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Permalink
https://escholarship.org/uc/item/16109559

Journal
FEBS Journal, 285(10)

ISSN
1742-464X

Authors
Silva, DA
Stewart, L
Lam, KH
et al.

Publication Date
2018-05-01

DOI
10.1111/febs.14394

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Peer reviewed
Structures and disulfide cross-linking of *de novo* designed therapeutic mini-proteins

Daniel-Adriano Silva¹,²,†, Lance Stewart¹,²,†, Kwok-Ho Lam³, Rongsheng Jin³ and David Baker¹,²,†

1 Department of Biochemistry, University of Washington, Seattle, WA, USA
2 Institute for Protein Design, University of Washington, Seattle, WA, USA
3 Department of Physiology and Biophysics, University of California, Irvine, CA, USA

Keywords
binding; computational; *de novo*; design; disulfide; folding; protein; stability

Correspondence
D. Baker, Department of Biochemistry, University of Washington, Seattle, WA 98195, USA
E-mail: dabaker@uw.edu
†These authors contributed equally to this work.

(Received 16 December 2017, accepted 26 January 2018)
doi:10.1111/febs.14394

Introduction

Recent advances in computational protein design now enable the massively parallel *de novo* design and experimental characterization of small hyperstable binding proteins with potential therapeutic activity. By providing experimental feedback on tens of thousands of designed proteins, the design-build-test-learn pipeline provides a unique opportunity to systematically improve our understanding of protein folding and binding. Here, we review the structures of mini-protein binders in complex with Influenza hemagglutinin and Bot toxin, and illustrate in the case of disulfide bond placement how analysis of the large datasets of computational models and experimental data can be used to identify determinants of folding and binding.

Two iterations of massively parallel design-build-test-learn cycles resulted in hundreds of high-affinity designed mini-protein binders [1] against influenza H1 hemagglutinin (HA) and the receptor-binding domain of botulinum neurotoxin B (BoNT H₃B). Mini-proteins Bot.671.2 (PDB: 5VID) and HB1.6928.2.3 (PDB: 5VLI) obtained in these experiments were co-crystallized with their respective targets. For Bot.671.2 (see Fig. 1A), the asymmetric unit of the crystal contains five copies of the target H₃B, four in complex with the mini-protein binder. All the copies of Bot.671.2 match the design closely (RMSD-monomerCₐ < 1.0 Å). For HB1.6928.2.3 (see Fig. 1B), the asymmetric unit contains one copy of the molecule in complex with HA which closely recapitulates the computational design (RMSD-monomerCₐ < 1.1 Å). Deep mutational scanning [2] revealed that the mini-protein binders can tolerate a large number of mutations without negatively affecting binding; the exceptions are amino acids at the binding interface and polar amino acids in the vicinity of the binding surface (likely due to electrostatic effects [3]). In agreement with control experiments in which lower design success was obtained with simple gly/ser loops than with Rosetta designed loops, amino acids in many of the loop positions cannot be indiscriminately replaced by glycines or serines.

We studied the effect of systematically introducing all geometrically allowable disulfide bridges [4] into each of the BoNT designs (3193 combinations of disulfides for 141 initial nondisulfided designs). We found that 23% of all the pairs tested improved binding (or stability), 51% did not affect binding, and 26% were detrimental. We found a strong correlation between the number of interaction hotspot residues spanned by the disulfide and binding (see Fig. 2B). For example, ~ 64% of the designs that have average binding energy per-residue < −3.25 kcal·mol⁻¹ and whose disulfides enclose at least four hotspots improved protein...
binding or stability. Starting from Bot.728.3 (see Fig. 2A), we used this information to synthesize an improved disulfide protein variant named Bot.2110.4 (disulfide C5-C28) and co-crystallized it with H2B (PDB: 5VMR, see Fig. 2C). The experimental X-ray structure closely resembles the design model (RMSD-monomerC<1.4 Å), has a correctly formed disulfide bond, and thermal denaturation experiments show that it is hyperstable (Tm > 95 °C). Thus, even for small mini-proteins [4,5], there are many potential ways to introduce covalent staples to further enhance stability and function.

Fig. 1. Structure of de novo designed binders. (A) The co-crystal structure of the binder Bot.671.2 in complex with BoNT H2B. Top: The binders (orange color) in the context of the target (pink color). Bottom: Close-up view of binding site. One (chains G and D) of the four copies of the complex present in the asymmetric unit of the crystal is shown. (B) Same as ‘A’, for the co-crystal structure of mini-protein binder HB1.6928.2.3 in complex with HA.

Fig. 2. Disulfide stabilization of de novo designed mini-protein binders. (A) The designed structure (cartoon representation) for the binder Bot.728.3 (the nondisulfide parent of Bot.2110.4, see ‘C’). (B) Effect on function of introduced disulfides on BoNT. X-axis, the Rosetta monomer energy (kcal-mol⁻¹ per-residue). Y-axis, the number of hotspot residues between the two cysteines forming the disulfide, green, disulfide designs with improved binding (or stability), red, disulfide designs with decreased binding, grey, designs with no change in binding. Disulfide designs with lower energy and more hotspots enclosed are more likely to be improved. (C) Co-crystal structure of binder Bot.2110.4 (cartoon representation) bound to H2B (surface representation). Two copies of Bot.2110.4 in the asymmetric unit are shown in different shades of green, the computational design is in pink. The disulfide (sticks representation) is very similar in the design model and crystal structure.
Methods

Crystallographic structures are available in the RCSB protein databank (www.rcsb.org) [6]. The dataset of computational (Rosetta) and experimental metrics is publicly available in the zenodo repository (www.zenodo.org, https://doi.org/10.5281/zenodo.838815). Data analysis was performed in Python (http://www.python.org) using the iPython shell [7] and the NumPy module [8]. For the systematic scan of disulfides in BoNT binders (see Fig. 2B), a given design was considered to improve (or vice-versa) if the change in binding was at least two categories (where the possible binding categories are as follows: not binding, 100, 10, 1, and 1 nM + protease resistant).

Acknowledgements

D-AS thanks MJ Countryman for help editing this manuscript. D-AS acknowledges support from the PEW Latin-American Fellowship in the biomedical sciences and Mexico’s CONACyT. The authors acknowledge funding support from NIH NIAID grants R01 AI125704, R01 AI091823, and R21 AI123920 to RJ, and the Washington Research Foundation, the Life Science Discovery Fund grant #9598385, and the Henrietta and Aubrey Davis Endowed Professorship in Biochemistry support to DB.

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