Protein Engineering: Advances in Phage Display for Basic Science and Medical Research

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Abstract—Functional Protein Engineering became the hallmark in biomolecule manipulation in the new millennium, building on and surpassing the underlying structural DNA manipulation and recombination techniques developed and employed in the last decades of 20th century. Because of their prominence in almost all biological processes, proteins represent extremely important targets for engineering enhanced or altered properties that can lead to improvements exploitable in healthcare, medicine, research, biotechnology, and industry. Synthetic protein structures and functions can now be designed on a computer and/or evolved using molecular display or directed evolution methods in the laboratory. This review will focus on the recent trends in protein engineering and the impact of this technology on recent progress in science, cancer- and immunotherapies, with the emphasis on the current achievements in basic protein research using synthetic antibody (sABs) produced by phage display pipeline in the Kossiakoff laboratory at the University of Chicago (KossLab). Finally, engineering of the highly specific binding modules, such as variants of Streptococcal protein G with ultra-high orthogonal affinity for natural and engineered antibody scaffolds, and their possible applications as a plug-and-play platform for research and immunotherapy will be described.

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In loving memory of Lev P. Ovchinnikov, great scientist, wise mentor, and dear friend

INTRODUCTION

“The power of evolution is revealed through the diversity of life” — this is the introductory sentence of the announcement for the 2018 Nobel Prize in chemistry awarded to Frances H. Arnold for the directed evolution of enzymes, and to George P. Smith and Gregory P. Winter for the phage display of peptides and antibodies. Now, scientists have taken control of evolution and can use it for purposes that bring the greatest benefit to humankind. Enzymes produced through directed evolution are used to manufacture everything from biofuels to pharmaceuticals, and antibodies evolved using phage display combat autoimmune diseases and cure cancer [1]. Although, due to the new algorithms and advances in computer performance, novel and improved synthetic protein structures and functions can now be designed entirely in silico by rational molecular and de novo design [2], synergistic combination of the computational design and in vitro evolutionary approaches produces variants superior to those that could be generated by the design only [3]. Thus, Protein Engineering, a multidisciplinary field that organically integrates chemistry, physical and molecular biology, protein and computer sciences, has arrived to complement and succeed to Gene Engineering, one of the major achievements in Molecular Biology.

Arnold’s seminal invention of directed evolution in 1996 [4-7] resulted in the tremendous boost to enzyme-engineering possibilities well surpassing the limits of
nature. Highly artificial selection environments, diverse catalytical reactions and non-natural substrates, incorpo-
ration of synthetic amino acids and chemical modific-
atations into proteins, all these achievements have dras-
tically broadened the boundaries of potential applications for the engineered enzymes: efficient and cost-effective,
green, and sustainable biocatalysts were produced by the
directed evolution for current industrial, pharmaceutical,
and research applications [8].

Traditionally, the term “directed evolution” has been
strictly referring to the Arnold’s method of enzyme
improvement through iterative cycles of protein diversifi-
cation at the gene level (using random mutagenesis) and
high-throughput screening of the encoded variants at the
protein level. Alternatively, in phage display, the DNA
variability is introduced by the initial library of billions
of protein or peptide variants, and then it undergoes a huge
reduction in the sequential rounds of panning and ampli-
fitation, resulting in a limited number (usually, dozens or
less) of unique phage clones. In both cases, amplification
of the selected molecular variants requires an unambigu-
ous physical genotype—phenotype linkage, which is read-
ily provided by a single cell or a phage particle. As men-
tioned above, principal distinction between the phage dis-
play and directed evolution is that DNA does not under-
go further diversification between the rounds of phage
display panning. Therefore, molecular evolution achieved
in phage display by the sequential enrichment of the
most-fitted variants is rather reminiscent of the genetic
bottleneck effect in nature, unlike the gradual canonical
directed evolution which, step by step, selects more and
more evolutionary advanced molecules. By its very
nature, based on molecular selection by the phage bind-
ing, the phage display technology cannot address modu-
lations and tuning of enzyme catalysis and is usually
directed toward enhancement of the structural stability of
the protein and binding affinities to antigens, partner pro-
teins, or ligands, demonstrating its tremendous power
particularly for specific peptide and antibody generation
and maturation.

Since Smith and Winter published their pivotal stud-
ies in 1980s-1990s [9-14], the scope and capabilities of
phage display have been enormously expanded [15-20].
Construction of large numbers of phage display peptide
and antibody libraries containing 10^{11-12} unique members
have been achieved [18, 21, 22] and are commercially
available (New England BioLabs, MoBiTec GmbH, and
Creative Biolabs, Shirley, NY, USA)

Synthetic antibodies, tailored to a specific antigen or
antigen epitope hugely helped to overcome protein struc-
ture determination challenges as well as powered up many
medical endeavors. Multiple modifications of the original
technology of peptide phage display, including cyclic and
artificially linked peptides [23], resulted in varieties of the
cancer-specific ligands validated in cancer diagnostics
and therapy [24].

