

Toward *de novo* design of immune silent protein and peptide therapeutics

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FDA – CERSI Collaborative Workshop "Predictive Immunogenicity for Better Clinical Outcomes" October 3 and 4, 2018

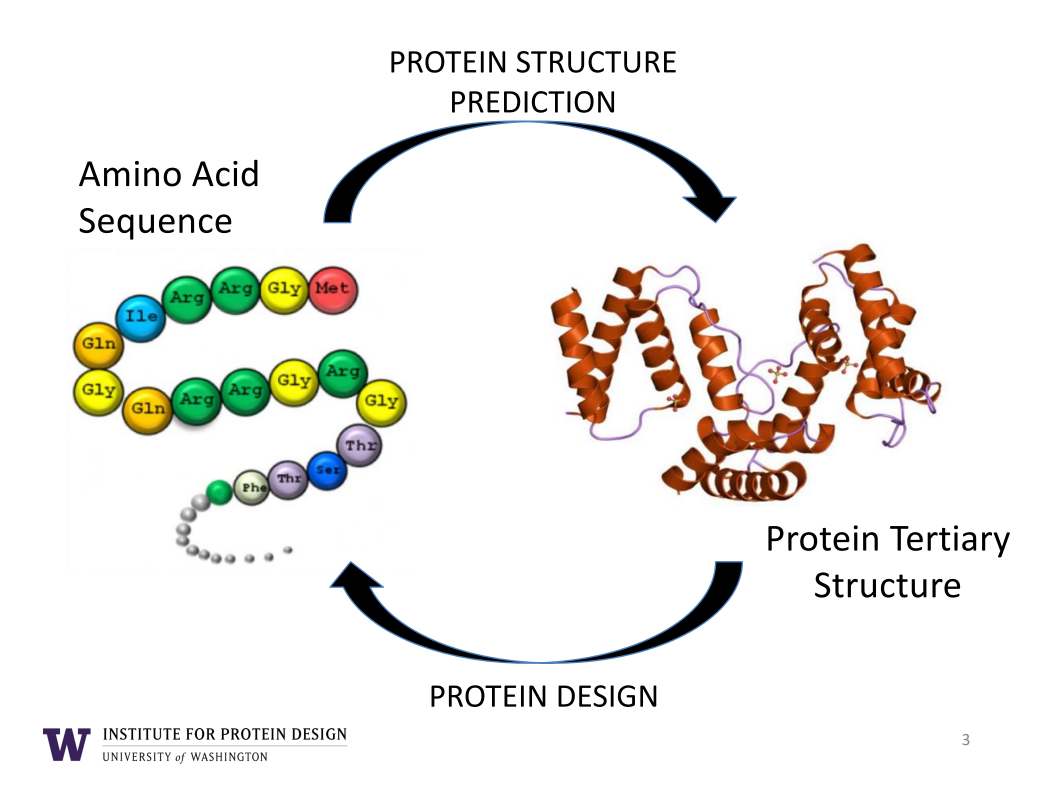
Design a new world of synthetic proteins

- Founded in 2012 by Dr. David Baker
- Organized within the School of Medicine, Biochemistry
- ~140 Person Umbrella Organization
 - Faculty PIs (Baker, DiMaio, King, Gu, Bradley)
 - WRF Innovation Fellows (21)
 - Translational Investigators (1)
 - Research Staff (17)
 - Postdocs (27)
 - Graduate Students (36)
 - Admin (6)
 - Undergrads (27)
 - High School Student (1)

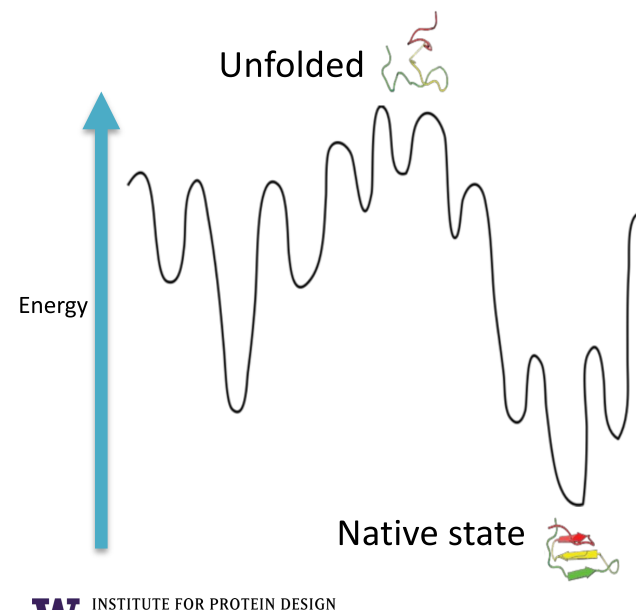








The folded states of proteins are likely global energy minima for their sequences





Protein Structure Prediction: find lowest energy structure for fixed sequence

Protein Design: find a sequence for which desired structure has lowest energy

Sample structures and sequences, and evaluate energies using **Rosetta molecular modeling suite**

15 Years Ago (2003): First De Novo Design

 First computational de novo design of a novel protein fold (Top 7) with atomic level accuracy.

Research Articles

Design of a Novel Globular Protein Fold with Atomic-Level Accuracy

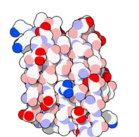
Brian Kuhlman,^{1*}† Gautam Dantas,^{1*} Gregory C. Ireton,⁴ Gabriele Varani,^{1,2} Barry L. Stoddard,⁴ David Baker^{1,3}‡

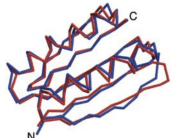
A major challenge of computational protein design is the creation of novel proteins with arbitrarily chosen three-dimensional structures. Here, we used a general computational strategy that iterates between sequence design and structure prediction to design a 93-residue α/β protein called Top7 with a novel sequence and topology. Top7 was found experimentally to be folded and extremely stable, and the x-ray crystal structure of Top7 is similar (root mean square deviation equals 1.2 angstroms) to the design model. The ability to design a new protein fold makes possible the exploration of the large regions of the protein universe not yet observed in nature.

