0
	.1 - 0.8			_GS031				5%						-	18.6 C		0.35 mg/M3 (sampl0
	.1 - 0.8			_GS031				1		Coastal		_			18.6 C	_	0.35 mg/M3 (sampl0
	.1 - 0.8			_GS031			Open	Ocean /		30%			Coastal		18.6 C	-	0.35 mg/M3 (sampl0
	.1 - 0.8			_GS031				5% /					Coastal upw		18.6 C		0.35 mg/M3 (sampl0
	.1 - 0.8			_GS031						1					18.6 C	-	0.35 mg/M3 (sampl0
	.1 - 0.8			_GS031									Estuary		18.6 C		0.35 mg/M3 (sampl0
	.1 - 0.8			_GS031									-lypersaline		18.6 C	24.2	0.35 mg/M3 (sampl0
	.1 - 0.8			_GS051				FI						. II-	27.3 C	34.2 ppt	
	.1 - 0.8			_GS051			Hype	ersaline					Open Ocean		27.3 C	34.2 ppt	C
and the second s	.1 - 0.8			_GS005				2%					Narm Seep	- II-	15 C	30.2 ppt	6 mg/M3 (sample n6
	.1 - 0.8			_GS005			1	210	- 0	oactal up	wolling				15 C	30.2 ppt	6 mg/M3 (sample n6
	.1 - 0.8			_GS011				Estuary /	C	pastal up	-				11 C		4.8 mg/M3 (sample 9
	.1 - 0.8			_GS011				6%		22%					11 C		4.8 mg/M3 (sample 9
	.1 - 0.8			_GS006				070							11.2 C	8	2.8 mg/M3 (sample 1
	.1 - 0.8			_GS006											11.2 C	2	2.8 mg/M3 (sample 1
	.1 - 0.8			_GS006											11.2 C	00.4	2.8 mg/M3 (sample1
51 0	.1 - 0.8	309 J	CVI_SITE	_GS033	10 13 42										37.6 C	63.4 ppt	

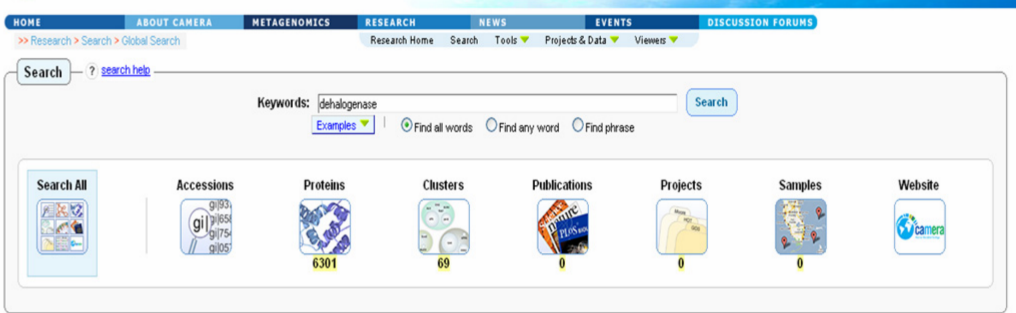
Camera >> research



Recent Searches [hide]
 lipase no matches
 glycerol kinase (about 4,239
matches)
Cellulase (about 3,163 matches)
ATP synthase epsilon (about
2,856
matches)
dehalogenase (about 4,963
matches)

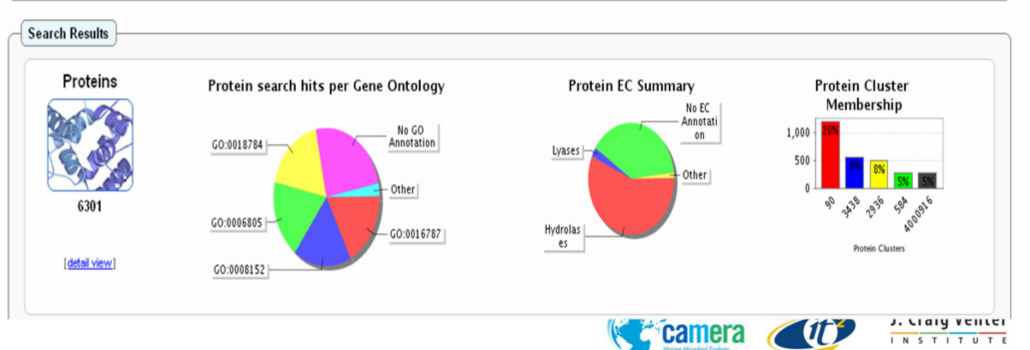
Recent BLAST Results [hide]							
Job Name	<u>Submit Date</u>	<u>Status</u>	# Hits	Subject Sequences			
test11 02/27/07 02:09 PM		completed	<u>2</u>	GOS: Assembled Sequences (N)			
tst10	02/27/07 02:09 PM	completed	0	GOS: Combined Assembly ORF Peptides (P)			
test09	02/27/07 02:08 PM	completed	<u>1</u>	GOS: All Metagenomic Sequence Reads (N)			
test06	02/27/07 02:05 PM	completed	<u>50</u>	All Metagenomic Sequence Reads (N)			