Undoubtedly, the progress in protein engineering
greatly affects the landscape of cancer- and immunothe-
reapeutics beyond the production of new potent peptide-
based ligands or synthetic and in vitro matured antibodies.
Applications of the antibody-drug conjugates, bispecific
antibody-based molecules, and other antibody derivatives
are empowered by the highly discriminative antibody-
driven targeted cell delivery and well-controlled effect on
the cells of choice. The targeted cell type is defined by the
antibody specificity to one of the cell–surface receptors,
and its fate — by the built-in effector moiety. The anti-
body-drug conjugates are able to kill cancer cells by bind-
ing to the tumor-specific receptors and directionally dis-
patching their “payloads” — potent cytotoxic small mole-
cules [25]. Cytolytic effect of the bispecific T-cell
engagers (BiTEs) depends on their ability to induce
synapse formation between the cancer and T cells by
bridging together their receptors targeted by two specifici-
ties [26, 27]. In addition of being a part of anti-cancer
therapy, the antibody-based reagents could specifically
inhibit inflammatory pathways and have been developed to
treat diseases and conditions caused by excessive inflam-
mation and autoimmunity [28, 29]. To date, hundreds of
different antibody-based formats have been engineered for
therapeutic purposes, with many representing BiTEs [30],
and new constructs are constantly emerging [31].

Here, we review the antibody phage-display technol-
ogy in detail – from the library construction methods to
synthetic antibody (sAB) production and characteriza-
tion. Using the phage-display pipeline established in the
Dr. Kossiakoff Laboratory (KossLab) at the University of
Chicago, as an example, the state-of-the-art progress in
sABs custom-tailoring for challenging applications, such
as protein structure determination of membrane proteins
is addressed. Then, the sABs high potency as energy and
functional probes, conformational and crystallization
chaperons, fiducial markers for single-particle cryo-elec-
tron microscopy (cryo-EM), and superb antigen-binding
reagents is discussed. Next, reciprocal maturation of the
uniquely specific affinity between the protein G (variant
GA1) and the engineered sAB scaffold (Fab^{LRT}) aiming
further higher-order construction efforts, such as multi-
valent and multi-specific antibody arrangements, is
described. The resulting novel, easily manipulatable,
multifunctional, and sturdy GA1-based plug-and-play
platform utilizing Fab^{LRT} molecules as the interchange-
able elements, and its possible applications in basic sci-
ence, biotechnology, and medicine are considered in the
conclusion.

ANTIBODY PHAGE DISPLAY
ANTIBODY FORMATS

Immunoglobulin G (IgG) is the most common type of
human antibodies. IgGs, which are complex, bi-valent,
multi-subunit proteins with many disulfide bonds and are unsuitable for bacterial expression. Therefore, smaller, mono-valent antigen-binding fragments such as scFv (single chain fragment variable) or Fab (fragment antigen-binding) are the most often used formats for phage display [32] (Fig. 1a). Fab molecule is a heterodimer containing a full-length light chain (Fab) and an Fd fragment of a heavy chain (VH and CH1 domains) usually linked together by a disulfide bond at their C-termini (this linkage is not essential for heterodimerization [33]. These chains, of approximately equal masses, comprise \( \sim 50 \text{ kDa} \) Fab. The scFv molecule is half a size (\( \sim 25 \text{ kDa} \)) and consists of two variable domains (VH and VL) fused together by a small flexible peptide linker of \( \sim 15 \) aa (Fig. 1a) Notably, while Fabs generally maintain full antigen-binding affinity upon reformattting into the full-size IgG and back (taking into account the 2-fold difference in the format valency), the scFv affinity can be drastically changed upon reformaattting and sometimes is influenced by the VH/VL domain mutual orientation and/or linker length, and, also, this format is prone to oligomerization [34, 35].

Both scFvs and Fabs have been displayed on the bacteriophage surface in the Winter’s pioneering works [24, 36] and are still the most popular for construction of sAB phage display libraries. However, combinatorial arrays [37] and some other new formats, such as nanobodies representing a single VH chain domain of an unusual homo-dimeric camelid IgG [38] (Fig. 1b), are successfully challenging the originals. A dozen of therapeutic antibodies have been commercialized that have been discovered by means of antibody-displaying phage libraries including the world best-selling rheumatoid arthritis treating antibody adalimumab [39], with many more in the pipeline [40].

In addition to phage display, a great variety of display platforms have been developed to better suit rationale and particularity of the selections. Large libraries of recombinant cell-surface proteins were constructed and successfully applied for display on the baculovirus and bacterial, yeast, insect, and mammalian cells [41-44]. Molecular displays on the surface of eukaryotic cells are highly permissive for multi-domain proteins and multi-molecular complexes, as well as for synthesis of functional mammalian proteins undergoing essential posttranslational modifications that are supported by the eukaryotic cytoplasm [45]. The alternative cell-free methods, like ribosome display [46] and mRNA display [47] offer their own advantages, such as speedy turnover, enormous library size, and potential incorporation of non-natural amino acids and chemical modifications into the proteins. However, despite of all the advantageous features of other display techniques mentioned above, phage display remains the most popular display platform today.

**M13 BACTERIOPHAGE**

This chapter is dedicated to M13 bacteriophage structure and physiology that have been studied for decades and can be skipped by the educated readers, however, it contains information important for better understanding of the M13 phage display design specifics, advantages, and limitations.

