CACAO training part I

Jim Hu and Suzi Aleksander For UW Parkside Fall 2014

Outline

- Part I: Gene Ontology and functional annotation
 - How known functions are used to reveal new knowledge
 - Gene Ontology
 - What is an annotation?
 - CACAO
- Part 2: Making annotations and challenges

LEVERAGING WHAT WE KNOW ABOUT FUNCTION

Leveraging what we know about function

- **Functional profiling**: For a list of genes, what functions are important?
 - Genes turned up or down together
 - Disease states
 - Environmental responses
 - Genotypes
 - ...
 - Genes encoding proteins that physically interact
 - Genes conserved in specific taxa
 - Genes found in specific microbial communities

Functional profiling example

Sister grouping of chimpanzees and humans as revealed by genome-wide phylogenetic analysis of brain gene expression profiles

Monica Uddin^{†‡}, Derek E. Wildman^{†‡}, Guozhen Liu^{†§}, Wenbo Xu[§], Robert M. Johnson¹¹, Patrick R. Hof^{||}, Gregory Kapatos^{†,††}, Lawrence I. Grossman[†], and Morris Goodman^{†‡,‡‡}

¹Center for Molecular Medicine and Genetics, Departments of ¹Anatomy and Cell Biology, ¹Biochemistry and Molecular Biology, and ¹¹Psychiatry and Behavioral Neurosciences, Wayne State University School of Medicine, 540 East Canfield Avenue, Detroit, Mi 48201; ⁵Bioinformatics Facility, 5107 Biological Science Building, 5047 Guilen Mall, Detroit, Mi 48202; and ¹Department of Neurobiology, Mount Sinal School of Medicine, One Gustave L. Levy Place, New York, NY 10029

Contributed by Morris Goodman, December 30, 2003

Gene expression profiles from the anterior cingulate cortex (ACC) of human, chimpanzee, gorilla, and macaque samples provide clues about genetic regulatory changes in human and other catarrhine primate brains. The ACC, a cerebral neocortical region, has humanspecific histological features. Physiologically, an individual's ACC displays increased activity during that individual's performance of cognitive tasks. Of ~45,000 prphe cats on microarray chins represent. more vulnerable to Alzheimer's disease than are other pyramidal neurons (17, 18). Physiologically, brain imaging results show increased activity in an individual's ACC when that individual is engaged in cognitive tasks (19–21). The ACC participates in decision making when interfering choices are present, a cognitive role involved in executive function (22). In view of these histological

cognitive tasks. Of ~43,000 preing transcripts of all or most h detected in human ACC samp 15,000, in gorilla and chimpan obtained from gene express expectation that the non-hun gorilla) should be more like humans. Instead, the chimpa human than like the gorilla; panzees are the sister group c biguous expression changes cesses and molecular functio represented in the data, the ch apparent regulatory evolutio important changes in the ance but to a greater extent in hur

Among important changes in the ancestry of both humans and chimpanzees, but to a greater extent in humans, are the up-regulated expression profiles of aerobic energy metabolism genes and neuronal function-related genes, suggesting that increased neuronal activity required increased supplies of energy.

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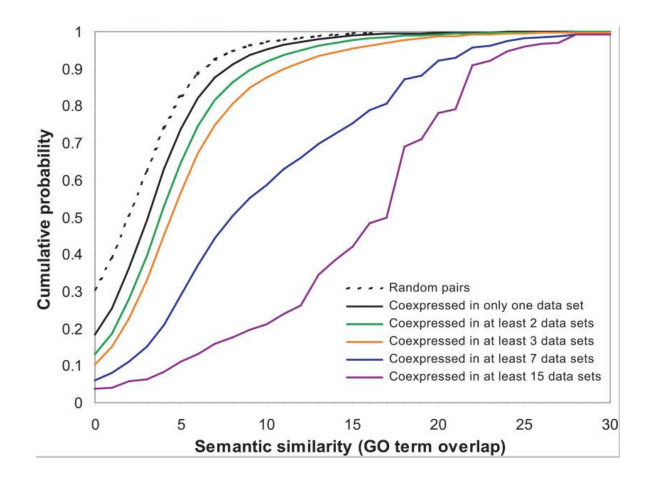
structed the phylogenetic history of the ACC gene expression profiles by treating each probe set as a single character, e.g., analogous to a single genomic locus or a single position in a

Uddin et al. (2004) PNAS 101:2957-2962

Leveraging what we know about function

- **Guilt by association**: For a gene of unknown function, can we infer its function from genes of known function:
 - that are coexpressed across many conditions
 - that are homologs
 - that are coinherited across evolution
 - that physically interact in a multiprotein complex

Coexpression correlates with common function



Lee et al. (2004) Genome Research 14:1085-1094

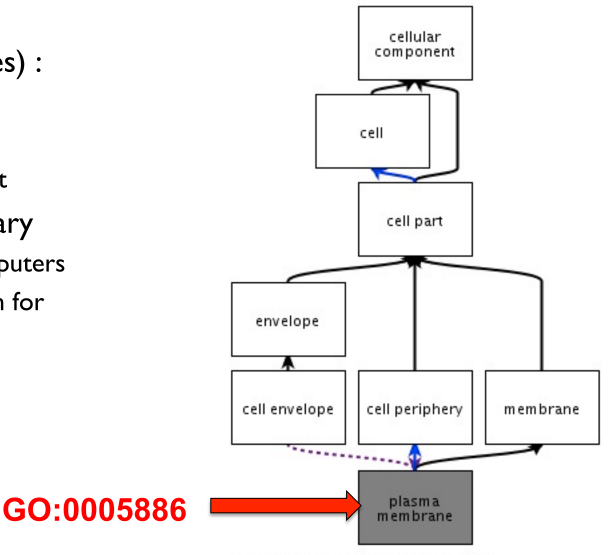
What do we mean by function?