Kuhlman et al, Science 2003

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Model X-ray Structure

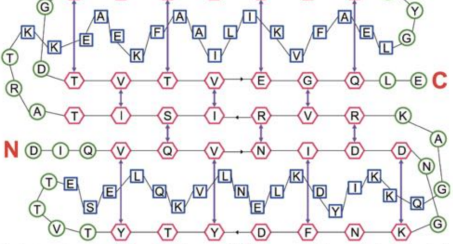
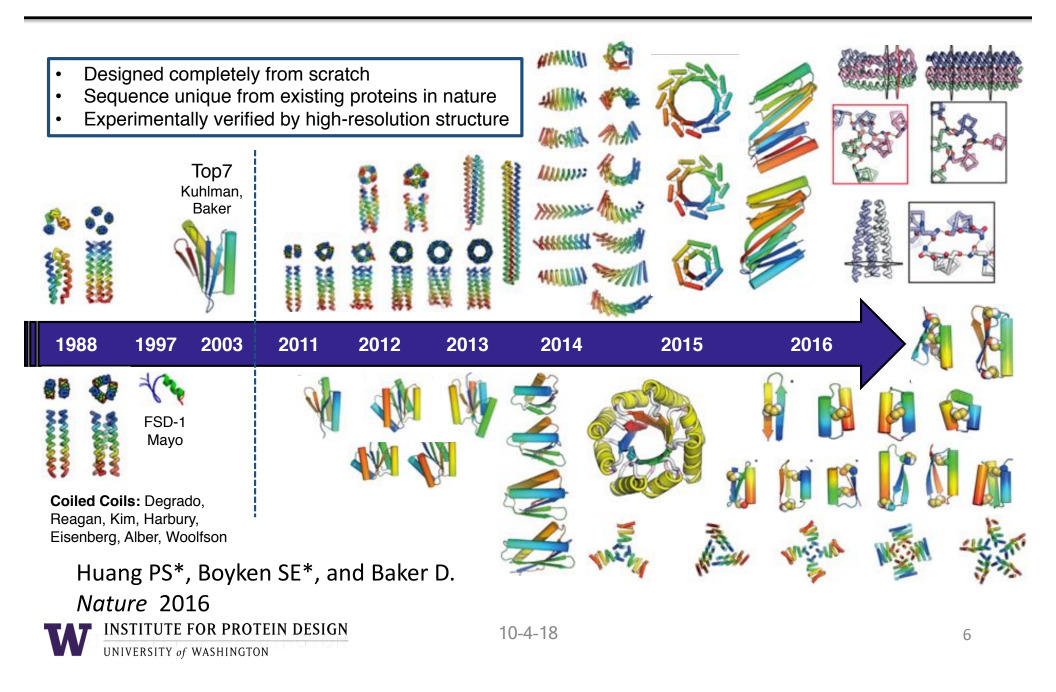
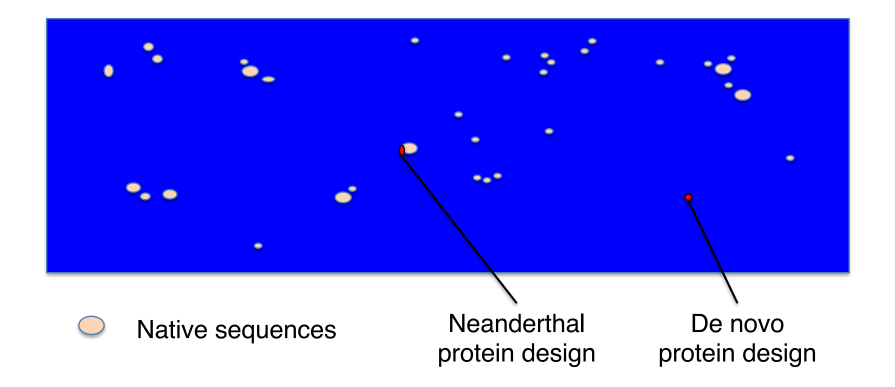


Fig. 1. A two-dimensional schematic of the target fold (hexagon, strand; square, helix; circle, other). Hydrogen bond partners are shown as purple arrows. The amino acids shown are those in the final designed (Top7) sequence.

Today: The Coming of Age of De Novo Protein Design



De novo protein design



Number of 100 residue amino sequences: $20^{100} = 1.3 \times 10^{130}$ Number of naturally occurring proteins: $\sim 10^{15}$

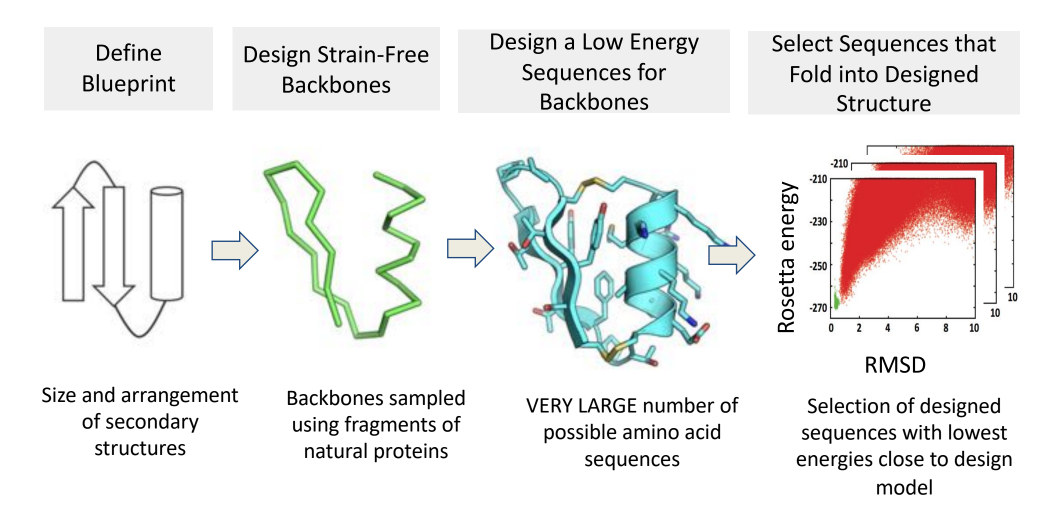
HISTORIC MOMENT IN PROTEIN DESIGN

- We've learned how to design proteins from scratch.
- There is finally enough computing power to do it.
- Genomics enables building and testing designs in the lab.