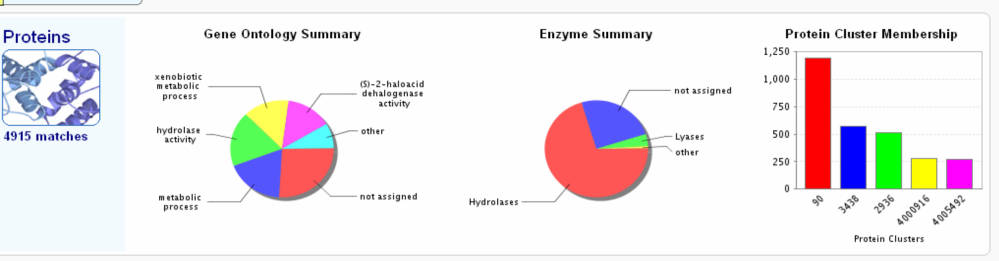




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All Matching Proteins

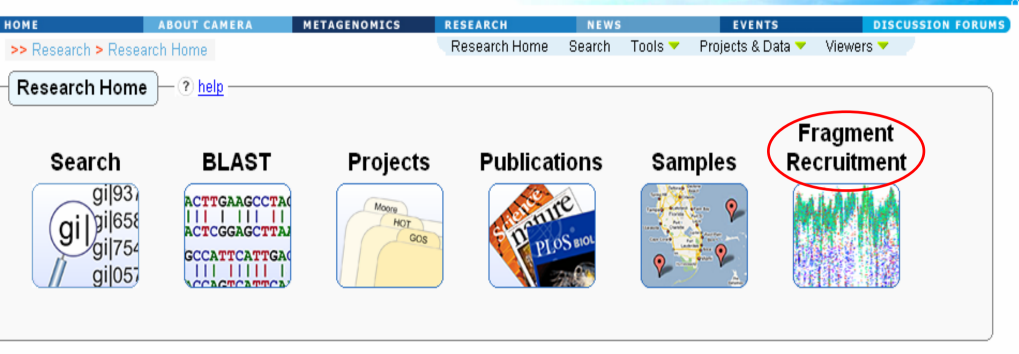
1 - 20 of 4915		Adva	inced Sort	⊲ prev 20 I g	oto 🔻 🛙 🗖	ext 20 🕨
Accession	External Acc	<u>NCBI GI</u>	Headline	DefLine	Length	Final
JCVI_PEP_1096665714389			Accession: CAM_CNAM_102139 c	JCVI_PEP_1096665714389 /source_dna_id=JCVI_ORF	147	CAM_C
JCVI_PEP_1096666313063			Accession: CAM_CNAM_102139 c	JCVI_PEP_1096666313063 /source_dna_id=JCVI_ORF	147	CAM_C
JCVI_PEP_1096668602421			Accession: CAM_CNAM_102139 c	JCVI_PEP_1096668602421 /source_dna_id=JCVI_ORF	155	CAM_C
JCVI_PEP_1096673421409			Accession: CAM_CNAM_102139 c	JCVI_PEP_1096673421409/source_dna_id=JCVI_ORF	148	CAM_C
JCVI_PEP_1096676798831			Accession: CAM_CNAM_102139 c	JCVI_PEP_1096676798831 /source_dna_id=JCVI_ORF	90	CAM_C
JCVI_PEP_1096680464767			Accession: CAM_CNAM_102139 c	JCVI_PEP_1096680464767 /source_dna_id=JCVI_ORF	147	CAM_C
JCVI_PEP_1096680522357			Accession: CAM_CNAM_102139 c	JCVI_PEP_1096680522357 /source_dna_id=JCVI_ORF	160	CAM_C
JCVI_PEP_1096691663293			Accession: CAM_CNAM_102139 c	JCVI_PEP_1096691663293 /source_dna_id=JCVI_ORF	146	CAM_C
JCVI_PEP_1096692584989			Accession: CAM_CNAM_102139 c	JCVI_PEP_1096692584989 /source_dna_id=JCVI_ORF	100	CAM_C
JCVI_PEP_1096697966445			Accession: CAM_CNAM_102139 c	JCVI_PEP_1096697966445 /source_dna_id=JCVI_ORF	151	CAM_C
JCVI_PEP_1105173466959			Accession: CAM_CNAM_102139 c	JCVI_PEP_1105173466959 /read_id=HOT_READ_857589	127	CAM_C
NCBI_PEP_37702690	AAR00932.1	37702690	Accession: CAM_CNAM_102139 c	>NCBI_PEP_37702690/ACCESSION=AAR00932.1/GI=3770	150	CAM_C
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NCBI_PEP_111022131	YP_705103.1	111022131	Accession: CAM_CNAM_104264 c	NCBI_PEP_111022131 /offset=0 /translation_table	276	CAM_C
JCVI_PEP_1096665787299			Accession: CAM_CNAM_104669 c	JCVI_PEP_1096665787299 /source_dna_id=JCVI_ORF	74	CAM_C
JCVI_PEP_1096670067379			Accession: CAM_CNAM_104669 c	JCVI_PEP_1096670067379 /source_dna_id=JCVI_ORF	71	CAM_C
JCVI_PEP_1096673037697			Accession: CAM_CNAM_104669 c	JCVI_PEP_1096673037697 /source_dna_id=JCVI_ORF	185	CAM_C
JCVI_PEP_1096674592091			Accession: CAM_CNAM_104669 c	JCVI_PEP_1096674592091 /source_dna_id=JCVI_ORF	175	CAM_C
JCVI_PEP_1096676803799			Accession: CAM_CNAM_104669 c	JCVI_PEP_1096676803799 /source_dna_id=JCVI_ORF	178	CAM_C
JCVI_PEP_1096688757069			Accession: CAM_CNAM_104669 c	JCVI_PEP_1096688757069 /source_dna_id=JCVI_ORF	121	CAM_C
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					Exp	ort 🔽
1 - 20 of 4915		Sho	w: <u>10</u> 20 <u>50</u>	Protein search	h results as	