- Massive body of published knowledge
 - Almost useless by itself!!
- We need
 - Knowledge that computers can analyze
 - Common vocabulary across different organisms
 - Disambiguation of synonyms
 - Connection of related ideas that are more or less specific
 - Examples:
 - polygon quadrilateral rectangle square
 - Enzyme kinase protein kinase protein tyrosine kinase

GENE ONTOLOGY

Gene Ontology (GO)

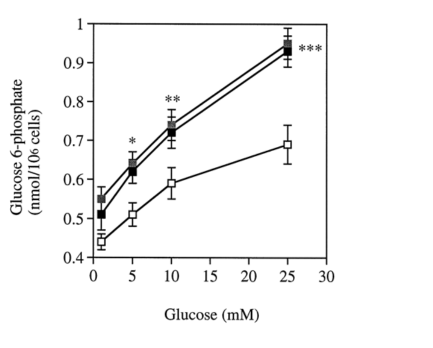
- 3 aspects (ontologies) :
 - Molecular Function
 - Biological Process
 - Cellular Component
- Controlled vocabulary
 - ID number for computers
 - Name and definition for humans
- Relationships



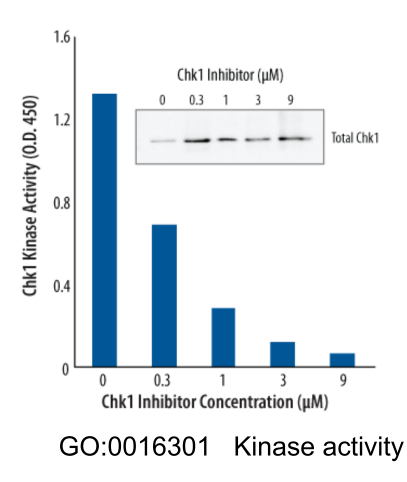
QuickGO - http://www.ebi.ac.uk/QuickGO

Molecular Function

 activities = what a protein can do by itself

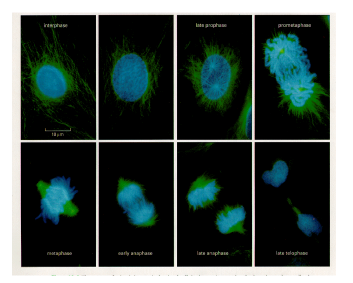






Biological Process

- a commonly recognized series of events
 - Including, but not just biochemical pathways



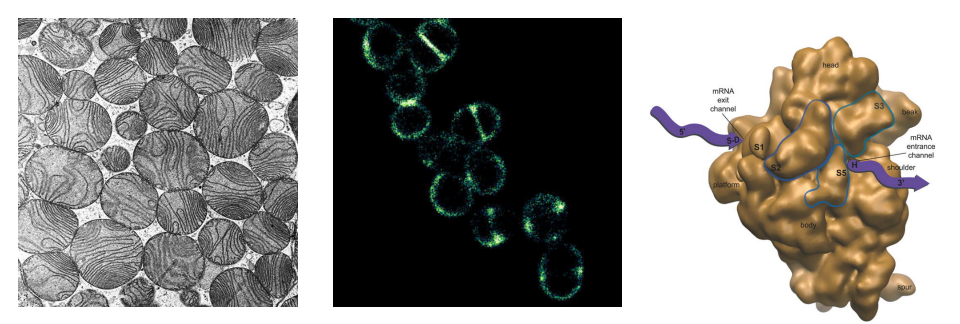
Promoter Factor sigma DNA helix RNA Polymerase Initiation Stop Elongation Termination SNA

GO:0051301 cell division GO:0006351 transcription, DNA dependent

From ridge.icu.ac.jp,

Cellular Component

- where a gene product acts
 - Subcellular location
 - Multicomponent complex

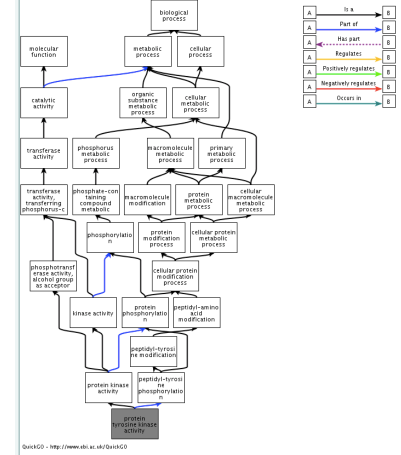


GO:0005739 GO:0009274 GO:0005840 mitochondrion peptidoglycan-based cell wall ribosome

From visualphotos.com, epmm.group.shef.ac.uk, wikimedia commons

GO terms

- ID numbers
- Definitions
- Relationships
 - Directed Acyclic Graph
- GO terms provide a way to describe functions, now we have to associate them with genes!
 - AKA GO annotation



GO ANNOTATION

Dictionary Thesaurus

Q annotation

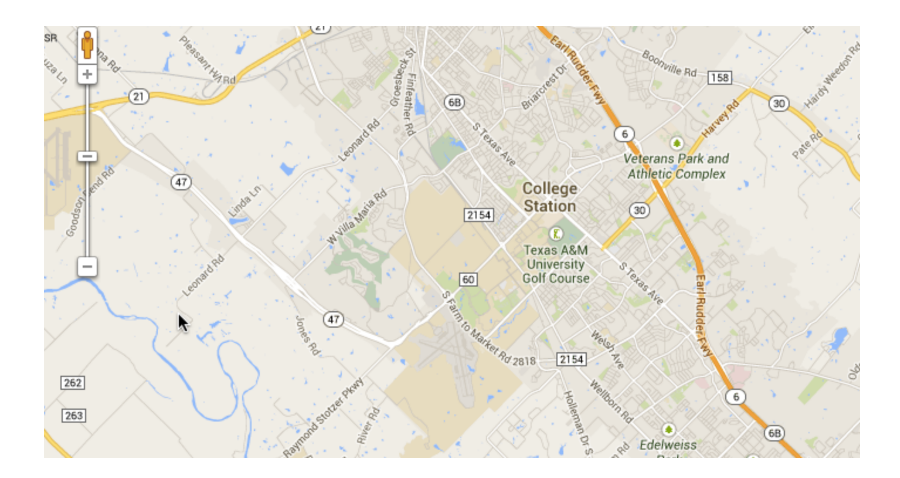
an•no•ta•tion | anə tā sH ən|

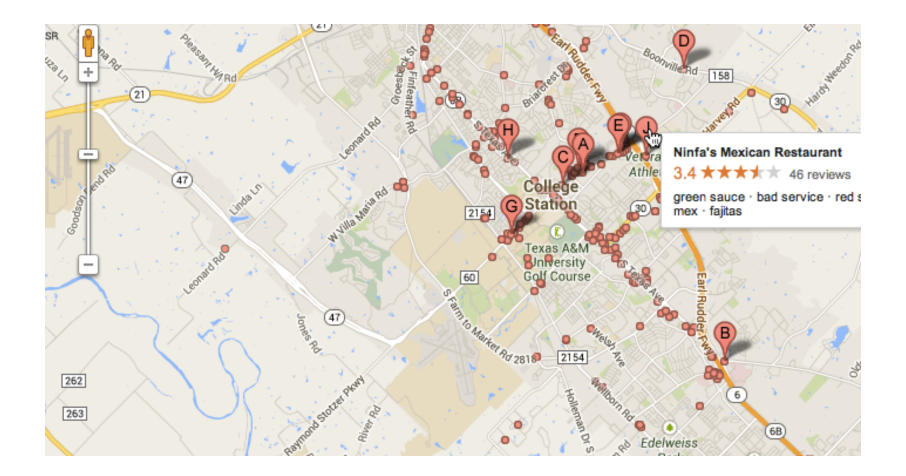
noun

a note of explanation or comment added to a text or diagram : marginal annotations.

the action of annotating a text or diagram : annotation of prescribed texts.