Bacteriophages M13, fd, and f1 belonging to Ff (F-pilus specific filamentous) phages and almost identical in
display libraries can be stored as frozen intact bacterial cell (although growth rate of the infected lytic: the progeny phage filaments are secreted from the infectious phage particles. M13 bacteriophage is non-lytic: the progeny phage replicative form (RF) and producing circular formable M13-based dsDNA phagemid vector, mimicking to accommodate large foreign DNA as a highly trans-display platform to date.

M13 phage has high capacity for replication and can accommodate large foreign DNA as a highly transformable M13-based dsDNA phagemid vector, mimicking phage replicative form (RF) and producing circular ssDNA phagemid molecules that can be packaged into infectious phage particles. M13 bacteriophage is non-lytic: the progeny phage filaments are secreted from the intact bacterial cell (although growth rate of the infected bacteria slows down). Consequently, M13-based phage display libraries can be stored as frozen E. coli stocks. The extraordinary stability of M13 bacteriophage particles in a wide range of temperature, pH, and solvent composition allows for prolonged incubations under various conditions of the phage-displayed antibody libraries during the experimental procedures, and the library storage in frozen or desiccated state for years [50].

The single-stranded circular phage M13 genomic DNA (6400 nt) encoding 11 phage-specific proteins in total is protected by the stable protein capsid composed of approximately 2,700 copies of the major coat protein gp8 forming the cylinder, and several copies of four other minor coat proteins at either end of the phage filament: gp3 (five copies) and p6 at one end; gp7 and gp9 at the other. The infectious M13 phage filament has a length of ∼1 μm and a diameter of ∼7 nm [51].

The first step of M13 infection involves recognition of the F pilus on the surface of the E. coli by gp3, followed by gp3 binding to the tolA-based bacterial complex that bridges between the outer and inner membrane of the host [32]. After penetration through the periplasm the phage genomic ssDNA is got stripped of gp8 and released into the host cytoplasm, where it is rapidly converted to a RF dsDNA by E. coli enzymes. This initiates transcription from 5 constitutive promoters, producing mRNAs encoding all 11 phage proteins. Hundreds of the single-stranded copies of the M13 genome are made by the mechanism called rolling-circle replication using RF DNA as a template: some of them are converted into more RF molecules, while the majority are destined to be packaged into the phage progeny. As freshly synthesized phage proteins accumulate, ∼750 gp5 dimers cooperatively cover the whole length of M13 circular ssDNA except for a short 77 base-long section, known as the packaging sequence. This gp5 shell packs genomic DNA into a flexible rod protected from nuclease, thus forming the intracellular precursor of the extracellular virion. Binding of the DNA packaging-signal at the forward end of the precursor by the assembly protein complex in the inner membrane initiates the phage extrusion. Next, the membrane-associated gp8 oligomers are built into a spiraling array around phage DNA, replacing gp5. This nascent filament passes through the periplasm-spanning channel and a porin-like structure composed of 14 sub-units of gp4. Finally, the inner membrane inserted gp3 and gp6 proteins are attached to the proximal end of the extruding virion particle completing the phage assembly.

The details of M13 phage replication and assembly are reflected in a vector construction for displaying of foreign polypeptides as coat-protein fusions on the phage. As a rule, the gene of interest is fused to the gene of one of the coat proteins cloned into the plasmid, which carries both plasmid dsDNA and f1 ssDNA origins of replication, f1 DNA packaging signal, and an antibiotic resistance gene [52]. This vector, called a phagemid, can be transformed and amplified in E. coli as a regular plasmid. Since this phagemid mimics RF of the phage DNA, the phagemid-transformed F pilus + cells can efficiently produce ss phagemid DNA and encapsidate it into the infectious phage progeny with the assistance of the M13 helper phage infection that provides all the phage proteins required for phage biogenesis. A mutation, that makes f1 DNA packaging signal of helper phage defective is disadvantageous for packaging of the helper-phage ssDNA, and thus favors formation of the phagemid-containing virions. Since these virions display the phagemid-encoded fusion protein, it provides physical link between the phage genotype and phenotype, realizing the key principle for phage-display as a molecular evolution-directed system [53]. In order to eliminate the need for the helper phage and avoid the helper-phage contamination in the phage progeny, bacterial “packaging” cell lines containing M13-phage based helper plasmids lacking the ssDNA replication origin, have been constructed [54], however, the classic way featuring the helper-phage superinfection is still very popular.

