Synthetic Biology = design and engineering of biological systems that aren't found in nature

Why would we want to do this?

- Want to understand natural systems. One of the best ways to understand a system is to change it or make new, related ones
- To fully "understand" a system, we should be able to predict the outcome when we change the system
- For molecular biology, this means:
 - designing new gene circuits and networks
 - modeling the designed systems & predicting their properties
 - making & testing the designs
 - updating our understanding from the model/test agreement

Engineers often look at biological systems & think that the systems are equivalent to electronic circuits

e.g,

fluorescent proteins light bulbs or LEDs transcription factors transistors or logic gates

repressors NOT gates activators OR/AND gates

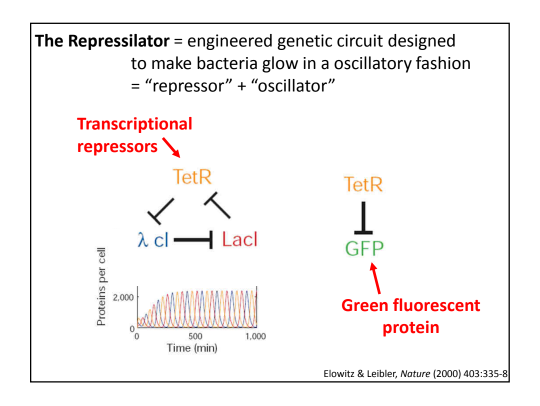
polymerases

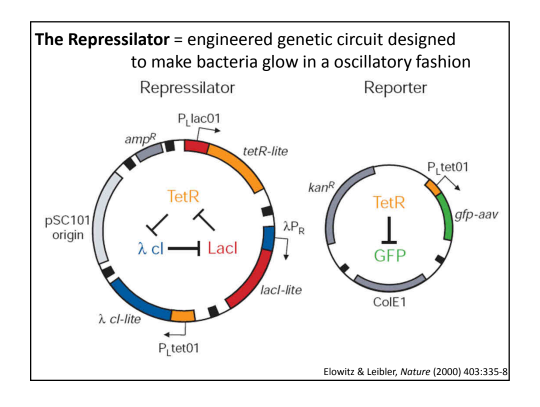
(transcriptional machinery) batteries

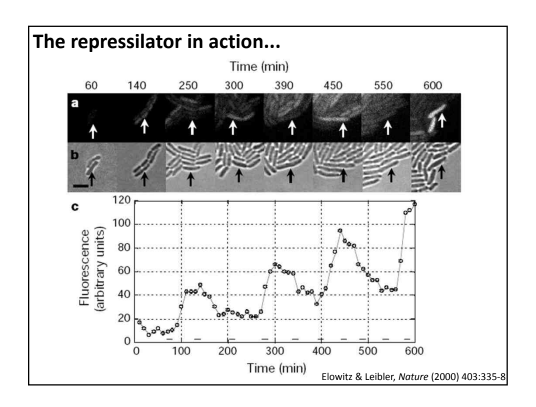
and so on...

Are they right?

→ raises the possibility that biological parts (genes, proteins, etc.) could be combined using the rules established for analog/digital circuits







iGEM: A synthetic biology contest

(from iGEM's web site)

Can simple biological systems be built from standard, interchangeable parts and operated in living cells? Or is biology simply too complicated to be engineered in this way?

iGEM's broader goals include:

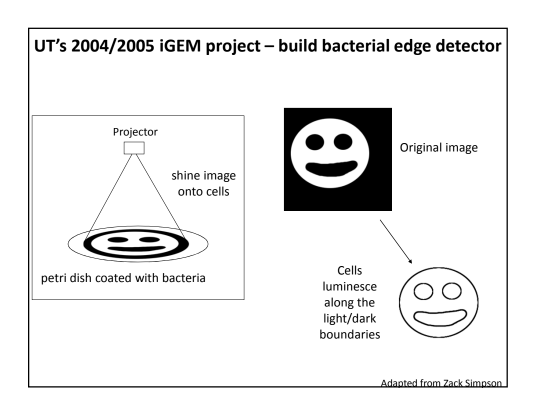
- To enable systematic engineering of biology
- To promote open & transparent development of tools for engineering biology
- To help construct a society that can productively apply biological technology