ORIGIN late Middle English : from French, or from Latin *annotatio(n-)*, from the verb *annotare* (see ANNOTATE).





Enter HAMLET. Full Summary

HAMLET

- 56 To be, or not to be: that is the question:
- 57 Whether 'tis nobler in the mind to suffer
- 58 The slings and arrows of outrageous fortune,
- 59 Or to take arms against a sea of troubles,
- 60 And by opposing end them? To die, to sleep-
- 61 No more—and by a sleep to say we end
- 62 The heart-ache and the thousand natural shocks
- 63 That flesh is heir to, 'tis a consummation
- 64 Devoutly to be wish'd. To die, to sleep;
- 65 To sleep: perchance to dream: ay, there's the rub;
- 66 For in that sleep of death what dreams may come
- 67 When we have shuffled off this mortal coil,
- 68 Must give us pause: there's the respect
- 69 That makes calamity of so long life;
- 70 For who would bear the whips and scorns of time,
- 71 The oppressor's wrong, the proud man's contumely,
- 72 The pangs of despised love, the law's delay,
- <u>73</u> The insolence of office and the spurns
- 74 That patient merit of the unworthy takes,
- 75 When he himself might his quietus make
- 76 With a bare bodkin? Who would fardels bear,
- 77 To grunt and sweat under a weary life,
- 78 But that the dread of something after death,
- 79 The undiscover'd country from whose bourn
- 80 No traveller returns, puzzles the will
- 81 And makes us rather bear those ills we have
- 82 Than fly to others that we know not of?
- 83 Thus conscience does make cowards of us all;

\varTheta 🔿 🔿 Notes to Haml...ct 3, Scene 1 🖉

57. suffer: endure patiently.

58. **slings:** *i.e.*, projectiles launched from slings.

60. To die, to sleep — / No more —: This sequence puzzles me. "To sleep" seems to be a comforting way of describing what it means "to die," but "No more" could mean "to dream no more"; remember that Hamlet said to Rosencrantz and Guildenstern, "I could be bounded in a nutshell and count / myself a king of infinite space, were it not that I / have bad dreams. On the other hand, "No more" could be all-encompassing: no more "slings and arrows"; no more "sea of troubles"; no more questions about what would be "nobler in the mind."

63. consummation: completion, end.

65. **rub:** *i.e.*, obstacle, catch. The term comes from the game Americans know as lawn bowling, in which "A rub is some fault in the surface of the green that stops a bowl or diverts it from its intended direction" (World Wide Words: Michael Quinton writes on International English from a British Viewpoint).

67. **shuffled off:** sloughed, cast off. **this mortal coil:** the turmoil of this mortal life.

- 68. respect: consideration.
- 69. of so long life: so long-lived.
- 70. bear the whips and scorns of time: *i.e.*, endure the punishments and insults that always come with the passage of time.

http://www.shakespeare-navigators.com/hamlet/H31.html

Levels of annotation for genomes

- Metadata
 - What is this genome?
- Features
 - Where are things in the sequence?
- Products
 - What do we know about the features?
- Systems
 - What do the products do?
 - individually?
 - Working together?

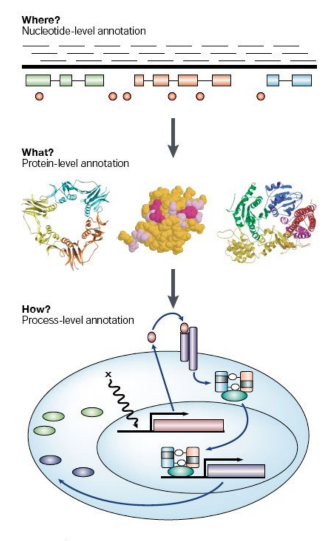


Figure 1 | The three layers of genome annotation: where, what and how?

L. Stein (2001) Nature Reviews Genetics 2:493

Functional Annotation w/GO

- Annotation: a note that is made while reading any form of text
- GO Annotation: a database entry in a specific format that associates a GO term with a gene product made based on evidence in a peer-reviewed paper
 - Specific format makes the annotations readable by both computers and humans
 - GO annotations capture the chain of evidence for how functions were inferred from experiments
 - More when we talk about CACAO

Where do annotations come from?

Journal home > Archive > Letters to Nature > Abstract

Letters to Nature

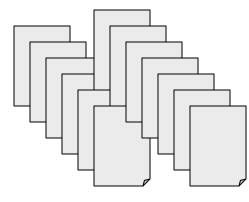
Nature 425, 628-633 (9 October 2003) | doi: 10.1038/nature02030

Basal body dysfunction is a likely cause of pleiotropic Bardet-Biedl syndrome

Stephen J. Ansleyl-å, Jose L. Badanol-å, Oliver E. Blacqueð-å, Josephine Hilf-, Bethan E. Hoskinsl-å, Carmen C. Leitch-, Jun Chul Kim³, Alison J. Ross¹, Erica R. Eichers³, Tanya M. Teslovich-, Allan K. Mah³, Robert C. Johnsen³, John C. Cavender², Richard Alan Lewis³.⁴, Michel R. Leroux³, Philip L. Beales⁴ and Nicholas Katanisl-⁴

Berdet – Biedl syndrome (BBS) is a genetically heterogeneous disorder characterized primarily by retinal dystrophy, obesity, polydactyly, renal maformations and learning disabilities. Although five BBS genes have been cloned^{1, 2, 3, 4, 5, 6}, the molecular basis of this syndrome remains elusive. Here we show that BBS is probably caused by a defect at the basel body of clited cells. We have cloned a new BBS gene, *BBS*, which encodes a protein with a prokaryotic domain, *pBP*, involved in pilus formation and twitching mobility. In one family, a prokaryotic domain, *pBP*, involved in pilus formation and twitching mobility. In one family, a prokaryotic domain, *pBP*, involved in pilus formation and twitching mobility. In one family, a symmetry, a known defect of the nodal clinum. We have easis found the tBBS localizes specifically to clinate structures, such as the connecting clium of the retina and columnar epithelial cells in the lung. In cells, BBS Blocalizes to centrosmes and basel badies and interacts with PCM1, a protein probably involved in clilogenesis. Finally, we demonstrate that all available caenorizability elevanosmologues are expressed exclusively in cliteted neurons, and contain regulatory elements for RFX, a transcription factor that modulates the expression of genes associated with clilogenesis and intraflegular transport.