Protein sequences as FASTA

CAMERA Accession: JCVL_PEP_1105173466959 Camera Accession: JCVL_PEP_1105173466959 Sample: HF10_10-07-02 - HOT station ALOHA, 10 m ORF: JCVL_ORF_1105173466958 Source DNA: HOT READ_85756924 Translation begin: 64 Translation table: 118 S'Stop: TAA 3'Stop: TAA JCVL_PEP_1105173466959 /read_id=HOT_READ_85758924 /begin=64 /end=445 /orientation=1 //Sprime_stop=TAA /orT_id=JCV_ORF_1105173466959 /read_id=HOT_READ_85758924 /begin=64 /end=445 /orientation=1 //Sprime_stop=TAA /orT_id=JCV_ORF_1105173466959 /read_id=HOT_READ_85758924 /begin=64 /end=445 /orientation=1 //Sprime_stop=TAA /orT_id=JCV_ORF_1105173466959 /read_id=HOT_READ_85758924 /begin=64 /end=445 /orientation=1 //Sprime_stop=TAA/orT_id=JCV_ORF_1105173466959 /read_id=HOT_READ_85758924 /begin=64 /end=445 /orientation=1 //Sprime_stop=Tathore JCVL_PEP_11005173466959		Peptide Details	[hide] [◄ back to search details] (?) help	
Bindby Type: Protein Sample: HF10_10-07-02 - HOT station ALOHA, 10 m ORF: JCVI_ORF_1105173466958 Source DNA: HOT_READ_85758924 Translation begin: 64 Translation end: 445 Orientation: forward Translation table: 11 11 5' Stop: TAA 3' Stop: TAA Defline: JCVI_PEP_1105173466959 /read_id=HOT_READ_85758924 /begin=64 /end=445 /orientation=1 /fs_prime_stop=TAA/3_prime_stop=TAA/3_prime_stop=TAA/ort_id=JCVI_ORF_1105173466958 /table=11 /length=127 /regatis_id=7702 /sample_id=Joi_SMPL_HF10_10-07-02 /sample_name=HF10_10-07-02 /simmes_o_i=1************************************	n	replice betails		
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ORF: JCVI_ORF_1105173466958 Source DNA: HOT_READ_85758924 Translation begin: 64 Translation end: 445 Orientation: forward Translation table: 11 % 5' Stop: TAA 3' Stop: TAA Defline: JCVI_PEP_1105173466959 /read_id=HOT_READ_85758924 /begin=64 /end=445 /orientation=1 /5_prime_stop=TAA /a_prime_stop=TAA /a_prime_stop=TAA /or_id=JCVI_ORF_1105173466958 /ttable=11 /length=127 /ergatis_id=7702 /sample_id=JGI_SMPL_HF10_10-07-02 /sample_name=HF10_10-07-02 /number_of_sites=1 /site_id_1=HAVXAIL_SITE_HOT_10 /location_1="Havaii Ocean Time-series station ALOHA, 10 m' /region_1="North Pacific Subtropical Gyre' /country_1=USA/site_depth_1="1" nm0/kg'' /dissolved_organic_carbon_1="78 um0/kg'' /dissolved_inorganic_phosphate_1="41 nm0/kg'' /dissolved_organic_carbon_1="78 um0/kg'' /dissolved_organic_carbon_1="38 put" /hemperature_1="26.4 C" /water_depth_1="4750 m' /core_cluster_id=CAM_CRCL_4671 /final_cluster_id=CAM_CL_10705 Seq Length: 127 Seq Length: 127 Sequence IANDITINTHSKFTGANTFFAQVIFQKNPKNSHFMGGKLVKTKEIFLNGQIRAGRTSKVKKQLIIGLRKILIKNTKLQSDFIWVYLEDLLPHOMIEYGEVLPKSGQ		Entity Type:	Protein	Download Protein sequence