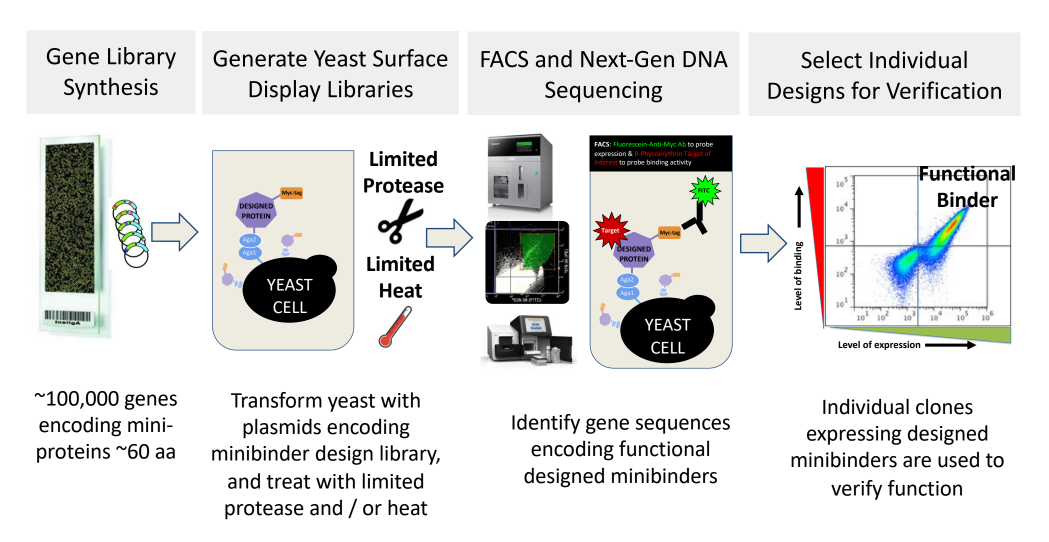




De novo protein design method

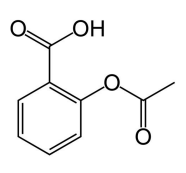


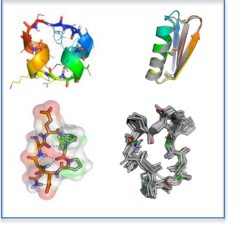
De novo protein design method

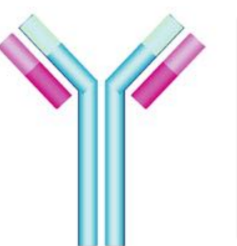


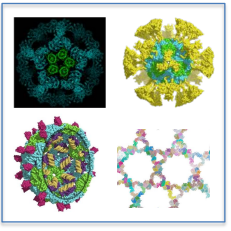
Protein Design Takes Us Beyond Traditional Small Molecules and Antibodies

• Computational design enables bottom up creation of totally new functional designer peptides, proteins, and nanomaterials.







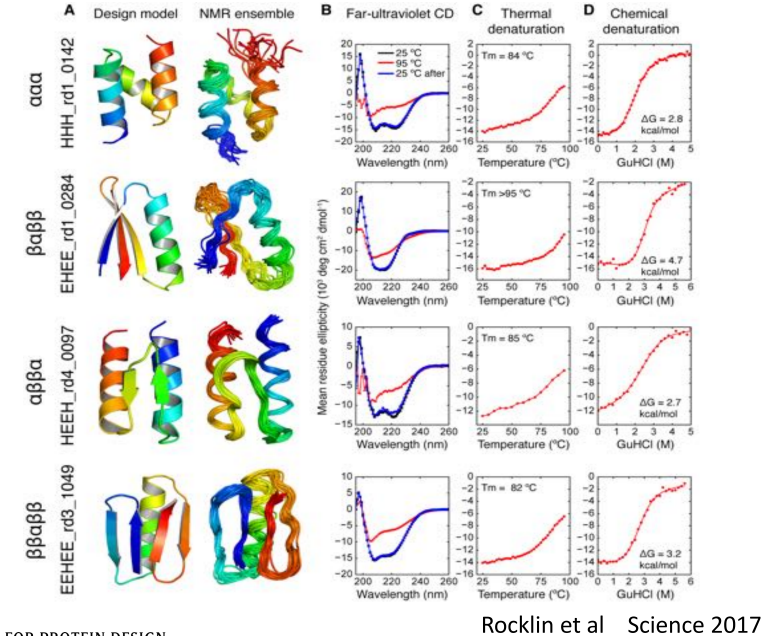


Traditional Small Molecule Drugs <1 KDa. Computationally Designed Mini-Proteins (<100 aa) Macrocycles (7-16 aa) 1-12 KDa. Traditional Antibodies or Smaller Ab Fragments 13-160 KDa.