Literature





Biocurators (rate limiting)

Datasets

Databases need help!

- >21 million peer-reviewed articles in PubMed
- Many millions of proteins recorded in UniProt

Search	Blast	Align	Retrieve ID Mapping *			
Search in		Query				
Protein Knowledgebase (UniProtKB)		human Search Advanced Search » Clear				
			by score descending ⊠			
Browse by taxonor						
Browse by taxonor						
Browse by taxonor	my, keyword, gene					
Browse by taxonor	my, keyword, gene					
	my, keyword, gene		by score descending ⊠ class or pathway I a to Reduce sequence redundancy to 100%, 90% or 50%			

http://www.uniprot.org



What is CACAO?

- Community Assessment of Community Annotation with Ontologies (CACAO)
 - Annotation of gene function
 - Competition
 - Within a class
 - Between teams at different schools
 - More details next week

How does CACAO work?

- Working in teams we will use the GONUTS website:
 - <u>http://gowiki.tamu.edu</u>
- Multiple innings: each is two weeks
 - Annotation week: you make annotations on the website to get points
 - Challenge week: you challenge annotations made by other teams to steal their points
- You can make as many annotations as you want.
 - You pick the topic
 - You have to convince us that they are correct.
 - The default is that they are wrong!!
- Your annotations could end up in databases used by researchers all over the world

How does CACAO work?

- Getting help is not cheating!
 - Talk to your teammates
 - Ask us questions
 - Talk to other professors
 - Email authors of papers

What to annotate

- You can start with a paper
 - Find the proteins discussed
 - Start with a GO term
- You can start with a protein
 - Find papers about the protein
- Either way, don't get stuck on what you started with
 - Your first paper may not have **experiments** about function
 - Reading about your initial protein may lead you to better information about other proteins

Functional Annotation w/GO

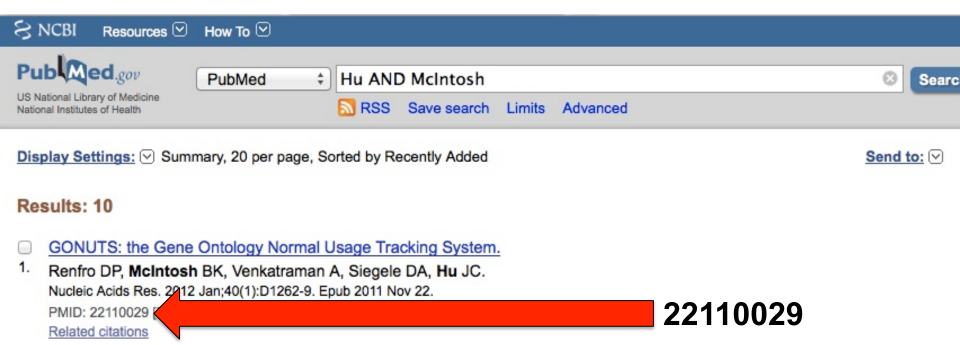
- Annotation: a note that is made while reading any form of text
- GO Annotation: a database entry in a specific format that associates a GO term with a gene product made based on evidence in a peer-reviewed paper

Starting with a paper

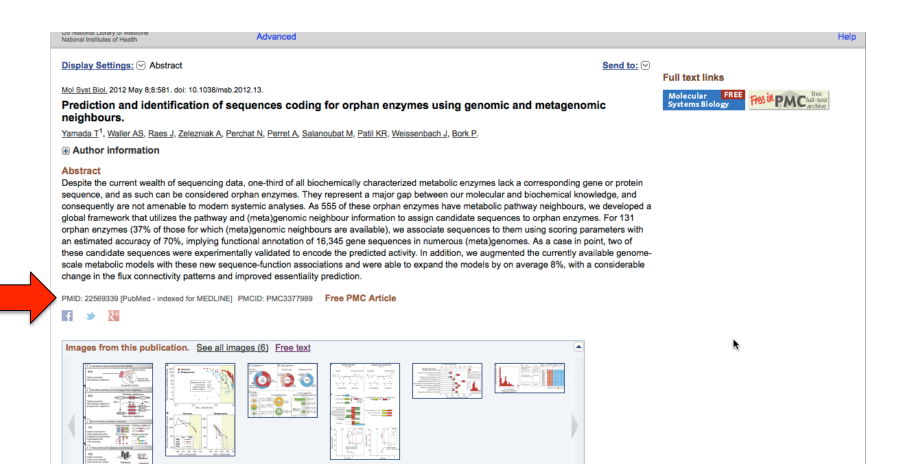
- Need a scientific paper with experimental data
 - Use PubMed: <u>http://www.ncbi.nlm.nih.gov/pubmed/</u>
 - Or use an alias like <u>http://pubmed.com</u>
 - No review articles, no books, no textbooks, no wikipedia articles, no class notes...
 - BUT you should start with those!
 - DON'T start with the first paper you see from a random PubMed search