Source DNA: HOT_READ_86758924 Translation begin: 64 Translation end: 445 Orientation: forward Translation table: 11 % Stop: TAA 3' Stop: TAA 3' Stop: TAA Defline: JCVI_PEP_1105173466959 /read_id=HOT_READ_85758924 /begin=64 /end=445 /orientation=1 /stop: TAA 3' Stop: TAA JCVI_PEP_1005173466959 /read_id=HOT_READ_85758924 /begin=64 /end=445 /orientation=1 /stop:site_id=702 /sample/id=JGI_8MPL_HF10_10-07-02 /sample_name=HF10_10-07-02 /sumble_of_sites=11 /site_id_1==HAWAIL_SITE_HOT_10 /location_1=="Hawail Ocean Time-series station ALOHA, 10 m" /region_1=="North Pacific SubtropHI_density_1==*0.08 ug/kg" /dissolved_inorganic_carbon_1=**1,967.60 umol/kg" /dissolved_inorganic_phosphate_1==*11 nmol/kg" /salinity_1=**3.08 psi/ nemperature_1=**26.4 C" water_depth_1=**4750 m* /core_cluster_id=CAM_CRCL_4671 /final_cluster_id=CAM_CL_10705 Seq Length: 127 Seq Length: 127 Sequence IANDITNTHSKFTGANTFFAQVIFOKNPKNSHFMGGKLVKTKEIFLINGQIRAGRTSKVKKQLILGLRKILIKNTKLQSDFIWVYLEDLLPHOMIEYGEVLPKSGQ		Sample:	HF10_10-07-02 - HOT station ALOHA, 10 m	
Translation begin: 64 Translation end: 445 Orientation: forward Translation table: 11 % 5' Stop: TAA 3' Stop: TAA Defline: JCVI_PEP_1105173466959 /read_id=HOT_READ_85758924 /begin=64 /end=445 /orientation=1 /5_ stop: TAA 0efline: JCVI_PEP_1105173466959 /read_id=HOT_READ_85758924 /begin=64 /end=445 /orientation=1 /ste_rgatis_id=7702 /sample_id=JGI_8MPL_HF10_10-07-02 /sample_inme=HF10_110-07-02 /number_of_sites=1 /site_id_1=HAWAIL_SITE_HOT_10 /location_1="Hawaii Ocean Time-series station ALOHA, 10 m" /region_1="North Pacific Subtropical Gyre" /country_1=USA /site_oth_1=="10"/sitesolved_organic_carbon_1="7.8 umol/kg" /dissolved_inorganic_bhosphate_1="41"/sitesolved_organic_carbon_1="7.8 umol/kg" /dissolved_inorganic_bhosphate_1="41"/sitesolved_organic_carbon_1="7.8 umol/kg" /missolved_mol/kg" //dissolved_inorganic_bhosphate_1="41"/sitesolved_organic_carbon_1="7.8 umol/kg" /water_depth_1="4750 m" /core_cluster_id=CAM_CRCL_4671 /final_cluster_id=CAM_CL_10705 Seq Length: 127 Sequence IANDITNTHSKFTGANTFFAQVIFQKNPKNSHFMGGKLVKTKEIFLNGQIRAGRTSKVKKQLILGLRKLILKNTKLQSDFIWVYLEDLLPHQMIEYGEVLPKSGQ		ORF:	JCVI_ORF_1105173466958	
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Translation table: 11 % 5' Stop: TAA 3' Stop: TAA JCVI_PEP_1105173466959 /read_id=HOT_READ_85758924 /begin=64 /end=445 /orientation=1 /stop: /5_prime_stop=TAA /3_prime_stop=TAA /orf_id=JCVI_ORF_1105173466958 /table=11 /length=127 /ergatis_id=7702 /sample_id=JGI_SMPL_HF10_10-07-02 /sample_name=HF10_10-07-02 /sample_rof_sites=1 /site_id_1=HAWAII_SITE_HOT_10 /location_1="Hawaii Ocean