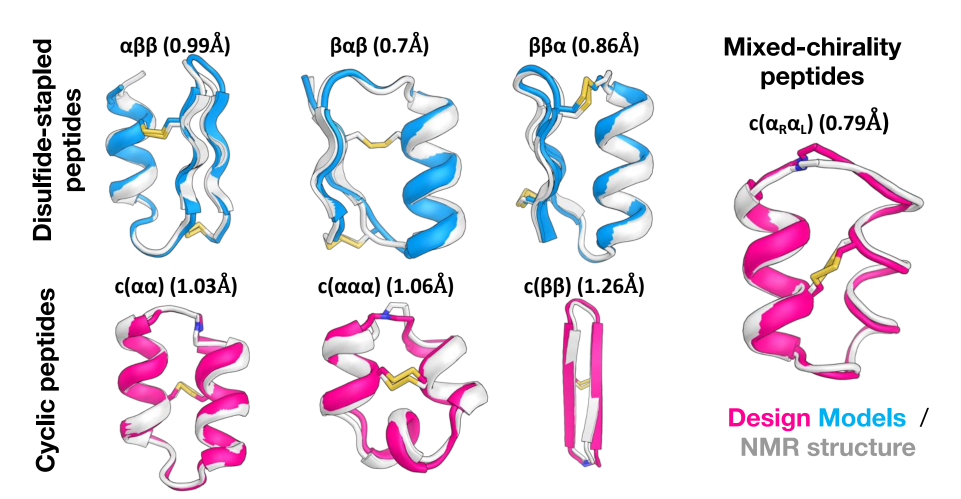
Computationally Designed Smart Nanomaterials 50 KDa. – 3 MDa.



Large Scale Design of Hyperstable Mini-Proteins



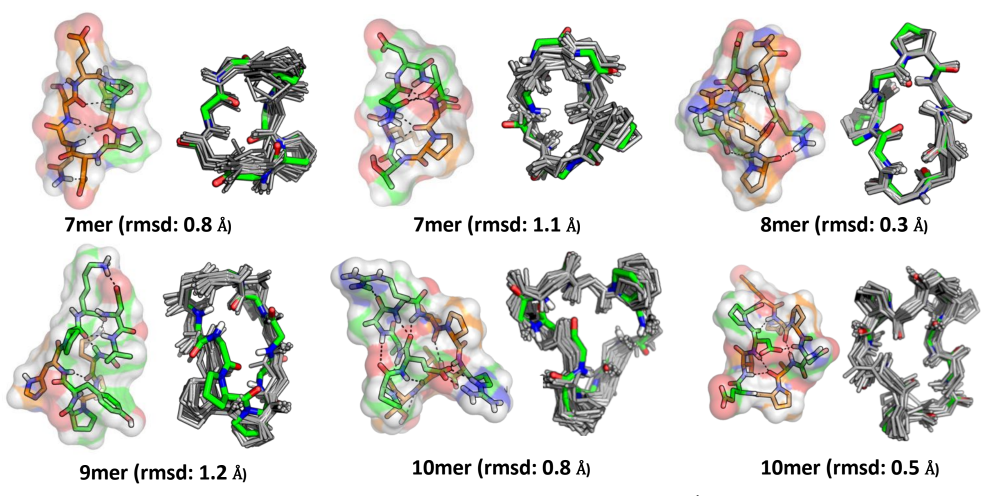
Design of Disulfide Stapled and Cyclic Mini-Proteins with Precise Control of Shape and Size



Bhardwaj, G*., Mulligan V.*, Bahl. C* et al., Nature (2016)



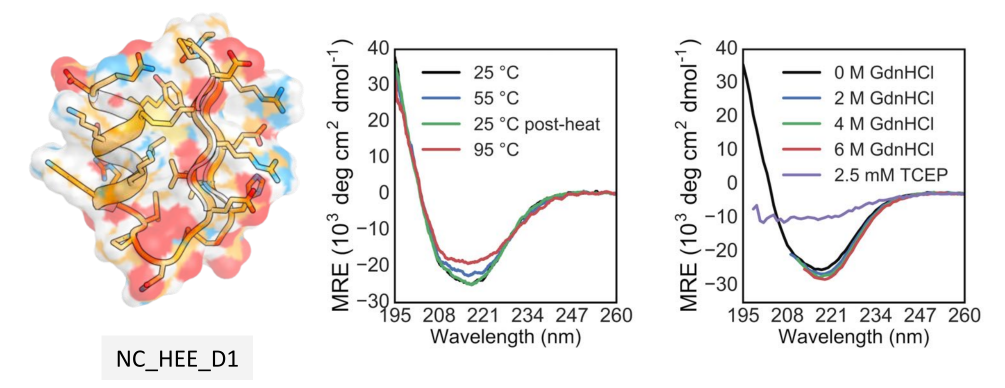
Rosetta can Design Peptide Macrocycles with Near Atomic Level Accuracy



Hosseinzadeh, P. *, Bhardwaj, G*., Mulligan V.* et al., Science (2017)

L-AA/D-AA computational model / NMR ensemble

Designed proteins show high thermal stability and resistance to chemical denaturation



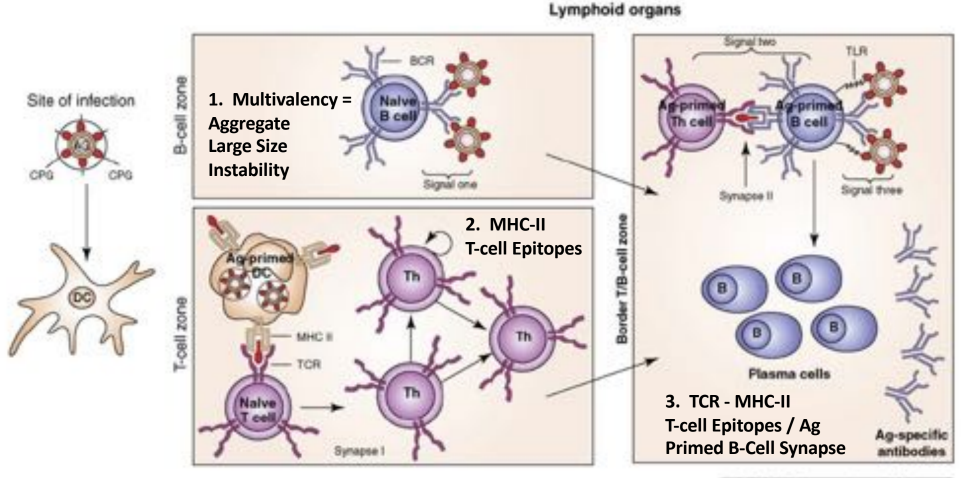
Bhardwaj, G*., Mulligan V.*, Bahl. C* et al., Nature (2016)



Immunogenicity ?