Starting with a paper

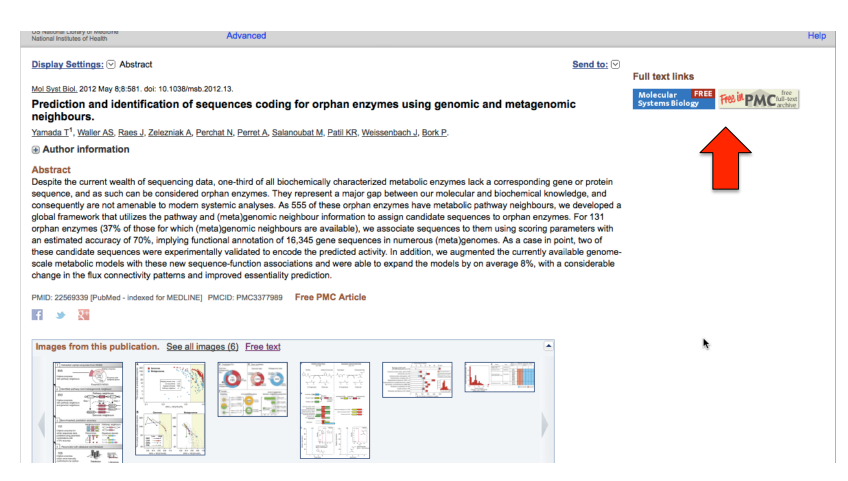
- Need a scientific paper with experimental data
 - PubMed review?
 - We refer to the paper through the PMID number
 - Not the full citation



Pubmed record



Getting the full text



- The abstract is not enough
 - But may be enough to reject a paper!!!

Getting the full text

- Some papers are open access
 - Pubmed Central
 - Journal sites
- Others are pay only
 - Don't pay real \$\$!
 - Your library may have subscriptions
 - Pick a different paper
 - Email the author and ask for a pdf
 - Send us a copy



ding gene or protein al knowledge, and bours, we developed a enzymes. For 131 ring parameters with se in point, two of ntly available genome-, with a considerable

Alternative path: Start w/Full Text

S NCBI Resources 🖂 How To 🖂							
US National Library of Medicine National Institutes of Health Save search Journal List Limits Advanced							
Display Settings: 🖓 Summary, 20 per page, Sorted by Default order							
Results: 1 to 20 of 596771 <pre></pre>							
 Recent advances in 2D and 3D in vitro systems using primary hepatocytes, alternative hepatocyte sources and non-parenchymal live cells and their use in investigating mechanisms of hepatotoxicity, cell signaling and ADME Patricio Godoy, Nicola J. Hewitt, Ute Albrecht, Melvin E. Andersen, Nariman Ansari, Sudin Bhattacharya, Johannes Georg Bode, Jennifer Bolleyn, Christoph Borner, Jan Böttger, Albert Braeuning, Robert A. Budinsky, Britta Burkhardt, Neil R. Cameron, Giovanni Camussi, Chong-Su Cho, Yun-Jaie Choi, J. Craig Rowlands, Uta Dahmen, Georg Damm, Olaf Dirsch, María Teresa Donato, Jian Dong, Steven Dooley, Dirk Drasdo, Rowena Eakins, Karine Sá Ferreira, Valentina Fonsato, Joanna Fraczek, Rolf Gebhardt, Andrew Gibson, Matthias Glanemann, Chris E. P. Goldring, María José Gómez-Lechón, Geny M. M. Groothuis, Lena Gustavsson, Christelle Guyot, David Hallifax, Seddik Hammad, Adam Hayward, Dieter Häussinger, Claus Hellerbrand, Philip Hewitt, Stefan Hoehme, Hermann-Georg Holzhütter, J. Brian Houston, Jens Hrach, Kiyomi Ito, Hartmut Jaeschke, Verena Keitel, Jens M. Kelm, B. Kevin Park Claus Kordes, Gerd A. Kullak-Ublick, Edward L. LeCluyse, Peng Lu, Jennifer Luebke-Wheeler, Anna Lutz, Daniel J. Maltman, Madler Matz-Soja, Patrick McMullen, Irmgard Merfort, Simon Messner, Christoph Meyer, Jessica Mwinyi, Dean J. Naisbitt, Andreas K. Nussler, Peter Olinga, Francesco Pampaloni, Jingbo Pi, Linda Pluta, Stefan A. Przyborski, Anup Ramachandran, Vera Rogiers, Cliff Rowe, Celine Schelcher, Kathrin Schmich, Michael Schwarz, Bijay Singh, Ernst H. K. Stelzer, Bruno Stieger, Regina Stöber, Yuichi Sugiyama, Ciro Tetta, Wolfgang E. Thasler, Tamara Vanhaecke, Mathieu Vinken, Thomas S. Weiss, Agata Widera, Courtney G. Woods, Jinghai James Xu, Kathy M. Yarborough, Jan G. Hengstler Arch Toxicol. 2013; 87(8): 1315–1530. Published online 2013 August 23. doi: 10.1007/s00204-013-1078-5 PMCID: PMC3753504 Article PubReader PDF-9.1M Supplementary Material 							
 Phosphoinositides: Tiny Lipids With Giant Impact on Cell Regulation Tamas Balla 							

Physiol Rev. 2013 July; 93(3): 1019–1137. doi: 10.1152/physrev.00028.2012 PMCID: PMC3962547 <u>Article PubReader</u>

Alternative path: Start w/Full Text

S NC	BI Resources 🕑 How To 🕑				
	al Library of Medicine stitutes of Health Limits Advanced	Journal list		Sea	
Journal	List > Mol Syst Biol > v.8; 2012 > PMC3377989 molecular Systems			PubReader fo click here to	
		dards in Systems Biology		Formats:	
	Mol Syst Biol. 2012; 8: 581. Published online May 8, 2012. doi: <u>10.1038/msb.2012.13</u> Prediction and identification of sequence	P	MCID: PMC3377989	Article <u>PubReader</u> <u>ePub (beta)</u> Related citations in PubMed The CanOE strategy: integrating geno across multiple prokaryote genomes to	
	genomic and metagenomic neighbours				
Mol Syst Biol	<u>Takuji Yamada, ¹ Alison S Waller, ¹ Jeroen Raes</u> , ^{2,3} <u>Alekse</u> <u>Salanoubat</u> , ^{5,6,7} <u>Kiran R Patil</u> , ¹ <u>Jean Weissenbach</u> , ^{5,6,7} ar <u>Author information ► Article notes ► Copyright and License information</u>	Links MedGen			
	This article has been cited by other articles in PMC.	Protein			
	Abstract	PubMed			
	Despite the current wealth of sequencing data, one-ti enzymes lack a corresponding gene or protein seque	Taxonomy			
		Taxonomy Tree			

• Good science \neq good for annotation

Second Extracellular Loop of Human Glucagon-like Peptide-1 Receptor (GLP-1R) Differentially Regulates Orthosteric but Not Allosteric Agonist Binding and Function^{*}

Received for publication, September 30, 2011, and in revised form, November 29, 2011 Published, JBC Papers in Press, December 6, 2011, DOI 10.1074/jbc.M111.309369

Cassandra Koole[‡], Denise Wootten[‡], John Simms[‡], Emilia E. Savage[‡], Laurence J. Miller[§], Arthur Christopoulos^{‡1}, and Patrick M. Sexton^{‡2}

From the [‡]Drug Discovery Biology, Monash Institute of Pharmaceutical Sciences and Department of Pharmacology, Monash University, Parkville, Victoria 3052, Australia and the [§]Department of Molecular Pharmacology and Experimental Therapeutics, Mayo Clinic, Scottsdale, Arizona 85259

Background: The ECL2 of the GLP-1R is critical for GLP-1 peptide-mediated selective signaling. **Results:** Mutation of most ECL2 residues to alanine results in changes in binding and/or efficacy of oxyntomodulin and exendin-4 but not allosteric agonists.