Time-series station ALOHA, 10 m" /region_1="North Pacific Subtropical Gyre" /country_1=USA /site_depth_1="10 m" /biomass_1="7.21+/2.68 ug/kg" None /chlorophyli_density_1="0.08 ug/kg" /dissolved_organic_carbon_1="780 umol/kg" //site_id=1=HAWAII_SITE_HOT_10 /location_1="Hawaii Ocean Time-series station ALOHA, 10 m" /region_1="North Pacific Subtropical Gyre" /country_1=USA /site_depth_1="10 m" /biomass_1="7.21+/2.68 ug/kg" /dissolved_inorganic_carbon_1="North Pacific Subtropical Gyre" /country_1=USA /site_depth_1="10 m" /biomass_1="7.21+/2.68 ug/kg" /dissolved_inorganic_carbon_1="North Pacific Subtropical Gyre" /country_1=USA /site_depth_1="10 m" /biomass_1="7.21+/2.68 ug/kg" /dissolved_inorganic_earbon_1="North Pacific Subtropical Gyre" /country_1="204.6 mmol/kg" /dissolved_organic_carbon_1="780 umol/kg" /dissolved_inorganic_earbon_1="204.6 mmol/kg" /dissolved_organic_carbon_1="350.8 psu" /lemperature_1="26.4 C" //dissolved_inorganic_earbon_1="4750 m" /core_cluster_id=CAM_CRCL_4671 /final_cluster_id=CAM_CL_10705 //dissolved_inorganic_earbon_1="204.6 mmol/kg" //dissolved_inorganic_earbon_1="204.6 mmol/kg" //dissolved_inorganic_earbon_1="204.6 mmol/kg" //dissolved_inorgani		Translation end:	445	
5' Stop: TAA 3' Stop: TAA Defline: JCVI_PEP_1105173466959 /read_id=HOT_READ_85758924 /begin=64 /end=445 /orientation=1 /5_prime_stop=TAA /3_prime_stop=TAA /or_id=JCVU_ORF_1105173466958 /table=11 /length=127 /ergatis_id=7702 /sample_id=JGI_SMPL_HF10_10-07-02 /sample_name=HF10_10-07-02 /number_of_sites=1 /site_id_1=HAWAIL_SITE_HOT_10 /location_1="Hawaii Ocean Time-series station ALOHA, 10 m" /region_1="North Pacific Subtropical Gyre" /country_1=USA /site_depth_1="10 m" /biomass_1="7.21+/-2.68 ug/kg" /chlorophyl_density_1="204.6 nmol/kg" /dissolved_organic_carbon_1="196.760 umol/kg" /dissolved_inorganic_phosphate_1="41 nmol/kg" /dissolved_organic_carbon_1="78 umol/kg" /dissolved_morganic_phosphate_1="41 nmol/kg" //sialinity_1="35.08 psu" /temperature_1="26.4 C" /water_depth_1="4750 m" /core_cluster_id=CAM_CRCL_4671 /final_cluster_id=CAM_CL_10705 Seq Length: 127 Seq Type: AA Sequence IANDITNTHSKFTGANTFFAQVIFQKNPKNSHFMGGKLVKTKEIFLNGQIRAGRTSKVKKQLILGLRKILIKNTKLQSDFIWVYLEDLLPHQMIEYGEVLPKSGQ		Orientation:	forward	
3' Stop: TAA Intellectual Property Notice Defline: JCVI_PEP_1105173466959 /read_id=HOT_READ_85758924 /begin=64 /end=445 /orientation=1 /5 Intellectual Property Notice Defline: JCVI_PEP_1105173466959 /read_id=HOT_READ_85758924 /begin=64 /end=445 /orientation=1 /5 Intellectual Property Notice None None None None Stop: JCVI_PEP_1105173466959 /read_id=HOT_READ_85758924 /begin=64 /end=445 /orientation=1 /5 None /site_id_1=HAWAIL_SITE_HOT_10 /location_1="THAWAII OCT-02 /sample_name=HF10_10-07-02 /number_of_sites=1 /> None /site_id_1=HAWAIL_SITE_HOT_10 /location_1="Hawaii Ocean Time-series station ALOAA, 10 mm' /region_1="North Pacific Subtropical Gyre" /country_1=USA /site_depth_1="10 m" /biomass_1="7.21+/2.68 ug/kg" //sitesolved_inorganic_phosphate_1=*41 nmol/kg" /dissolved_organic_carbon_1="78 umol/kg" None Seq Length: 127 Seq Length: 127 Seq Type: A A Sequence IANDITNTHSKFTGANTFFAQVIFQKNPKNSHFMGGKLVKTKEIFLNGQIRAGRTSKVKKQLILGLRKILIKNTKLQSDFIWVYLEDLLPHQMIEYGEVLPKSGQ		Translation table:	<u>11</u>	
Defline: JCVI_PEP_1105173466959 /read_id=HOT_READ_85758924 /begin=64 /end=445 /orientation=1 /5_prime_stop=TAA /3_prime_stop=TAA /or_id=JCVI_ORF_1105173466958 /ttable=11 /length=127 /ergatis_id=7702 /sample_id=JGI_SMPL_HF10_10-07-02 /sample_name=HF10_10-07-02 /number_of_sites=1 /site_id_1=HAWAII_SITE_HOT_10 /location_1="Hawaii Ocean Time-series station ALOHA, 10 m" /region_1="North Pacific Subtropical Gyre" /country_1=USA /site_depth_1="10 m" /biomass_1="7.21+/-2.68 ug/kg" /chlorophyll_density_1="0.08 ug/kg" /dissolved_inorganic_carbon_1="1,967.60 umol/kg" /dissolved_oxygen_1="204.6 nmol/kg" /nitrate+nitrite_1="1 nmol/kg" /salinity_1="35.08 psu" /temperature_1="26.4 C" /water_depth_1="4750 m" /core_cluster_id=CAM_CRCL_4671 /final_cluster_id=CAM_CL_10705 Seq Length: 127 Seq Type: AA Sequence IANDITNTHSKFTGANTFFAQVIFQKNPKNSHFMGGKLVKTKEIFLNGQIRAGRTSKVKKQLILGLRKILIKNTKLQSDFIWVYLEDLLPHQMIEYGEVLPKSGQ		5' Stop:	TAA	
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Seq Type: AA Sequence IANDITNTHSKFTGANTFFAQVIFQKNPKNSHFMGGKLVKTKEIFLNGQIRAGRTSKVKKQLILGLRKILIKNTKLQSDFIWVYLEDLLPHQMIEYGEVLPKSGQ		Defline:	<pre>/5_prime_stop=TAA /3_prime_stop=TAA /orf_id=JCVI_ORF_1105173466958 /ttable=11 /length=127 /ergatis_id=7702 /sample_id=JGI_SMPL_HF10_10-07-02 /sample_name=HF10_10-07-02 /number_of_sites=1 /site_id_1=HAVVAIL_SITE_HOT_10 /location_1="Hawaii Ocean Time-series station ALOHA, 10 m" /region_1="North Pacific Subtropical Gyre" /country_1=USA /site_depth_1="10 m" /biomass_1="7.21+/-2.68 ug/kg" /chlorophyll_density_1="0.08 ug/kg" /dissolved_inorganic_carbon_1="1,967.60 umol/kg" /dissolved_inorganic_phosphate_1="41 nmol/kg" /dissolved_organic_carbon_1="78 umol/kg" /dissolved_oxygen_1="204.6 nmol/kg" /nitrate+nitrite_1="1 nmol/kg" /salinity_1="35.08 psu" /temperature_1="26.4 C"</pre>	None
Sequence IANDITNTHSKFTGANTFFAQVIFQKNPKNSHFMGGKLVKTKEIFLNGQIRAGRTSKVKKQLILGLRKILIKNTKLQSDFIWVYLEDLLPHQMIEYGEVLPKSGQ		Seq Length:	127	
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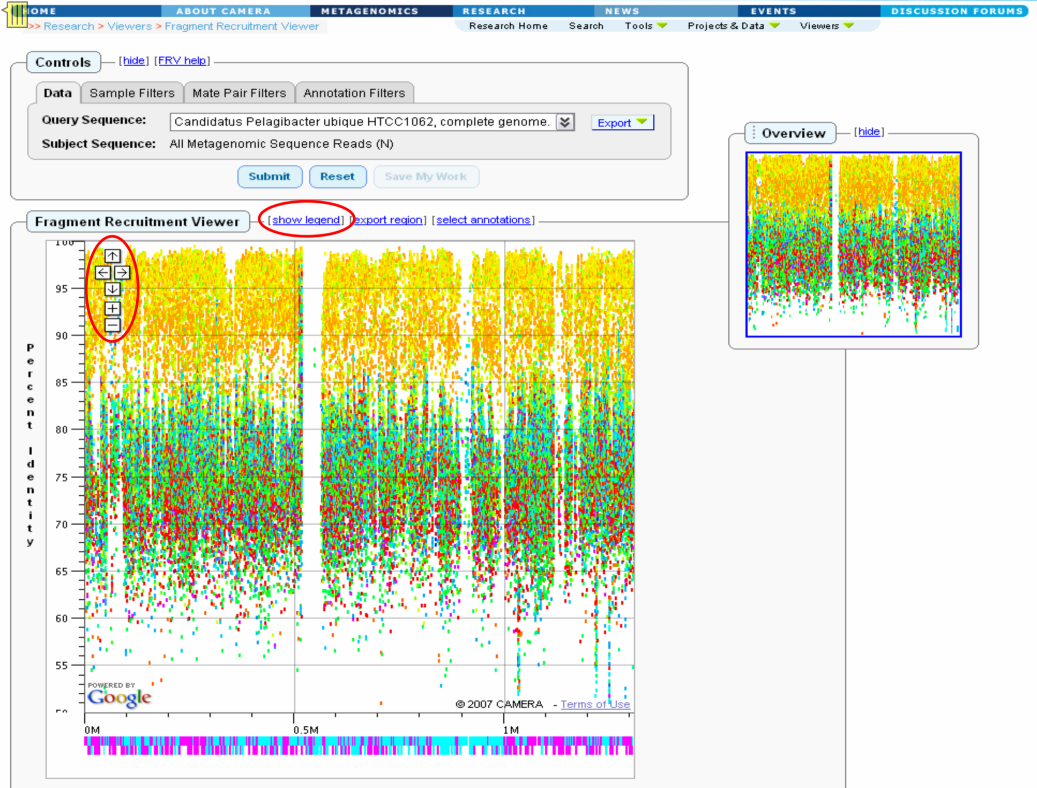