Causes of Immune Responses to Proteins



TRENDS in Pharmacological Sciences

Sauerborn M, Brinks V, Jiskoot W, Schellekens H. Immunological mechanism underlying the immune response to recombinant human protein therapeutics. Trends Pharmacol Sci. 2010 Feb;31(2):53-9.

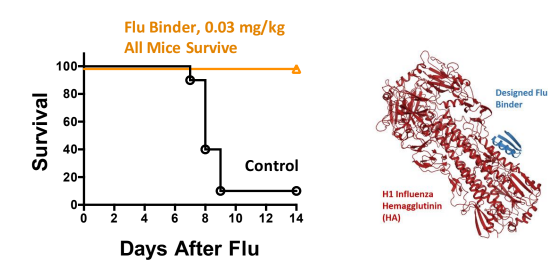


Features of Immunogenic Substances vs. *De Novo* Designed Mini-Binders

| Immunogenic Protein | Designed Mini-Binder / Macrocycle |
|---|--|
| Large size (> 10 KDa.) | Small (< 10 KDa.) |
| Multivalent = B-cell receptor crosslinking | Monomeric |
| Poor stability = Denaturation = Aggregation = Multivalent | Hyper-Stable (>80 °C, Protease resistant) |
| Not Self | Not Self |
| T-cell epitopes (MHC-II) require processing and presentation. | Hard to digest or D-handed un-natural amino acids make it hard to process. |
| Re-design (deimmunize) hard | Re-design (immune silence) easier |
| Long T1/2 resident time (weeks) | Short T1/2 (minutes to hours) |
| Delivery is often I.V. or S.C. (systemic) | Delivery options, I.V., S.C., Aerosol (localized) |
| Excipient formulations | Simple formulations (PBS) |
| TA7 INSTITUTE FOR PROTEIN DESIGN | 10-4-18 18 |

Designed Influenza Therapeutic Mini-Binder

- Low cost potent, inhalable, long-lived, broadly neutralizing anti-viral therapeutic.
- 40 amino acids (synthetic or recombinant)
- 2 disulfide bonds, $Tm > 95^{\circ}C$, $Kd > \sim 5 nM$.
- In vitro Neutralization EC50 < 0.003 ug/ml.
- Not immunogenic in mice.





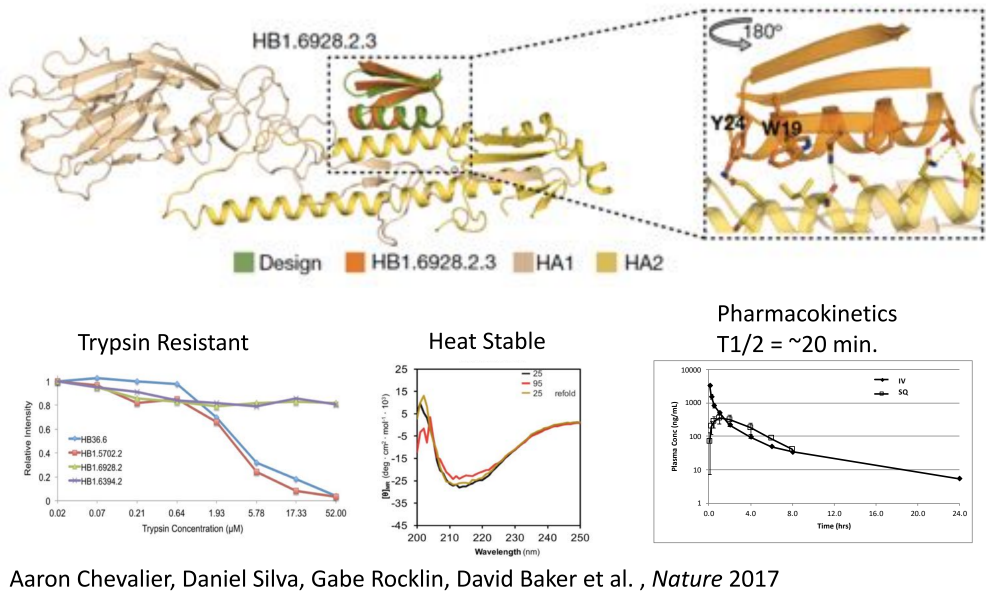
Heat Stable Aerosol



Image by IPD and Cognition Studio

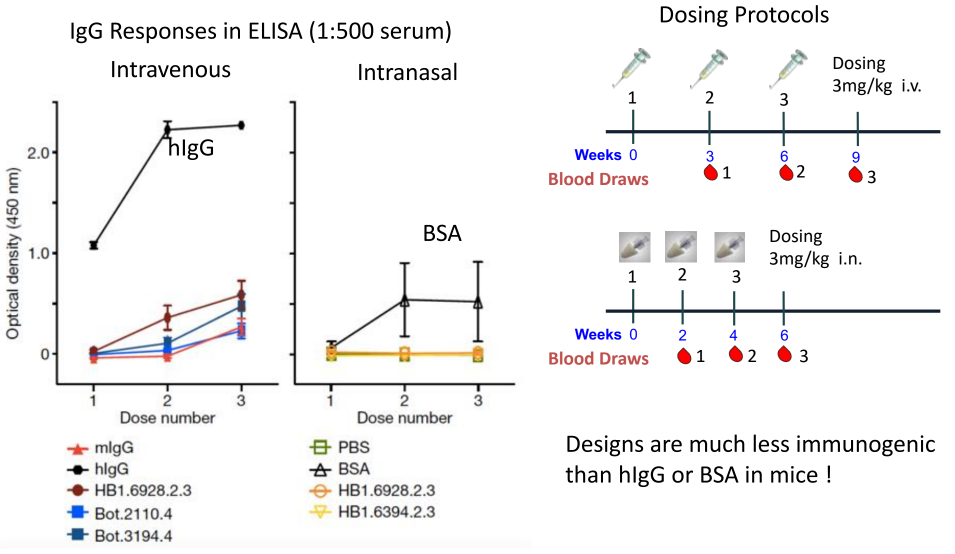
Aaron Chevalier, Daniel Silva, Gabe Rocklin, David Baker et al., Nature 2017