Conclusion: ECL2 of the GLP-1R has ligand-specific as well as general effects on peptide agonist-mediated receptor activation. **Significance:** This work provides insight into control of family B GPCR activation transition.

• Good science \neq good for annotation

Robust design and optimization of retroaldol enzymes

Eric A. Althoff,^{1,2} Ling Wang,¹ Lin Jiang,^{1,3} Lars Giger,⁴ Jonathan K. Lassila,⁵ Zhizhi Wang,¹ Matthew Smith,¹ Sanjay Hari,¹ Peter Kast,⁴ Daniel Herschlag,⁵ Donald Hilvert,⁴ and David Baker¹*

¹Department of Biochemistry, University of Washington and HHMI, Seattle, Washington 98195

²Arzeda Corp., Seattle, Washington 98102

³Department of Biological Chemistry, UCLA, Los Angeles, California 90095

⁴Laboratory of Organic Chemistry, ETH Zurich, 8093 Zurich, Switzerland

⁵Department of Biochemistry, Stanford University, Stanford, California 94305

• Good science \neq good for annotation





Vitamin C Enhances the Generation of Mouse and Human Induced Pluripotent Stem Cells

Miguel Angel Esteban,^{1,6} Tao Wang,^{1,6} Baoming Qin,^{1,6} Jiayin Yang,¹ Dajiang Qin,¹ Jinglei Cai,¹ Wen Li,¹ Zhihui Weng,¹ Jiekai Chen,¹ Su Ni,¹ Keshi Chen,¹ Yuan Li,¹ Xiaopeng Liu,¹ Jianyong Xu,¹ Shiqiang Zhang,¹ Feng Li,¹ Wenzhi He,¹ Krystyna Labuda,² Yancheng Song,³ Anja Peterbauer,⁴ Susanne Wolbank,² Heinz Redl,² Mei Zhong,⁵ Daozhang Cai,³ Lingwen Zeng,¹ and Duanqing Pei^{1,*}

¹Stem Cell and Cancer Biology Group, Key Laboratory of Regenerative Biology, South China Institute for Stem Cell Biology and Regenerative Medicine, Guangzhou Institutes of Biomedicine and Health, Chinese Academy of Sciences, Guangzhou 510663, China ²Ludwig Boltzmann Institute for Clinical and Experimental Traumatology, Austrian Cluster for Tissue Regeneration, Vienna 1200, Austria

• Good science \neq good for annotation

10624 • The Journal of Neuroscience, August 11, 2010 • 30(32):10624-10638

Neurobiology of Disease

Excess Phosphoinositide 3-Kinase Subunit Synthesis and Activity as a Novel Therapeutic Target in Fragile X Syndrome

Christina Gross,¹ Mika Nakamoto,^{2*} Xiaodi Yao,^{1*} Chi-Bun Chan,³ So Y. Yim,¹ Keqiang Ye,³ Stephen T. Warren,^{2,4,5} and Gary J. Bassell^{1,6}

Departments of ¹Cell Biology, ²Human Genetics, ³Pathology and Laboratory Medicine, ⁴Biochemistry, ⁵Pediatrics, and ⁶Neurology, Emory University School of Medicine, Atlanta, Georgia 30322

Functional Annotation w/GO

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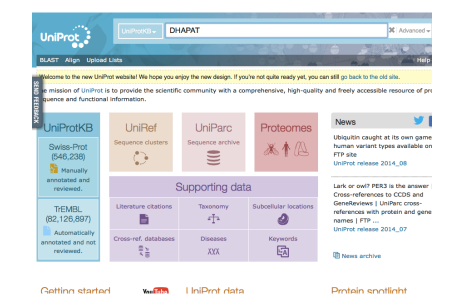
Finding proteins

- Search UniProt for something interesting
- Look in UniProt for the protein(s) in the paper you are reading.

No matter what, you will need to find the protein's accession on UniProt (http://uniprot.org) Use that accession to make a page for that protein on GONUTS (http://gowiki.tamu.edu) Add your GO annotations to the protein's page on GONUTS

UniProt (http://www.uniprot.org)

- If you have a paper, look for an accession
 - UniProt accession
 - NCBI Gene ID
- If you don't have an accession, search by name/keyword



UniProt search results

- Multiple entries
 - Find the right one
 - Icons
 - Gold = Swissprot = reviewed
 - Plain = TrEMBL = automated

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	P98192	GNPAT_MOUSE	<mark>≿</mark>	Dihydroxyacetone phosphate acyltran	Gnpa
	Q9ES71	GNPAT_RAT		Dihydroxyacetone phosphate acyltran	Gnpa
0	P32784	GPT1_YEAST	.	Glycerol-3-phosphate O- acyltransfer	SCT1 GPT1, YBL03 YBL03
	P36148	GPT2_YEAST	÷	Glycerol-3-phosphate O- acyltransfer	GPT2 YKR06

UniProt records

- Lots of information to help you
 - Summary of existing GO annotations
 - Link to QuickGO for complete set of existing annotations
 - Information about the protein



Acyl-CoA + glycerone phosphate = CoA + acylglycerone phosphate.

Pathwayⁱ

Membrane lipid metabolism; glycerophospholipid metabolism.

GO - Molecular function

- glycerone-phosphate O-acyltransferase activity Source: UniProtKB -
- > palmitoyl-CoA hydrolase activity & Source: UniProtKB > receptor binding & Source

GO - Biological process

cellular lipid metabolic process
 Source: Reactome
 cerebellum morphogene

Make sure you have the right protein

- Right species/strain
- Not a fragment
- Sometimes UniProt has multiple entries for the same protein
 - Gold star = SwissProt = reviewed
 - Blank star = TrEMBL = computational entry
- Sometimes the protein you want is not in UniProt
 - May want to find another paper/protein
- Ask for help
 - OK to email the UniProt help desk
 - check your reasoning with us!