By now, every M13 coat protein has been tried for molecular display as both N- and C-terminal fusions [55]. Nature of the coat protein to be used as the fusion partner as well as size and structural characteristics of the proteins chosen to be displayed should be considered for the display success. As a rule, large, structurally complex, and multi-subunit proteins, like antibodies, cannot be expressed in bacterial cells, and this automatically makes them inappropriate for bacteriophage display. Additionally, fusion proteins could be rejected after the synthesis at different stages of phage maturation. Such obstacles as
steric hindrance or wrong surface texture could prevent some bulky, or highly charged, or overly hydrophobic fusion proteins from proper incorporation into the phage particles both prior to phage assembly — upon membrane association, or during recognition by the assembly complex, or after the capsid completion — at the phage extrusion step due to insufficiently wide gp4 channel opening for the particle decorated with a fully folded and assembled fusion proteins. While the major coat protein gp8 supports high multivalency of the display and is frequently used for the display of peptide libraries, even antibody fragments are too large for gp8 display. Historically, gp3 was the first display support that was successfully demonstrated to work in the antibody libraries displaying scFvs or Fabs [24, 36] and is the most commonly used today. To be displayed, usually the antibody fragment is fused to the N-termini of gp3 or its C-terminal domain, which itself assembles into a capsid tip, while it is the N-terminal domain of gp3, which recognizes the E. coli F-pilus. The wild-type gp3, product of the copy of unmodified g3 in the helper-phage DNA, is incorporated into the pentameric gp3 arrays at a higher proportion over the fusion protein, thus ensuring infectivity of the progeny virions [56, 57]. Moreover, the ratio of the fusion to wild type gp3 available for the phage assembly is such, that not every progeny particle displays the antibody. However, this is not usually problematic for the library performance, since the standard antibody phage library has a density of 10^{13} cfu/ml, which is 100 times greater than the typical antibody libraries.

**M13 PHAGE ANTIBODY LIBRARIES**

Natural antibody libraries. Antibody-displaying phage libraries can be either natural or synthetic, or both, depending on the origin of the diversity component, incorporated into the antibody-fragment scaffold. Diversity of the natural libraries is provided by the antibody repertoire acquired in the live immune system by gene rearrangement [12]. Each of the three major human immunoglobulin gene loci (one for HC and two — lambda and kappa, for LC, light chain), contains multiples of variable (V), diversity (D, only for HC) and joining (J) genes. V(D)J recombination, different in each individual lymphocyte, causes one random copy of each type of the gene segment to be sequentially joined together, generating 1x10^{11} unique heavy-chain sequences and an enormous potential number of naïve paired antibody perhaps in the range of 10^{16}-10^{18} unique antibody sequences [58]. Real diversity of the naïve human antibody repertoire is estimated to be at least 10^{12} unique combinations. Since the number of peripheral blood B cells in a healthy adult human is on the order of 5x10^{9}, the circulating B cell population samples only a microscopic fraction of this diversity.

Antibody diversity is, mostly, present in their six complementarity determining regions (CDRs), highly variable structural loops undergoing clonal selection in the immune system. Three of them: L1, L2, and L3 are located in VL, and three: H1, H2, H3 — in VH. In general, the CDR loops in HC are more frequently involved in antigen binding than those in the LC. Contribution of each of the six CDR loops to antigen recognition is individual and each residue position within a single CDR may play a completely different role depending on the antigen—antibody complex [59]. Five of the six CDRs of a restricted variety and length from 3 to 15 residues adopt a limited number of backbone conformations, while H3 is longer, more compositionally and structurally diverse [60] and is considered to be the most important for antigen recognition [61].

Standard natural combinatorial antibody libraries are constructed by random pairing of VH and VL repertoires from cDNA of the pool of mammalian B cells into a phagemid vector using PCR cloning. The antibody assortment can be enriched for the desired-target binders by prior immunization. The resulting immune library does not need to be as large as naïve libraries, since the immunized pool of lymphocytes would contain multiple proliferated B-cell clones targeting the antigen. Some naïve antibody libraries represent 10^{11} individual clones or more [18, 21, 22], reaching practical limit of the phage antibody library size preset around 10^{12} value, due to the technical limitation of bacterial transformation, culturing, and storage [62]. While theoretical diversity of the naïve combinatorial libraries is much higher than their achievable size, there is significantly less variety of the naturally paired VH-VLs in the mammalian immune system. A miniature phage library containing ~10^{9} naturally paired scFvs derived from cDNA of the individual B cells was constructed using a single-cell microfluidics that allowed to mine for an extremely rare antibody variants as well as to display natural antibody paratopes from hundreds of donors in one phage library [63]. This technique was advanced to high-throughput single-cell activity-based screening and sequencing platform of thousands of antibodies from the mice immune cells and activated human memory cells [64]. At present, there are dozens of commercially available natural antibody libraries in multiple formats, including chicken scFvs and camelid VHVs (Creative Biolabs).

Natural antibody libraries have been proven to be excellent suppliers of high-quality antibodies suitable for therapeutic applications in medicine [65] and veterinary [66], and many such antibodies are in clinical use or at various stages of therapy development [40].

Synthetic antibody libraries. Most of the sAb libraries use the natural Ig framework, however, a number of modules representing protein-binding polypeptides from natural sources were engineered into the stable spatial arrangements of several diversified loops fea-
turing compact antigen-binding structures [67]. When implemented in the phage display libraries, they yielded high-quality non-Ig binders that could be of great interest for basic and applied scientific research due to their robust folding, high solubility, and small size. A number of non-Ig binders for health-relevant targets are, presently, at different stages of clinical trials [68]. Nonetheless, despite the constant challenge from other emerging formats, the classic libraries based on the Ig-fragment scaffolds remained the most desirable for sAb production [69].