Recent Searches [hide]
 <u>lipase</u> no matches glycerol kinase (about 4,239
matches)
 <u>Cellulase</u> (about 3,163 matches) <u>ATP synthase epsilon</u> (about
2,856 matches)
 <u>dehalogenase</u> (about 4,963 matches)

Recent BLAST Results [hide]							
Job Name	<u>Submit Date</u>	<u>Status</u>	# Hits	Subject Sequences			
test11 02/27/07 02:09 PI		completed	<u>2</u>	GOS: Assembled Sequences (N)			
tst10	02/27/07 02:09 PM	completed	0	GOS: Combined Assembly ORF Peptides (P)			
test09	02/27/07 02:08 PM	completed	<u>1</u>	GOS: All Metagenomic Sequence Reads (N)			
test06	02/27/07 02:05 PM	completed	<u>50</u>	All Metagenomic Sequence Reads (N)			





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Region

Base Pair range: [min]

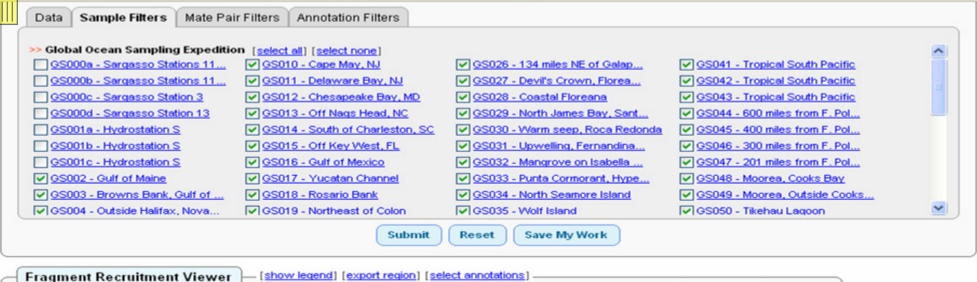
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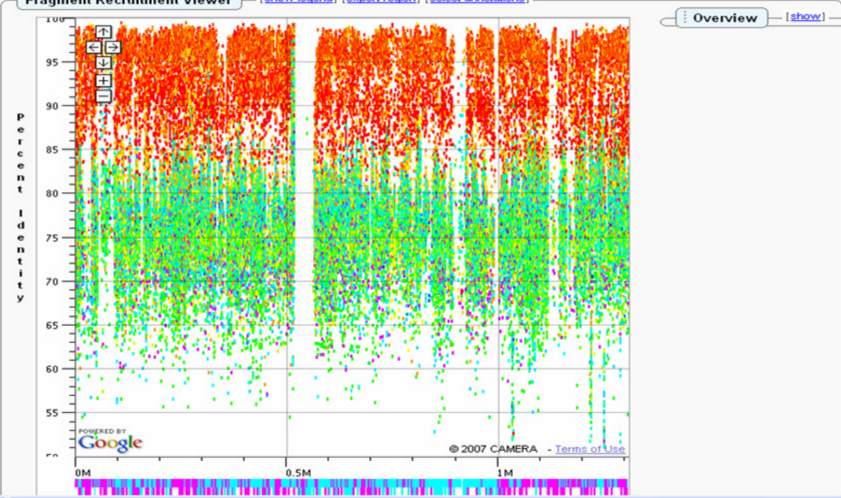
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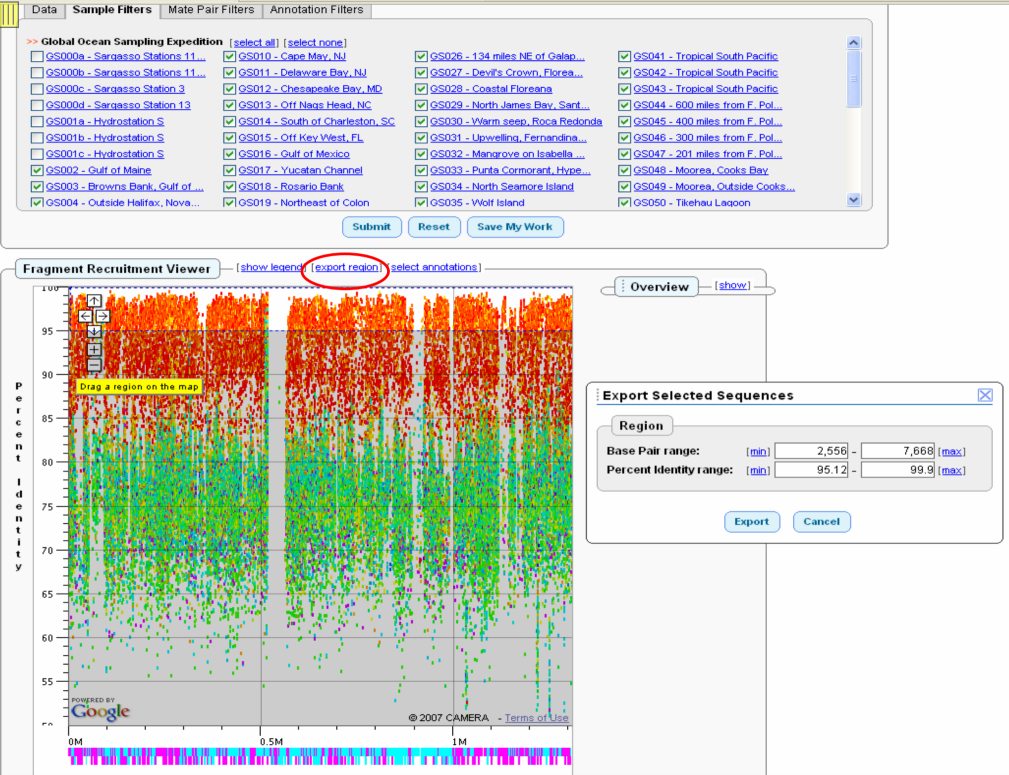
1 - 50 of 61

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🚽 prev 50	aoto 🔻	II next 50 🛾