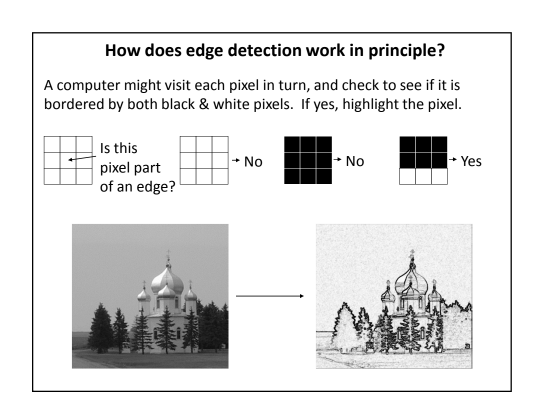
2004: MIT, UT, Princeton, Boston University, Cornell

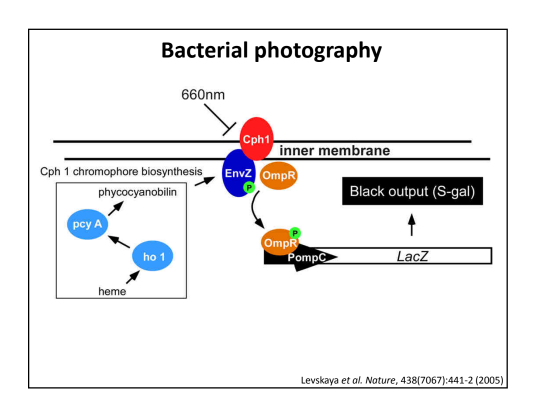
2005: 13 teams (the above + UK, Germany, more...)

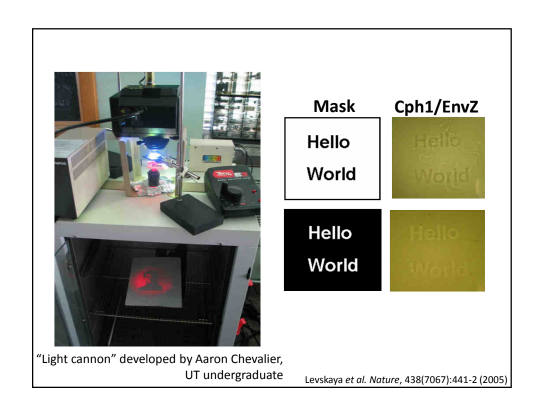
2006: 32 teams, incl. Japan/Latin America/Korea/India/more Europe

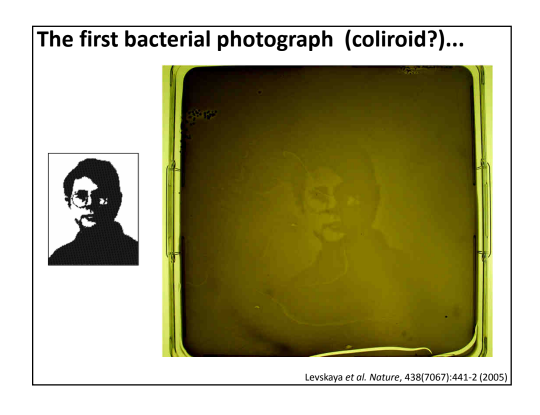
54 teams in 2007, 84 teams in 2008, 112 teams in 2009, 130 teams in 2010, 165 teams in 2011, and 245 teams in 2012 and 2013...