The CDR loops of a sAb library need to be designed so that the resulting library is enriched with diverse, yet nature-like sequences, different in length and composition [70]. As discussed above, the pre-existing variety of CDRs incorporated into the natural antibody libraries originates from the operational immune system in vivo [12], therefore, quality of the natural library depends, mostly, on the efficiency of the variety incorporation into the library, i.e., on the library size. On the contrary, theoretical number of the possible combinations in the case of synthetic CDR variants has almost no upper limit: e.g., complete randomization of just a single piece of 10 codons produces more than $10^{13}$ (20$^{10}$ including stop codons) of clone variations — amount, practically unattainable in a phage library. Therefore, smart diversification strategy considering each CDR length variability and randomization method as well as their magnitude became the key issue in the design of a quality sAb library [69].

The early examples of sAb libraries incorporated random CDR sequences of different length by means of degenerate oligonucleotides synthesizes from different nucleotide mixtures at different codon positions [71]. Due to genetic code redundancy, all 20 amino acids are encoded by NNK or NNS degenerate codon (N is any of the four deoxyribonucleotides; K = G or T; S = G or C), removing two out of three stop codons (TGA and TAA, but not TAG) and reducing the number of possible codons from 64 to 32, thus, decreasing the number of unique DNA sequences by two-fold. Other combinations of nucleotide mixtures can further reduce the number of codons and encoded amino acids in the sets. Examples of the commonly used degenerate codons in CDR design include KMT (M = A or C) that encodes Ala, Asp, Ser or Tyr; WMC (W = A or T) for Asn, Ser, Thr, or Tyr; and RRT (R = A or G) for Asn, Asp, Gly, or Ser. Degenerate oligonucleotides are easy to design and are cost-efficient — highly functional antibody libraries of $10^8$-$10^9$ unique clones have been constructed using this method [72]. Although the smaller nucleotide sets are highly desirable due to the library size limitations, there are many mutually exclusive amino acids in these sets. This problem is resolved by the more expensive but fully codon-specific synthetic methods, such as trinucleotide phosphoramidite synthesis [73] or dsDNA triplet-based Slonomics [74], dictating incorporation of only the chosen codons at the desired ratio into each position. Consequently, CDRs can be designed to have a smaller number of variants, while featuring more nature-like distribution of amino acids, thus making the sABs to better resemble natural antibodies and improving the clone-representing potential of the library. Aiming for the library size $\leq10^{10}$ unique CDR combinations can significantly improve the clone coverage and sampling power of the phage display.

As was stated above, theoretical size of the library depends on its design: the number of CDRs touched, the length range of the CDR loops and quality, and quantity of the codon-randomization events in each CDR. Following the natural antibody tendency favoring the highest variability and paratope involvement in L3 and H3 CDRs, most sAB libraries that have only H3 and L3 diversified and just one of the canonical natural sequences incorporated in place of each of other four CDRs, proved to be very efficient [73, 75, 76]; however, some high-quality libraries have all six CDRs diversified [72, 77, 78]. Surprisingly, the minimalist libraries, containing sequences of only two residues, Tyr and Ser, in their CDRs, were proven highly effective in generating specific antibodies against a wide array of antigens providing a rationale for high abundance of these two residues in the CDRs of the natural immune repertoire [62, 79]. Due to the simpler design, these libraries are relatively easy to construct, and, due to the smaller theoretical size, they possess higher sampling power. Involvement of more of CDRs and CDR positions into deeper diversification, while producing an exponentially increasing number of possible variants, allows to build more enhanced and sterically extended paratopes with higher conformational diversity, thus, facilitating efficient recognition of various epitopes of different types of antigens.

As was mentioned above, natural antibody libraries have a huge advantage over any sAb library since they contain only expressible and stable CDR variants that have passed multiple control points of the immune system. In addition, chemical synthesis of CDRs is inherently compromised by synthetic errors, introducing all sorts of mutations that generate unwanted amino-acid substitutions and stop codons resulting in nonfunctional antibody clones [72, 77] To eliminate poorly expressed clones and non-productive frame shifts, a proofreading step, such as selection of ampicillin-resistant sAB — β-lactamase genetic fusions, has been added as the final step of library construction [77].

Another important aspect defining functionality of the sAB phage display is a balance between the theoretical number of possible variants set by the design, and the number of unique phage clones achieved at a construction step, i.e., the library sampling power. Clone under-representation in a library can also be caused by clone-
specific impediments, like slower replication or assembly rates, reducing or even eliminating the pool of most troubled variants. The higher-order diversification design may account for such astronomical calculated number of the unique variants that the realistically limited of manageable phage library variety of $10^{12}$ clones rationalized above would constitute only a microscopic fraction of the theoretical prospects. Due to this problem, more extensive and deeper diversification would not allow for selection of the ideal antibody variants, covered by the design but not even present during the phage panning, however, there would be a greater variety of the second- and third-best clones, suitable for the task. sAB phage libraries of this kind have produced a wide range of high-quality antibodies for a broad spectrum of antigens, although their theoretical variety is hugely under-sampled. The well-designed sAB libraries have several distinct advantages, which include optimal variability, size, clone representation and density of the display, high sAB expression, solubility and stability, and ease of further engineering and optimization.