Potent Anti-Flu Mini-binder is Hyperstable



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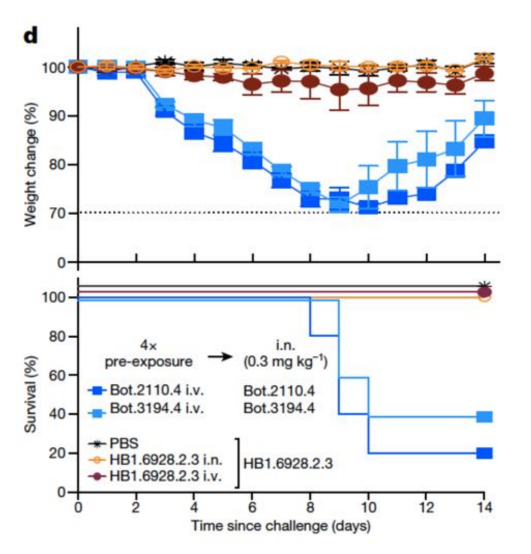
Designed Mini-Binders Elicit Little or No Antibodies in Mice

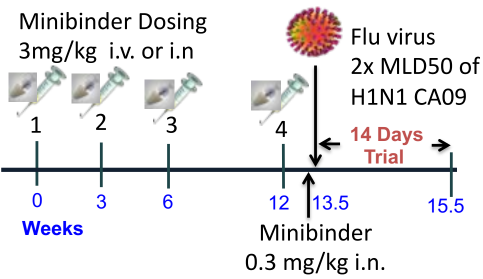


Aaron Chevalier, Daniel Silva, Gabe Rocklin, David Baker et al., Nature 2017

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Repeat Dosing of Mini-Binders Does Not Alter Prophylactic Efficacy in Mice





Complete prophylactic protection after 4 repeated doses from 8 weeks to 1 week prior to lethal flu virus challenge !

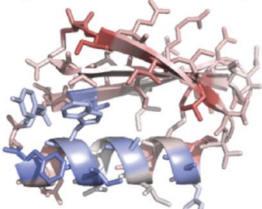
Aaron Chevalier, Daniel Silva, Gabe Rocklin, David Baker with Deb Fuller's Lab et al., Nature 2017 W INSTITUTE FOR PROTEIN DESIGN UNIVERSITY of WASHINGTON
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~Congruence Between Computational and Experimental Saturation Site Mutagenesis (SSM)

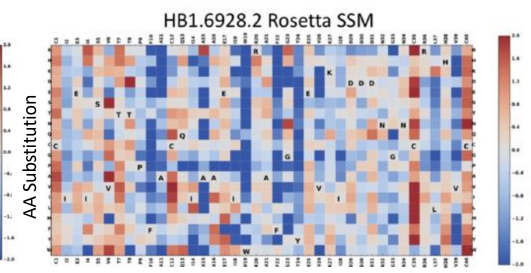
HB1.6928.2 Experimental SSM AA Substitution

Position in Designed Sequence

Experimental SSM Average

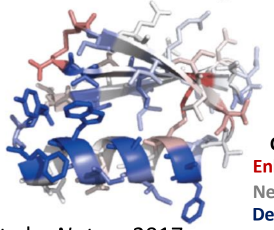


Can Re-Design Sequence to **Reduce T-Cell Epitope Liability if** Needed !



Position in Designed Sequence

Rosetta SSM Average

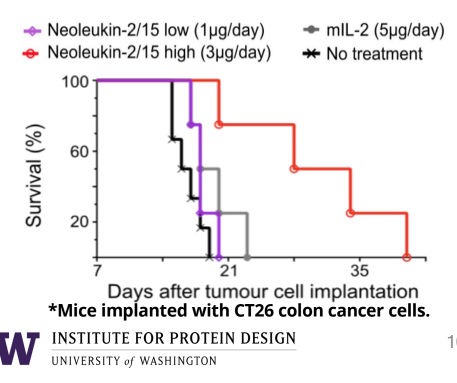


Change Effect Enhance Function Neutral **Destroy Function**

Aaron Chevalier, Daniel Silva, Gabe Rocklin, David Baker et al., Nature 2017 INSTITUTE FOR PROTEIN DESIGN

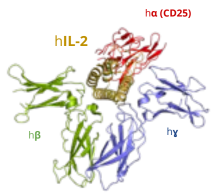
De Novo Designed Interleukin 2 (IL-2) Mimetics

- Less toxic than IL-2 •
- Easier of manufacture
- Heat stable, 90 aa mini-protein
- Wide range of immuneoncology applications
- No immunogenicity observed • in mice.

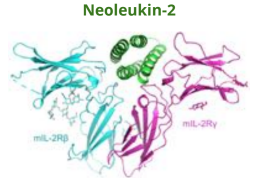




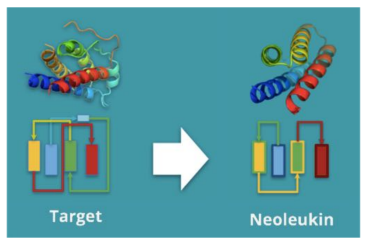
Multi-Dose Monotherapy or Combination Therapy For Challenging Cancers