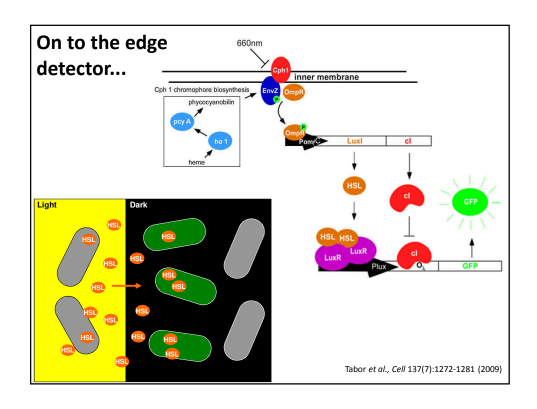


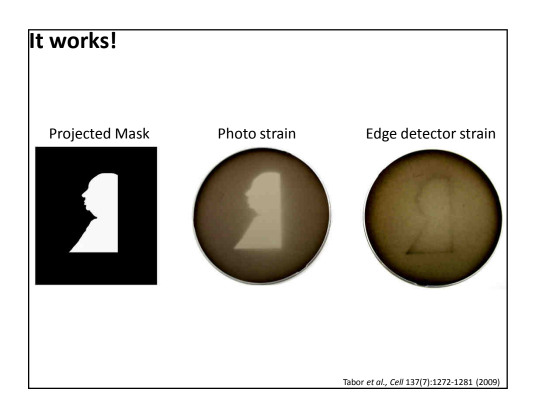


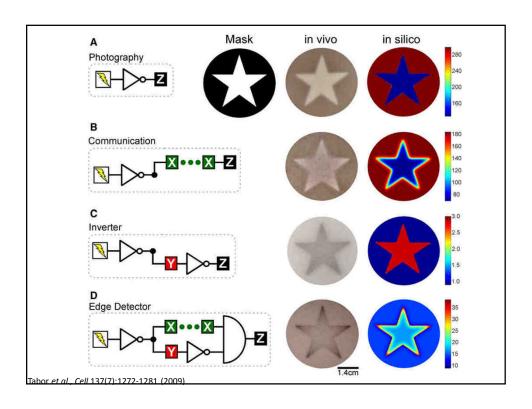


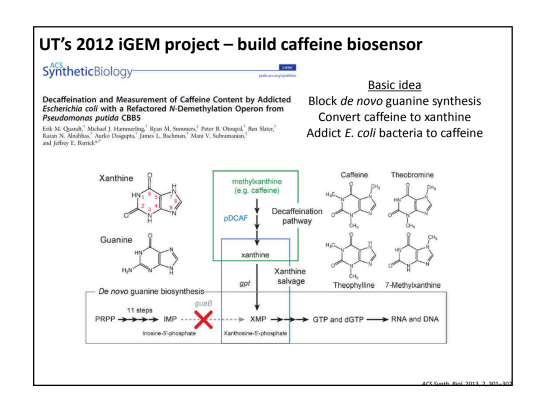


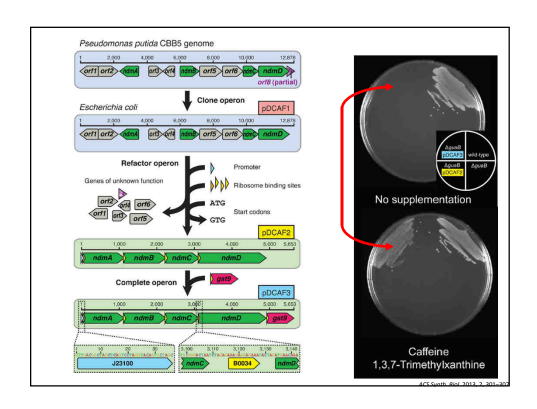


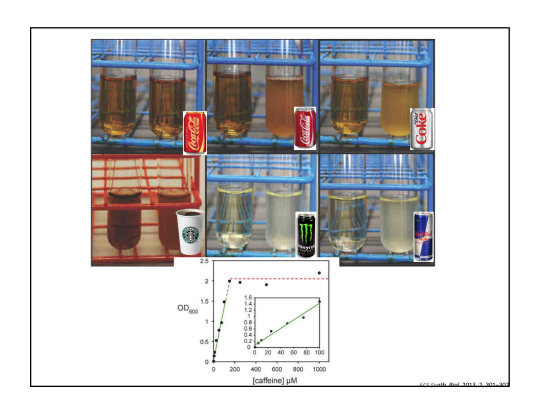


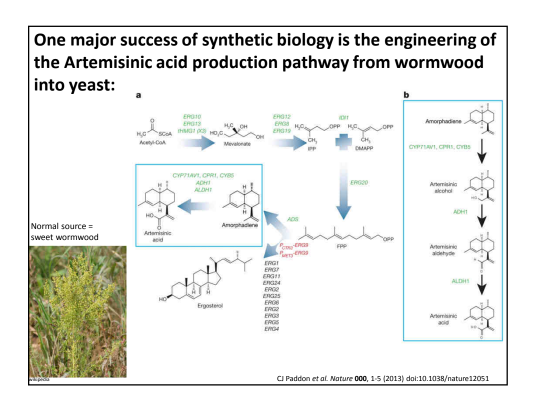


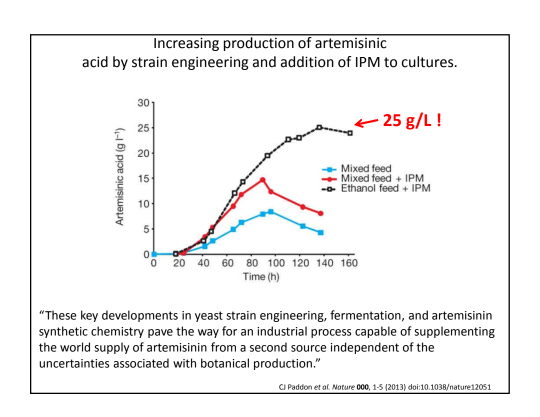












Who needs nature? Made-to-order, designer organisms

Largest Gene Synthesis Supplier in USA

- 100% sequence accuracy guaranteed
- Fastest turnaround: as few as 4 business days
 lowest price: starting at \$0.23/bp



We can now manufacture a complete genome from commodity chemicals

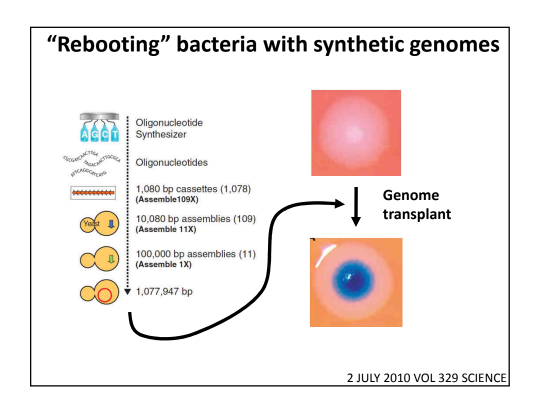
Therefore, we can program whatever changes we want, assuming we can get it into cells...

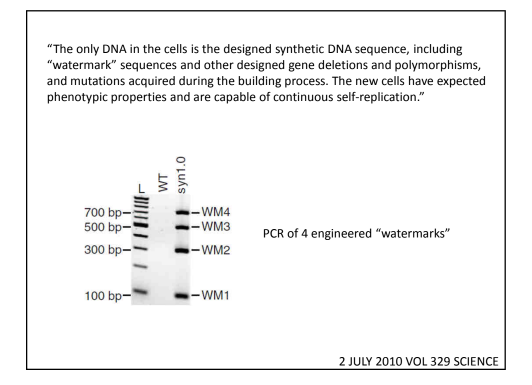
Creation of a Bacterial Cell Controlled by a Chemically Synthesized Genome