THE KOSSLAB PIPELINE

**Synthetic-antibody library construction.** The KossLab pipeline have been constructed based on the scaffold from a humanized Fab 4D5 Herceptin framework (here, Fab)$^3$ engineered for high stability and efficient phage display. Combination of the mixed nucleotide and trinucleotide phosphoramidite based methods were used for synthesis of randomized DNA oligonucleotides that were next cloned into the Fab framework using Kunkel mutagenesis method [80].

In the library, limited amino acid diversity is introduced into all three HC CDRs (H1, H2, and H3), as well as the third CDR of LC: L3, while L1 and L2 have fixed canonical loops: SVSSA and SASSLYS, respectively. L3 length varies from four to six residues. In H3, the loop lengths range from 6 and 20 residues (sub-library A), 7 and 15 residues (sub-library B) or 6 and 17 residues (sub-libraries C and D). H3 design in the most complex sub-library D (6 to 17 residues) was based on a tailored oligonucleotide synthesis strategy that biased the sequences in favor of Y, S and G but allowed for 19 of the 20 genetically encoded amino acids (only C was excluded) and potentiated high level of diversity amongst the H3 loops in terms of both length and sequence. The pooled library consists of $10^{10}$ unique variants, huge under-sampling of the theoretical diversity (by more than 20 orders of magnitude) [62]. Extensive exploitation of the library over the years in hundreds of successful selections resulted in multitudes of high-affinity antibodies with dissociation constants in the sub-nanomolar to single-digit nanomolar range against a wide variety of protein antigens [81-100] confirming that this fully synthetic library with the limited diversification has essentially recapitulated capacity of the natural immune system.

**Target immobilization and phage elution.** Production and immobilization of the target-protein in an accessible and native functional state is the next critical step in the selection-project progress after obtaining high-quality phage-displayed sAB library. The original and still frequently used way of antigen immobilization for biopanning is passive protein adsorption on the activated plastic surfaces in random orientation through multiple non-specific non-covalent interactions [101]. However, adsorption is usually accompanied by some degree of structural change of the antigen, and there is a possibility that the antibodies isolated against the passively adsorbed antigen may not recognize the free native antigen in solution. Nevertheless, passive immobilization is appealing due to simplicity of the method and frequent success of selections that yield clones that recognize and bind functional protein.

At present, the most common immobilization method avoiding adsorption-induced conformational changes of the antigen is based on the extremely tight biotin–streptavidin interactions [102] that are widely used in various research and biotechnology applications requiring tight and specific intermolecular interactions (detection and isolation of proteins, nucleic acids and lipids, protein purification, new-generation DNA-sequencing, mass-spectrometry based proteomics and many others).

Protein antigen could be chemically conjugated with biotin [103] using commercially available biotinylation reagents targeting variety of specific functional groups or residues, including primary amines, sulfhydryls, carboxyls, and carbohydrates [104]. Chemical biotinylation can result in multiplicity of protein modification spots and masking or altering natural epitopes of the antigens by biotin moieties. Another, the least structure-disturbing way for biotin attachment, is through a genetic linkage of the antigen to Avi-tag, a peptide (GLNDIFEAQKIEWHE) specifically recognized and efficiently biotinylated by the biotin ligase BirA. BirA could modify Avi-tagged proteins in vitro, after purification [105], or in vivo [106], during expression in *E. coli* bearing BirA plasmid; in every case high-level of antigen biotinylation has been demonstrated (70-90%). This modification level is sufficient for efficient capture of the antigen on the streptavidin–coated microplates or streptavidin-coupled magnetic beads followed by the library panning.

There are other affinity tags options for tethering proteins in a relatively uniform orientation with minimal disruption of native conformation [107] including metal ion-[108, 109,] antibody- [110], and protein—ligand capture technologies [111] that have been used for the target immobilization. Depending on the antigen immobilization technique, elution methods also vary widely. The most-common eluants are diluted solutions of HCl or
Glycine (adjusted to pH 2–3) that disrupt molecular interactions between the antibody displayed and the antigen immobilized (and the biotin–streptavidin complex too). Although there is almost no risk of eluting phage incompletely, the prolonged exposures to low pH could affect phage infectibility. If this becomes a problem, a gentle and highly specific alternative would be competitive elution with the antigen excess. This mostly releases the target-bound phage limiting selection of the non-specific clones [112, 113], but requires additional amounts of the purified antigen. The most common affinity-independent elution method entails insertion of a unique proteolytic cleavage site next to the affinity tag [114]. Cutting-off the affinity elution method would trigger release of the target–bound phage exclusively, enhancing specificity of the selected phage clones compared to the competitive elution. In addition, phage elution of any kind can be skipped, if no phage tittering is needed, instead, enterohemorrhagic E. coli cells could be directly infected with the antigen–bound phage in selection wells or by addition of the phage pulled out on magnetic beads.

Although there are many other types and varieties of protein-binding chemistry, the exceptionally-high femtomolar affinity of a biotin–streptavidin pair makes this technique generally superior to others, since dissociative loss of the target during multiple washing steps at lower immobilization affinities could ultimately impede quality of the selection. In addition, the biotin–streptavidin binding, as many other receptor–ligand interactions, immune complexes included, is easy to eliminate at low pH, which could be useful during the system set-up and testing.