Co-crystal structure of IL-2

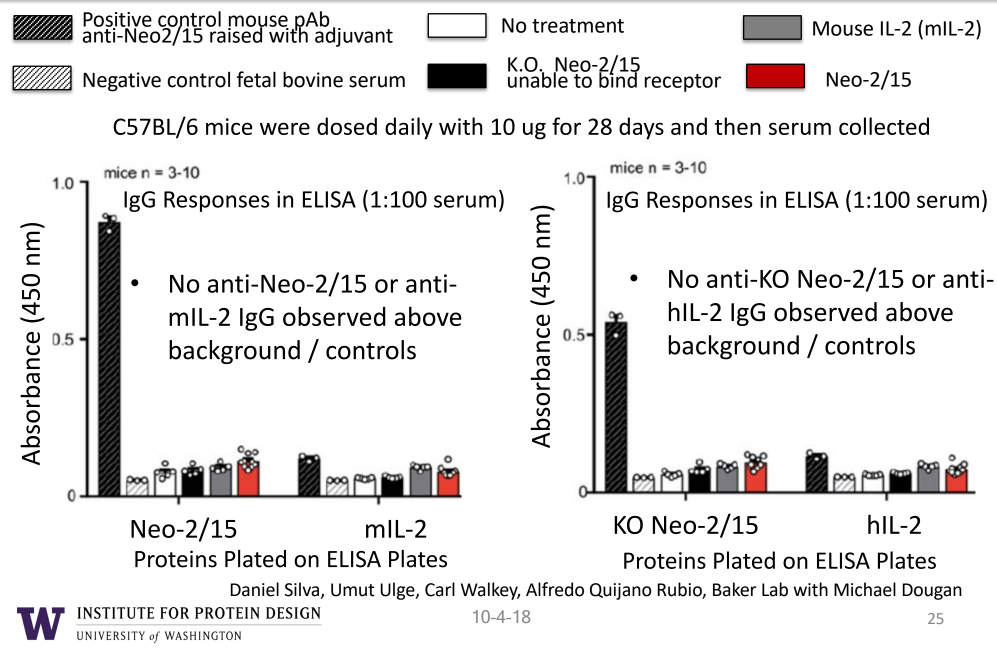


Co-crystal structure Neoleukin-2



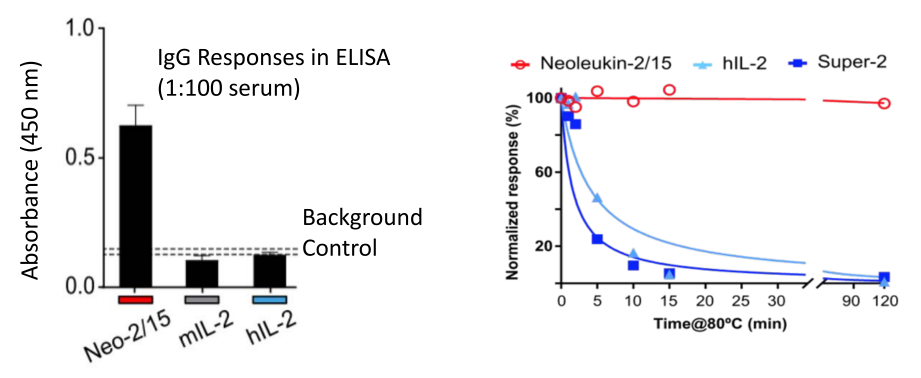
Daniel Silva, Umut Ulge, Carl Walkey, Alfredo Quijano Rubio, Baker Lab with Michael Dougan and Chris Garcia 10-4-18

De Novo Designed Interleukin 2 (IL-2) Mimetics Have No Anti-Design Antibody Response



Mouse polyclonal anti-Neo-2/15 raised with adjuvant do not react with mouse IL-2 or human IL-2

Neo-2/15 still binds hIL-2RβX receptor after Incubation @80°C



Proteins Plated on ELISA Plates

Daniel Silva, Umut Ulge, Carl Walkey, Alfredo Quijano Rubio, Baker Lab with Michael Dougan

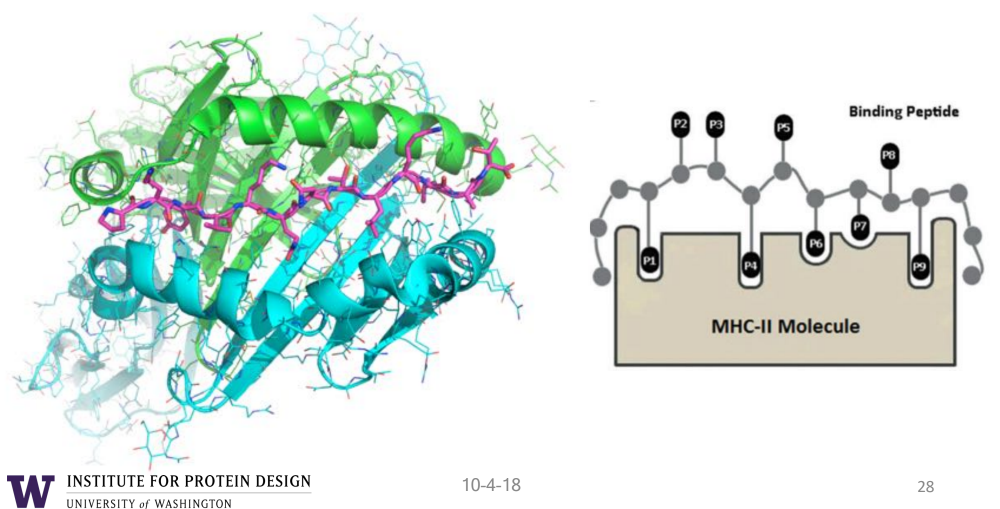


Structure and Function By Design

- Through design, we can maintain structural features through design of an astronomical number of different amino acid sequences.
- As such, numerous desired target product profile features are achievable through *de novo* design.
 - Size
 - pl
 - Stability
 - H-bonding networks = hydration sphere
 - Others.
- By definition *de novo* designed protein sequences do not exist in nature and could be recognized as foreign!
- How can we design immune silence ?