Daniel G. Gibson, ¹ John I. Glass, ¹ Carole Lartigue, ¹ Vladimir N. Noskov, ¹ Ray-Yuan Chuang, ¹ Mikkel A. Algire, ¹ Gwynedd A. Benders, ² Michael G. Montague, ¹ Li Ma, ¹ Monzia M. Moodie, ¹ Chuck Merryman, 1 Sanjay Vashee, 1 Radha Krishnakumar, 1 Nacyra Assad-Garcia, 1 Cynthia Andrews-Pfannkoch, Evgeniya A. Denisova, Lei Young, Zhi-Qing Qi, Thomas H. Segall-Shapiro, ¹ Christopher H. Calvey, ¹ Prashanth P. Parmar, ¹ Clyde A. Hutchison III, ² Hamilton O. Smith, ² J. Craig Venter, ^{1,2}*

"We report the design, synthesis, and assembly of the 1.08-megabase pair Mycoplasma mycoides JCVI-syn1.0 genome starting from digitized genome sequence information and its transplantation into a M. capricolum recipient cell to create new M. mycoides cells that are controlled only by the synthetic chromosome."

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But, wait! They only changed DNA, not the rest of the cell!

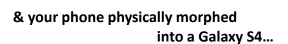
However...

In biology, <u>software encodes the hardware</u>. Most (all?) of the cell is specified by the DNA.

It's as though you bought a Blackberry...



installed the Android operating system...