Create a protein page in GONUTS

Textpres	so Dev Prod		i/index.php/LAMBD:VLYS GONUTS EcoliWiki - annotation = Goo	gle BugTracker Hub bl	og E.coli Database Portal News (47	L) = Popular =	RSS C	Q* Google	
LAMBD:VLYS - 0						Bmcinto	sh <mark>talk</mark> prefe	rences watchlist o	contributions log
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Entering/editing annotations

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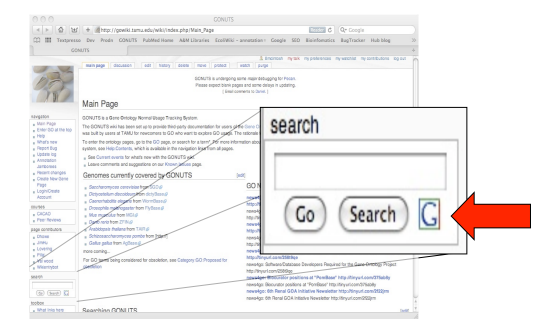
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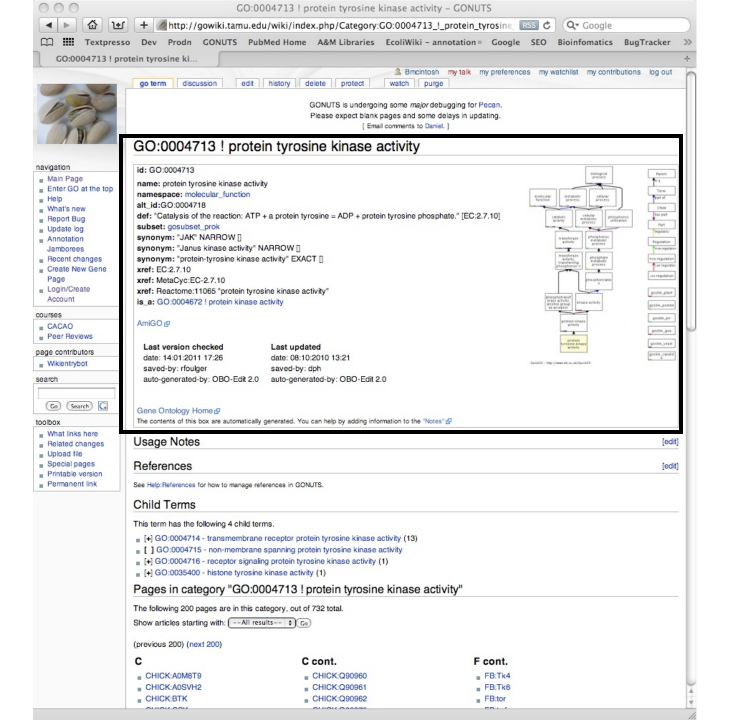
Functional Annotation w/GO

- Annotation: a note that is made while reading any form of text
- GO Annotation: a database entry in a specific format that associates a GO term with a gene product made based on evidence in a peer-reviewed paper

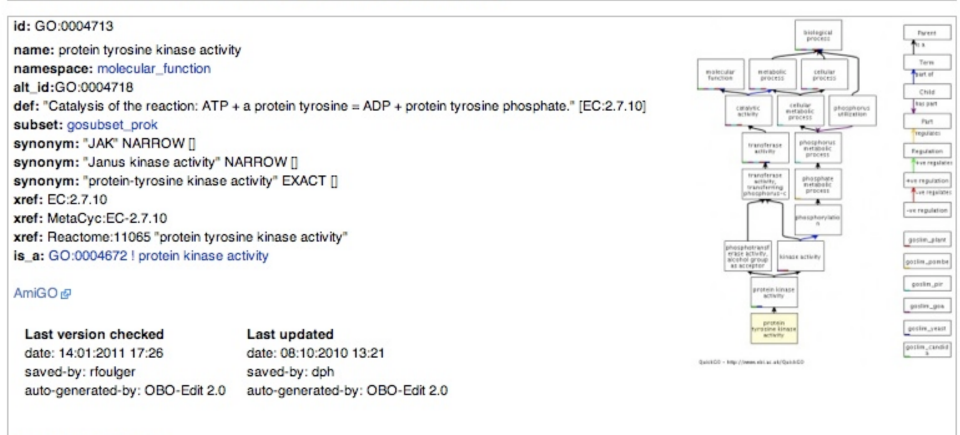
Finding GO terms

- GONUTS: <u>http://gowiki.tamu.edu</u>
- QuickGO: <u>http://www.ebi.ac.uk/QuickGO</u>
- AmiGO: <u>http://amigo.geneontology.org</u>



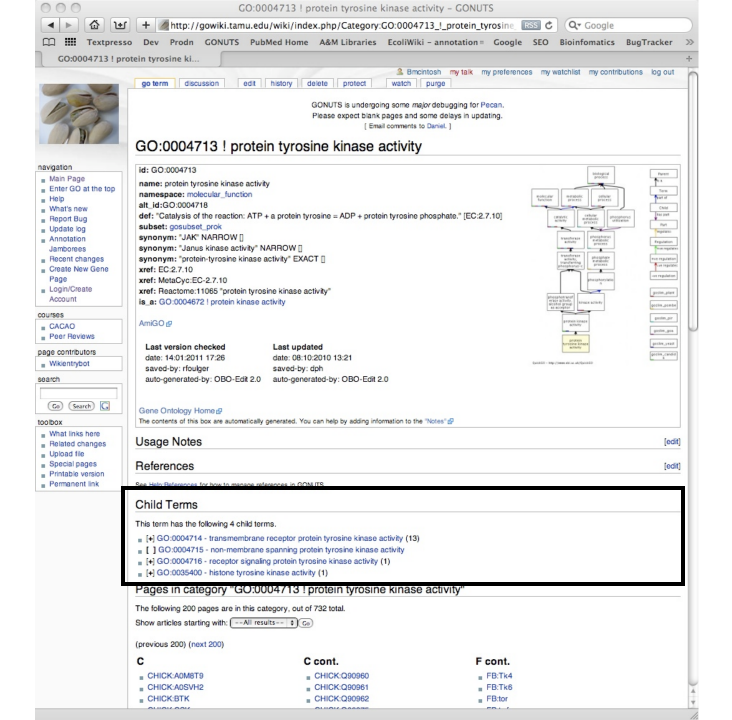


GO:0004713 ! protein tyrosine kinase activity



Gene Ontology Home

The contents of this box are automatically generated. You can help by adding information to the "Notes" @