Several modifiers and tags for target biotinylation have been applied in the immobilization protocols at KossLab pipeline at different periods and for specific antigens (i.e., natural non-recombinant proteins could be only biotinylated in a chemical reaction), including the SNAP-mediated biotinylation and immobilization that is discussed next.

**SNAP-tagged targets: immobilization and specific proteolytic elution.** Although the biotin–streptavidin binding is not covalent, it serves all intents and purposes in biopanning, since the binding is almost irreversible ($K_D \approx 10^{-15}$ M). However, there are several popular novel protein tags that could be attached covalently to the specific chemical moieties of the insoluble matrix among multiple other useful soluble carriers. Such enzyme derivatives, as, HaloTag (Promega, USA) [115, 116], SNAP-tag [117-119], and CLIP-tag [119, 120] (both – New England Biolabs) represent self-labeling tags that catalyze covalent attachment of the exogenously added specifically designed synthetic ligand. These systems have been exploited for years in vitro and in vivo for protein labeling, localization, and fluorescence, super-resolution, and electron microscopy imaging. HaloTag is a derivative of a bacterial enzyme haloalkane dehalogenase covalently linked through a reactive chloroalkane linker to a functional group of choice. Both SNAP-tag and CLIP-tag are engineered variants of the human DNA repair enzyme O(6)-alkylguanine-DNA alkyltransferase that catalyzes transfer and attachment of the alkyl group via a thioether bond to the reactive cysteine of the enzyme. SNAP-tag has been engineered to react with O(6)-benzylguanine, while CLIP-tag – with O(2)-benzylecytosine derivatives. The discriminatory substrate binding and choice of the fluoroscently labeled substrates, makes these two tags highly applicable for orthogonal labeling of proteins in living cells [120].

Availability of various other SNAP substrates, besides fluorescent, like SNAP-biotin and SNAP-capture magnetic beads (NEB), allowed us to develop and successfully apply several variants of the antigen-immobilization technique based on the SNAP-tagging of target proteins. In addition to the classic biotin–streptavidin based immobilization of the SNAP-biotinylated target, the novel way of direct covalent attachment onto the SNAP-capture magnetic beads has been tested. Covalent nature of the direct capture and extreme substrate specificity of the engineered SNAP-tag allowed for efficient immobilization of not only highly or partially purified targets but even immobilization from the crude lysates of the cells expressing SNAP-tagged proteins. This direct covalent capture could be hugely beneficial for the proteins with IMAC-incompatible characteristics or complicated purification protocols, despite the fact, that, in our hands, the SNAP-capture beads demonstrated reproducibly somewhat lower antigen-binding capacity and specificity for phage binding, compared to the best-performing Streptavidin-coupled magnetic beads (Dyno beads, Invitrogen, USA). Also, further improvement and optimization of the surface chemistry and ligand density of the SNAP-capture beads should eliminate this problem in future.

Biotinylation of the SNAP-tagged protein is a bimolecular reaction, simple, fast, completed within a short time, and irreversible that does not require any substantial excess of the SNAP-biotin reagent and, subsequently, is free of the purification step prior to the binding to streptavidin magnetic beads. We also have inserted a Thrombin-cleavage site between the tag and the antigen in the commercial SNAP-vector (New England Biolabs) to ensure fast and gentle elution of the enriched antigen-bound phage from the magnetic beads without acidic treatment. Thrombin-cleavage site was chosen because it had the highest cleavage rate, compared to some other highly specific proteases [121], like TEV, PreScission, or 3C protease; their cleavage sites were introduced into the experimental SNAP-fusion proteins as well and the cleavage rates were tested.

Theoretically, cleavage should release only the target-specific phage, while the SNAP-tag-bound phage should remain attached to the beads, however, natural spontaneous phage dissociation could significantly con-
taminate the eluted phage. Therefore, to avoid phage binding to the immobilized SNAP-tag, a soluble SNAP-tag competitor is added during the panning step. In addition, we found, that like with other solubility-enhancing tag proteins (MBP, NusA, thioredoxin, GST, SUMO and Fh8 tag [107, 122]), fusion to the SNAP-tag (182 amino acid-long, well-structured, highly soluble, and stable protein) can drastically improve stability, solubility, and production of the hard-to-express, poorly folded, and structurally unstable proteins.

The modified library-panning protocol developed in the KossLab for the SNAP-tagged targets (Fig. 2) has been successfully used, as demonstrated below, for sAB generation against dozens of targets, including viral proteins [89, 123] as well as for reciprocal maturation of the engineered-protein G and Fab-scaffold interface.