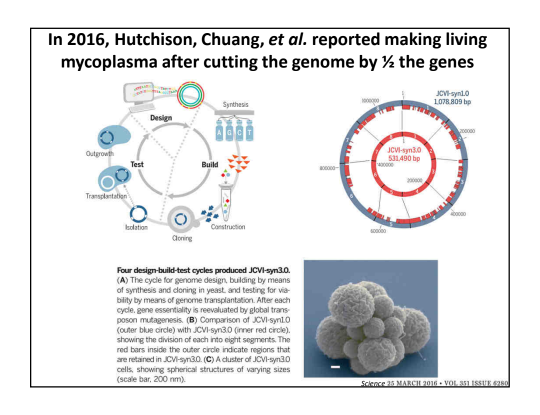


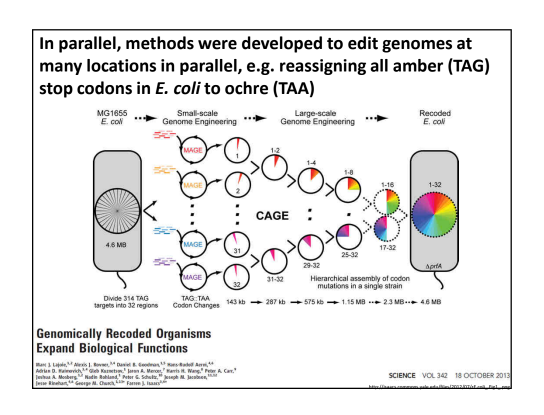
Some good quotes from the paper:

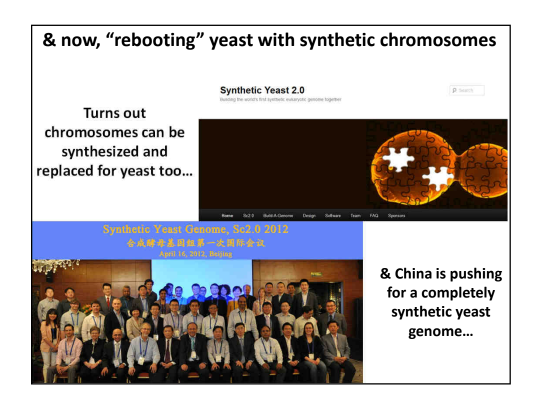
"If the methods described here can be generalized, design, synthesis, assembly, and transplantation of synthetic chromosomes will no longer be a barrier to the progress of synthetic biology."

"We expect that the cost of DNA synthesis will follow what has happened with DNA sequencing and continue to exponentially decrease. Lower synthesis costs combined with automation will enable broad applications for synthetic genomics."

"As synthetic genomic applications expand, we anticipate that this work will continue to raise philosophical issues that have broad societal and ethical implications."







Science April 4, 2014: Vol. 344 no. 6179 pp. 55-58

Total Synthesis of a Functional Designer Eukaryotic Chromosome

Narayana Annaluru, ^{1*} Héloïse Muller, ^{1,2,3,4}, Leslie A. Mitchell, ^{2,5} Sivaprakash Ramalingam, ¹ Giovanni Stracquadanio, ^{2,6} Sarah M. Richardson, ⁶ Jessica S. Dymond, ^{2,7} Zheng Kuang, ² Lisa Z. Scheifele, ^{2,8} Eric M. Cooper, ² Yizhi Cai, ^{2,9} Karen Zeller, ^{2,8} Neta Agmon, ^{2,5} Jeffrey S. Han, ¹⁰ Michalis Hadjithomas, ¹¹ Jennifer Tullman, ⁶ Katrina Caravelli, ^{2,12} Kimberly Cirelli, ^{1,12} Zheyuan Guo, ^{1,13} Viktoriya London, ^{1,13} Apurva Yeluru, ^{1,13} Sindurathy Murugan, ⁶ Karthikeyan Kandavelou, ^{1,14} Nicolas Agier, ^{15,16} Gilles Fischer, ^{15,16} Kun Yang, ^{2,6} J. Andrew Martin, ^{2,6} Murat Bilgel, ³ Pavlo Bohutskyi, ¹³ Kristin M. Boulier, ²² Brian J. Capaldo, ¹³ Joy Chang, ¹³ Kristie Charoen, ³³ Woo Jin Choi, ¹⁵ Peter Deng, ¹¹ James E. Dicarlo, ¹³ Judy Doong, ¹³ Jessiylyn Dunn, ¹³ Jason I. Feinberg, ¹² Christopher Fernandez, ²² Charlotte E. Floria, ¹² David Gladowski, ¹² Pasha Hadidi, ³³ Isabel Ishizuka, ¹² Javaneh Jabbari, ¹² Calvin Y. L. Lau, ¹³ Pablo A. Lee, ¹³ Sean Li, ¹³ Denise Lin, ¹² Matthias E. Linder, ¹² Jonathan Ling, ¹³ Jaime Liu, ¹³ Jonathan Liu, ¹³ Mariya London, ¹² Henry Ma, ³¹ Jessica Mao, ³¹ Jessica E. McDade, ³³ Alexandra McMillan, ³² Aaron M. Moore, ²² Won Chan Oh, ³³ Yu Ouyang, ³⁸ Renus Wong, ³³ Nerina Paul, ¹² Laura C. Paulsen, ¹³ Judy Qiu, ³³ Alex Rhee, ³³ Matthew G. Rubashkin, ³³ Ina Y. Soh, ¹² Nathaniel E. Sotuyo, ¹² Venkatesh Srinivas, ¹³ Allison Suarez, ¹³ Andy Wong, ¹³ Remus Wong, ¹³ Wei Rose Xie, ²² Yijie Xu, ³³ Allen T. Yu, ²² Romain Koszul, ³⁴ Joel S. Bader, ^{2,6} Jef D. Boeke, ^{2,11,5} Srinivasan Chandrasegaran ¹†