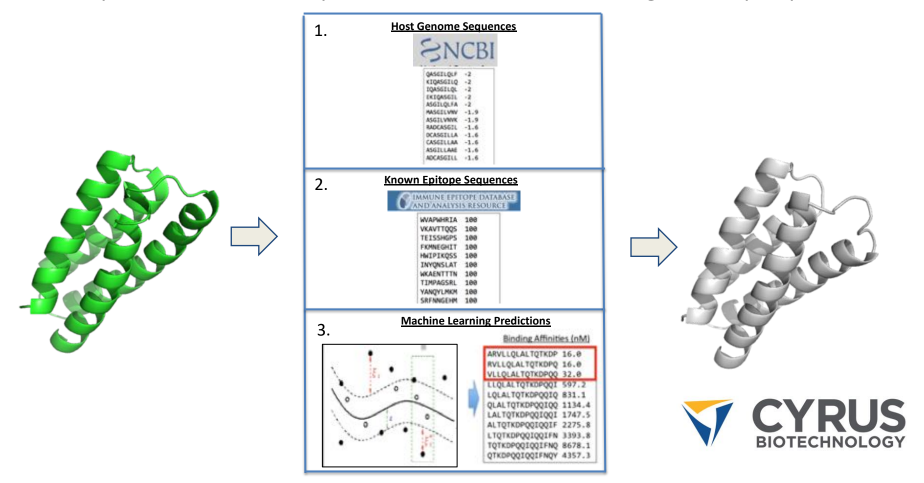
MHC-II displays peptides on the surface of cells for T-cell receptors

- Peptide binding cleft between 2 domains
- binds 15-24mers, 9mer core
- P1,P4,P6,P9 "pocket" positions



Reducing the Liability of T-cell Epitopes by Design

• Through Rosetta design, we can maintain structural features and alter the amino acid sequence to "silence" predicted or known offending T-cell epitopes.



King C, Garza EN, Mazor R, Linehan JL, Pastan I, Pepper M, Baker D. Removing T-cell epitopes with computational protein design. Proc Natl Acad Sci U S A. 2014 Jun 10;111(23):8577-82.



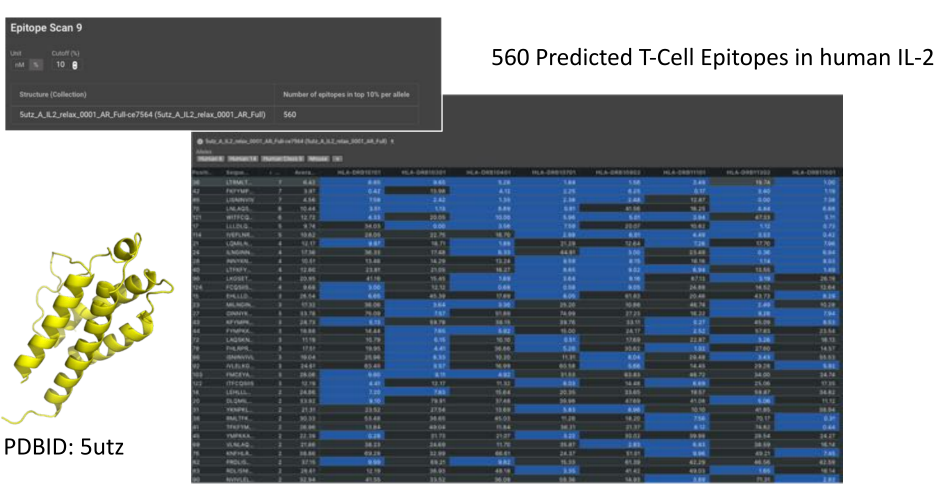
Current Immunogenicity Testing Paradigm for De Novo Designed Proteins

- **Step 1.** Since MHC-II T-cell help is key to IgM to IgG class switching and strong long lived antibody responses, we scan *de novo* designed sequence through T-cell epitope prediction software (~\$10 per protein, in compute time).
- **Step 2.** Any T-cell epitopes identified in Step 1 should be synthesized and tested in Naïve Primary T Cell Assay (e.g. Proimmune) which covers 40 human donors looks at CD4+ T-cell responses (~\$5K per peptide = expensive).
- **Step 3.** Any T-cell epitopes of Step 2 that are found to activate CD4+ T-cells should be computationally re-designed (Iterative FlexDesign with position specific score matrix and EpitopeScan Indigo King / Cyrus) while preserving structure and function of original design (~\$100 per protein, in compute time).
- **Step 4.** Make and test activity of ~10 new "T-cell epitope silenced" *de novo* designed proteins from Step 3 (\$5,000 per design). Downselect the best variant(s).
- **Step 5.** Test a limited number of *de novo* designs from Step 4 in pooled donor DC-cell presentation / T-Cell proliferation assay (\$30,000 per design, SUPER EXPENSIVE)
- Step 6. Repeat Steps 1-5 as needed. But ultimately need to move a candidate into safety / tox and into the clinic.



Caution: Not All T-cell Epitopes Are "Offensive"

- Most Current Computational Methods only "Identify" T-cell Epitopes
- Native human IL-2 has quite a few "predicted" MHC-II T-cell Epitopes



Presumably central tolerance is dealing with these epitopes 10-4-18



18.53

14.83

18.54

Considering Anti-Drug Antibodies (ADAs) Observed for Approved Biologics vs. *De Novo* Designs

- Most approved / licensed biologics are known to elicit ADAs (Stats)
- Most of these ADA's have no effect on PK / PD of biologic therapy
- Sometimes there is a dangerous auto-immune reaction (e.g. Factor VII) / Rare
- Conclusions:
 - Approved biologics aren't necessarily a good proxy for *de novo* designed proteins
 - Of the approved biologics, it is primarily the foreign (not mAb, not human protein) that are known to elicit ADAs.
 - De novo designed mini-proteins tend to be immune silent due to their stability, small size, monomeric nature, short serum half-life.
 - *De novo* designed mini-proteins can be re-designed to reduce the liability of offensive T-cell epitopes.
 - Since *de novo* designed proteins are foreign, but do not have sequence similarity to human proteins it is unlikely an ADA immune response would have an adverse effect (needs to be tested).

Acknowledgements !



Thank You !