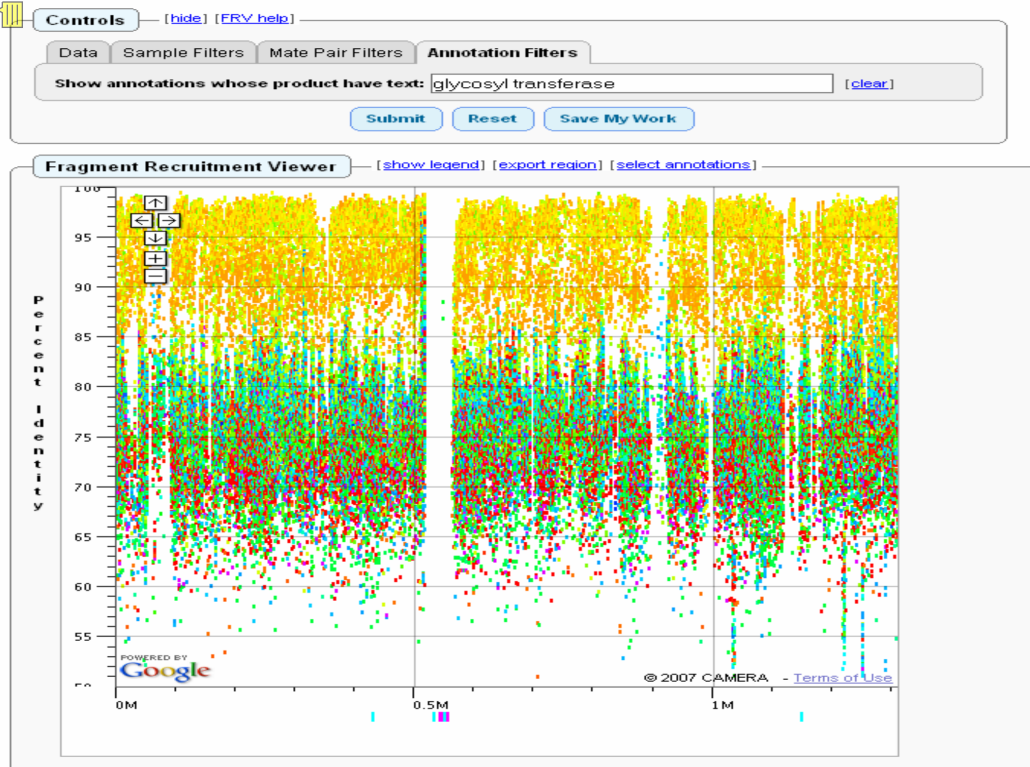
Begin 🔺	End	Strand	Length	<u>Gl Number</u>	Protein ID	Product
512931	513007	+	76	GenelD:3517178		tRNA-IIe
513017	513092	+	75	GenelD:3517179		tRNA-Ala
513101	515997	+	2896	GenelD:3517180		23S ribosomal RNA
516442	516975	+	533	GenelD:3517181	YP_265947.1	glucose-6-phosphate isomerase-like protein
516956	517402	+	446	GenelD:3517182	YP_265948.1	hypothetical protein
517419	518273	+	854	GenelD:3517183	YP_265949.1	NAD dependent epimerase/dehydratase family
518270	519133	-	863	GenelD:3517184	YP_265950.1	NAD dependent epimerase/dehydratase family
519133	520869	-	1736	GenelD:3517185	YP_265951.1	acetolactate synthase II large chain
520922	521581	+	659	GenelD:3517186	YP_265952.1	Protein-L-isoaspartate(D-aspartate) O-methyltransferase (PCMT) family
521587	522627	+	1040	GenelD:3517152	YP_265953.1	zinc-containing alcohol dehydrogenase
522650	523831	+	1181	GenelD:3517153	YP_265954.1	cyclopropane-fatty-acyl-phospholipid synthase (CMAS family)
523831	524610	+	779	GenelD:3517154	YP_265955.1	FabG-like protein
524613	525977	-	1364	GenelD:3517155	YP_265956.1	FAD oxidase family
526036	526809	-	773	GenelD:3517156	YP_265957.1	methyltransferase
526790	528139	+	1349	GenelD:3517157	YP_265958.1	aminotransferase
528210	529664	-	1454	GenelD:3517158	YP_265959.1	deoxynucleotidase
529658	530587	-	929	GenelD:3517159	YP_265960.1	transketolase family
530587	531378	-	791	GenelD:3517160	YP_265961.1	transketolase
531418	532356	+	938	GenelD:3517161	YP_265962.1	UDP-glucose 4 epimerase
532349	533857	+	1508	GenelD:3517162	YP_265963.1	cytidylyltransferase/carbohydrate (or pyrimidine) kinase
533858	534580	+	722	GenelD:3517163	YP_265964.1	spsA-like glycosyl transferase
534588	535577	+	989	GenelD:3517164	YP_265965.1	dTDPglucose 4,6-dehydratase
535581	537269	-	1688	GenelD:3517165	YP_265966.1	unknown membrane protein
537246	537701	+	455	GenelD:3517166	YP_265967.1	unknown membrane protein
537702	538163	+	461	GenelD:3517167	YP_265968.1	unknown membrane protein
538166	538732	+	566	GenelD:3517168	YP_265969.1	phosphoheptose isomerase
538725	539216	-	491	GenelD:3517169	YP 265970.1	D-alvcero-D-manno-hentose 1 7 bisphophate phosphatase







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Base pair position

CAMERA http://camera.calit2.net

- Website: <u>http://camera.calit2.net</u>
- Questions: <u>camera-info@calit2.net</u>

- Saul A. Kravitz
- Rekha Seshadri

skravitz@jcvi.org rseshadri@jcvi.org

Camera (