"Here, we report the synthesis of a functional 272,871–base pair designer eukaryotic chromosome, synIII, which is based on the 316,617–base pair native *Saccharomyces cerevisiae* chromosome III. Changes to synIII include TAG/TAA stop-codon replacements, deletion of subtelomeric regions, introns, transfer RNAs, transposons, and silent mating loci as well as insertion of loxPsym sites to enable genome scrambling."

Changes engineered into chromosome III

~2.5% of sequence changed

Recoded all amber (TAG) stop codons to ochre (TAA)
Introduced 98 Cre/Lox recombination sites
Introduced unique sequences for PCR and new restriction enzyme sites
Standardized telomeres

Reduced size from 316,617 bp to 272,871 bp (~14% reduction)

Deleted 10 tRNA genes, 21 Ty elements/LTRs, silent mating loci (only one tRNA was essential, moved to a plasmid)

Removed leucine biosynthesis gene LEU2 to be an auxotrophic marker Deleted all introns (affected 7 genes)

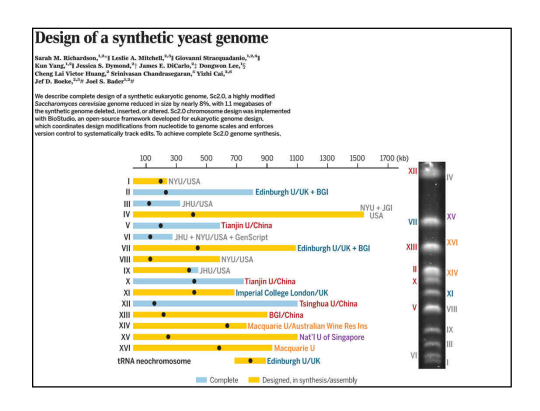
Deleted subtelomeric DNA

Only 10 errors in assembly: 9 single base changes and 1 lost recombinase site

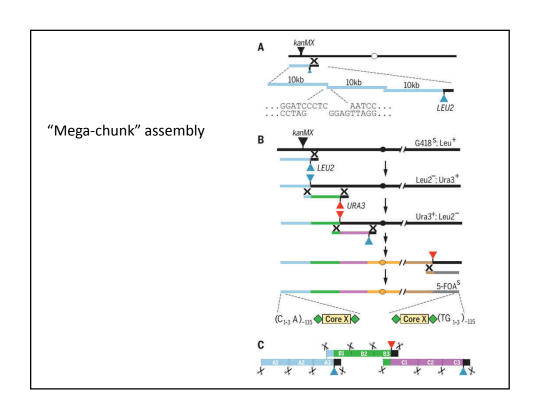
Last year, the Synthetic Yeast Genome Project (Sc2.0) reported on five newly constructed synthetic yeast chromosomes:



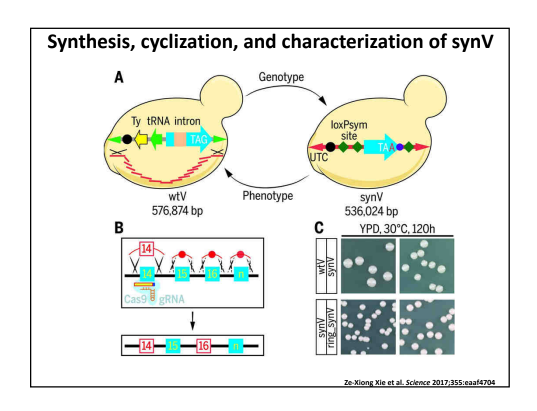
How the cover was made: http://science.sciencemag.org/content/355/6329/eaan1126



Design challenge or amendment	Policy adopted by design team			
Subtelomeric repeats of varying copy number on multiple chromosomes	Delete and monitor for phenotypes as chromosomes are combined. Exception: vitamin biosynthesis genes retain one copy.			
Dispersed repeated genes of high copy number, as well as high-copy COS and seripauperin genes	Delete and monitor for phenotypes as chromosomes are combined.			
loxPsym sites <300 bp apart when inserted algorithmically (not especially useful and more difficult to synthesize)	loxPsym thinning to eliminate the loxPsym site closer to the centromere.			
Stop codon overlaps a second CDS; insertion of loxPsym site would disrupt second CDS; also TAG recoding to TAA could disrupt CDS	Favor preservation of "verified ORFs" over "dubious ORFs" and "uncharacterized ORFs"; always add loxPsym site to a verified ORF in this case			
Tandem repeats inside CDSs (34)	Use GeneDesign's RepeatSmasher module to recode such genes to minimize DNA level repetitiveness, making DNA easier to synthesize and assemble.			
Homopolymer tracts, including frequent A and T tracts, are difficult to synthesize	In synthesis phase, permit 10% length variation for homopolymer tracts >10 bp provided they are in a noncoding region.			
Introns	Delete pre-mRNA introns precisely, except from genes with evidence of a fitness defect caused by intron deletion (35, 36). The HAC1 intron, which uses separate splicing machinery and is known to play a critical role in regulation of the unfolded protein response, was not deleted (9). Delete all tRNA introns precisely.			
Intronically embedded snoRNAs	These are individually nonessential and were deleted with their host introns. They could be "refactored" by insertion into the array of snoRNAs on chr II.			



	WT size	SYN size	No. of stop codon swaps	No. of loxP sites added	bp of PCRTag recoded	bp of RE sites recoded	No. of tRNA deleted	bp of tRNA deleted	bp of repeats deleted
chr01	230208	181030	19	62	3535	210	4	372	3987
chr02	813184	770035	93	271	13651	1215	13	993	7030
chr03	316617	272195	44	100	5272	250	10	794	7358
chr04	1531933	1454671	183	479	25398	2298	28	2261	11674
chr05	576874	536024	61	174	8760	813	20	1471	11181
chr06	270148	242745	30	69	4553	369	10	835	9297
chr07	1090940	1028952	126	380	17910	1572	36	2887	13284
chr08	562643	506705	61	186	9980	714	11	878	19019
chr09	439885	405513	54	142	7943	436	10	736	11632
chr10	745751	707459	85	249	12582	1102	24	1853	7523
chr11	666816	659617	68	199	11769	1017	15	1243	4214
chr12	1078177	999406	122	291	15129	1539	19	1646	10843
chr13	924431	883749	100	337	15911	0	21	1691	7673
chr14	784333	753096	96	260	13329	1113	14	1152	5115
chr15	1091291	1048343	147	399	18015	2058	20	1612	9542
chr16	948066	902994	127	334	15493	1374	17	1338	10048
Total	12071297	11352534	1416	3932	199230	16080	272	21762	149420



Let's end the lectures on a fun note, with some speculative near-future synthetic biology experiments



Science fiction? or not? You be the judge!

"De-extincting" extinct species



Remember Dolly, the cloned sheep?

What if the cells being cloned came from an extinct animal and were put into a surrogate mother?

Would that resurrect the species?

This was tried in 2009 for the Pyrenean ibex, and almost worked...



Cloned goat dies after attempt to bring species back from extinction

Groundbreaking experiment fails, but scientists pave way for 'return of other creatures

But now there's another way!