Strategies

- Search for a keyword and browse the ontology for the right term
 - In GONUTS only search categories if you get too many hits
 - Look at the parents, children, and relatives
 - Use Google, Wikipedia etc. to find alternative search terms
- Look at terms suggested by others for your protein
 - Computational with the IEA evidence code
 - Curators with TAS or IC
- Look at terms used for homologous proteins in model organisms

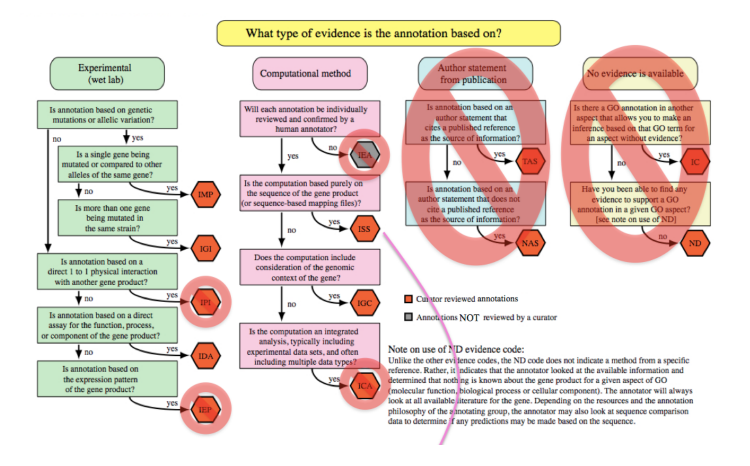
Functional Annotation w/GO

- Annotation: a note that is made while reading any form of text
- GO Annotation: a database entry in a specific format that associates a GO term with a gene product made based on evidence in a peer-reviewed paper

Evidence Codes for CACAO

- Evidence codes describe the type of work or analysis done by the authors
 - IDA: Inferred from Direct Assay
 - IMP: Inferred from Mutant Phenotype
 - NOT just for mutations! Includes inferred from inhibition in vivo by drugs, RNAi, etc.
 - IGI: Inferred from Genetic Interaction
 - ISO: Inferred from Sequence Orthology
 - ISA: Inferred from Sequence Alignment
 - ISM: Inferred from Sequence Model
 - IGC: Inferred from Genomic Context
- Expert biocurators get to use others, but we restrict them for CACAO. If it's not one of these 7, your annotation is incorrect!!!
- <u>http://gowiki.tamu.edu/wiki/index.php/evidence_codes</u>

Decision tree to choose evidence



Evidence pull-down menu

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Public rows can be edited or deleted by any user who can edit Private rows can be edited or deleted by their creator, or by admins

search

Some evidence types require more information

- With/from
- Evidence from sequence comparison
 - With the protein accession for the protein you are comparing to
 - That protein must have experimental annotation to the same GO term
- Evidence from computational analysis
 - With the reference for the analysis tool
- Evidence from genetic interaction
 - With the other gene(s) your protein is interacting with

Evidence Codes for CACAO

- Picking the right evidence code is important
- Use the evidence code decision tree
 - <u>http://gowiki.tamu.edu/wiki/images/3/32/CACAO_decisiontree.pdf</u>
- Use the evidence code guidelines at the GO consortium website:
 - <u>http://www.geneontology.org/GO.evidence.shtml</u>
- Discuss!

Note required for CACAO

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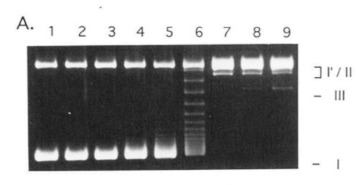
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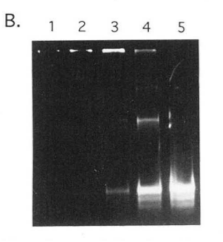
What they did

- Finding the proteins
- Do these tell us about the function?
 - Figure I: sequenced ParC and Part of ParE
 - Figure 2: SDS page of purified proteins
 - Figure 3: Relaxation and decatenation activities of TopolV

- ...

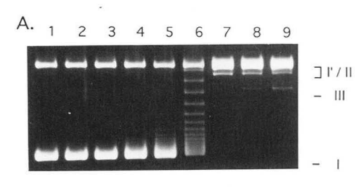
Figure 3

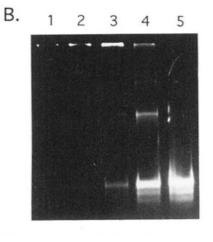




- Panel A: relaxation
- Panel B: decatenation
- What do these mean?

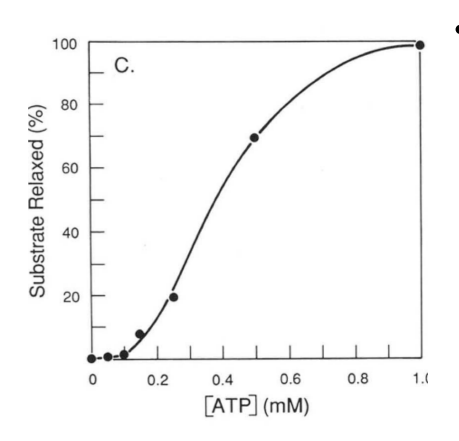
Figure 3





- Panel A: relaxation
- Panel B: decatenation
- What do these mean?
- Panel A shows GO: 0003916 ! DNA topoisomerase activity but does not show what kind
- Panel B shows GO: 0061505 ! DNA topoisomerase II activity

Figure 4



Shows ATP dependence: GO: 0003918 ! DNA topoisomerase type II (ATP-hydrolyzing) activity

GO annotation for E. coli ParC

TableEdit

ECOLI:PARC

Qualifier	÷	
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GO ID	GO:0003918	
GO term name	DNA topoisomerase type II (ATP-hydrolyzing) activity	
Reference	PMID: \$ 8227000	
Evidence Code	IDA: Inferred from Direct Assay \$	
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Notes	Topoisomerase assay in Fig 3. ATP dependent decatenation means it is a Type II from Fig 4	
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