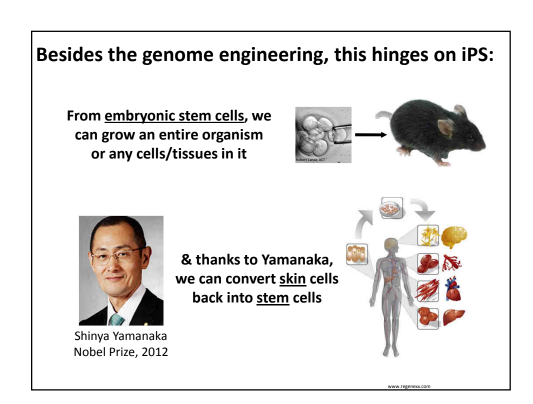
- > We can sequence a genome in a few days for a few \$K
- > We can synthesize or alter big pieces of the DNA
- ➤ We can (almost) "reboot" cells with this DNA
- We can convert cells to stem cells to embryos
- > We can in vitro fertilize animals

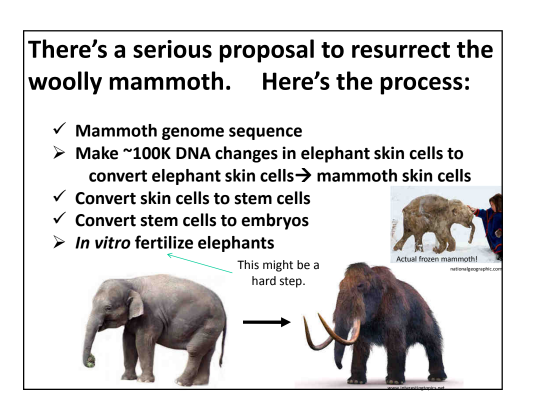
So why not just "edit" the genomes of the closest living animals to be like their extinct relatives?

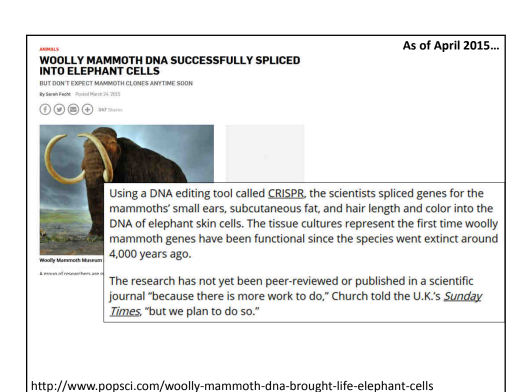


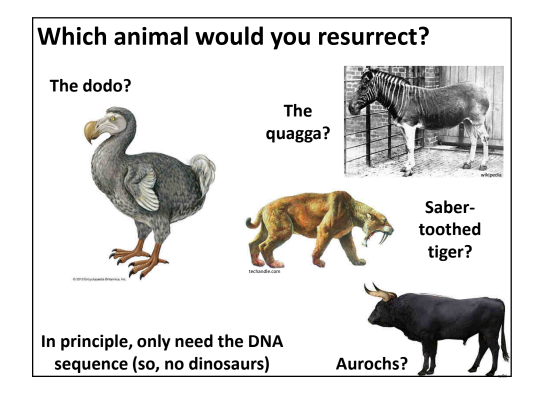
Sound familiar?

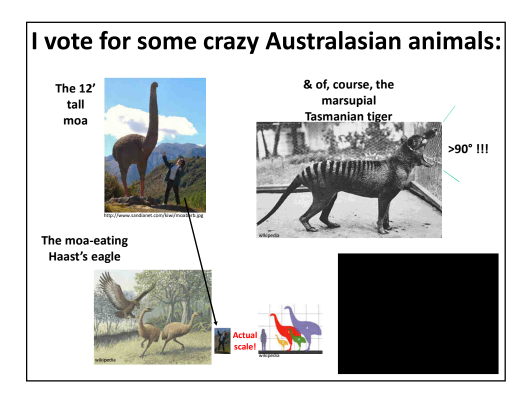
tp://jurassicpark.wikia.com











What about neanderthal? Should we do it?

Svante Pääbo



- \checkmark Human and neanderthal genome sequence
- ➤ Edit DNA in human skin cells to convert convert human skin cells → neanderthal skin cells
 - → I give this step 10 years max before we can do this
- ✓ Convert skin cells to stem cells
- √ Convert stem cells to embryos
- ✓ In vitro fertilize a surrogate mother