SYNTHETIC ANTIBODIES. WHAT IS SPECIAL ABOUT THEM?

sABs are superior to the naturally produced antibody in many special features and applications. To start with, sAB production is performed in vitro, free of demanding and expensive dependence on laboratory animals and can allow for isolation of very highly-selective antibody reagents confirmed to bind the native protein, or their certain conformations, or post-translationally modified forms. Also, sABs variety is not affected by such inherent limitations of the mammalian immune system as intolerance, or presence of immunodominant antigens, or clonal drift. Moreover, while natural monoclonal antibody reproduction requires continuous maintenance and storage of the hybridoma cell line, sAB clones can be indefi-
nitely preserved in a form of frozen or lyophilized DNA. Due to the same AB scaffold used for cloning of all the variants, sABs are easy to sequence and reformat into a full antibody or expression vectors for production in bacterial or mammalian cells. This has driven development and validation of the high-throughput sAB expression platforms [124].

sABs could compete with monoclonal antibodies almost in any standard laboratory applications, like ELISA and related homogeneous and heterogeneous immunoassays, flow cytometry, immunostaining, immunoprecipitation, chromatin immunoprecipitation, and many others. However, methods, requiring recognition of SDS-denatured proteins (such as Western blotting) that are highly feasible for monoclonal antibody, could be problematic for sAB. Unlike the most of natural-antibody CDRs that should recognize a short antigenic peptide (8-18 aa long) presented on the immune cell and thus are specific for linear epitopes readily accessible upon antigen denaturation, sABs are selected by binding to the native-state proteins. In the native form, many such linear epitopes are often hidden or interrupted, and outcompeted by the three-dimensional epitopes exposed and more potent for in vitro selection. Moreover, the well-structured proteins with pronounced surface architecture and limited number of the defined low-energy states comprise the best sAB targets, while flexible, poorly folded, and unstable proteins are not appropriate targets for sABs selection but are as good as well-folded proteins for natural antibody production. Advantageously, in vitro selection potentiates unparalleled antibody customization, unachievable by natural production of immunoglobulins. By the means of precise control over the antigen composition and state under the highly-manipulatable panning conditions sABs can be tailored to: (i) target specific regions of the surface of a protein, (ii) recognize distinct conformational states of the antigen, (iii) induce desired conformational changes, and (iv) capture and stabilize transient states and multi-protein complexes [81]. These unique sAB capabilities are opening new possibilities for investigation of macromolecular structures and processes inaccessible to the traditional affinity-reagent technology.

**sAB customization strategies. Different epitope targeting sABs and translational medicine.** Like the natural process of antibody production, in vitro selection usually generates sAB variants recognizing antigen epitopes of high “immunogenicity”. However, unlike natural immunogenicity, which is contingent also on the epitope-specific antibody progression through the controls of the immune system, the in vitro antigen epitope dominance depends, mostly, on the binding affinity of the sAB phage clone, and, to a lesser extend, on the share of the clone in the phage pool under the panning.

By binding to different epitopes of the protein, sABs can affect protein function as inhibitors or activators, promote or suppress ligand binding and homo- and hetero-protein interactions. sAB-based activators and inhibitors of definite immune checkpoints have proved their clinical efficacy as immunomodulatory therapeutics activating immune responds in tumorigenesis or reducing inflammation in autoimmunity. Besides the checkpoint blockade, similarly to the natural monoclonal antibodies, sABs could impact a variety of other areas, such as, allergy, transplantation, and T-cell immunotherapies.

The KossLab pipeline production also have confirmed an undeniable potential of sABs targeting different antigen determinants in translational medicine. For example, among the selected binders that modulate isocitrate dehydrogenase enzymatic activity, the explicit one has demonstrated ability to reactivate the enzyme mutant form associated with brain tumors [90]. “Activator drugs” of this kind could be the key remedy for many genetic disorders or malignant cell transformations caused by mutations disrupting protein function. Furthermore, a panel of sABs binding to different regions of paramyxovirus envelope glycoproteins and affecting different processes of the viral entry into the cell has been used to understand the steps in viral membrane fusion leading to acute respiratory infections [125].

Among the set of generated actin-filament pointed-end binders, three sABs have demonstrated unique properties toward the actin-dynamic probing: one binder caps the pointed end, the second one crosslinks actin filaments, and the third severs actin filaments and promotes disassembly. This study may provide insights into the details of actin-filament rearrangements occurring in healthy and diseased human tissues, in innate and adaptive immune cells, during metastatic cancer and viral and bacterial infection [94].

A rapid and efficient screening to tailor phage-display for the selection of neutralizing antibody was set up in the KossLab for the Anthrax model. The high-affinity neutralizing sABs selected against edema toxin, an adenylate cyclase and a major mediator of anthrax pathogenesis, has demonstrated efficacy of the synthetic alternatives to the traditional antibody therapeutics: its potency to inhibit edema-toxin catalyzed cAMP production in human cells is comparable with that reported for the best-performing monoclonal antibodies [126].

There are also sAB applications, where it is advantageous to deal with a variety of sAB specificities targeting non-redundant epitopes on the protein [127]. Although these are often obtained spontaneously and can be easily characterized by epitope-binning methods [128], sometimes, there is one major epitope prevailing in the selection. Then, the epitope-masking strategy directing the sAB binding on the protein away from the immunodominant hot spot could be implemented during the selection step. In this case, the sABs targeting the predominant protein epitope is pre-bound to the protein, thus, excluding the epitope from the competition for phage binding
and allowing for selection of the sAB variants specific for the secondary epitopes.

In the KossLab, this trick has been efficiently applied in many instances, in particular, for generation of the sABs targeting non-redundant epitopes in two viral proteins, C-terminal domain of Ebola Zaire Nucleoprotein (EBOV NP$^{\text{CT}}$) and Zika methyltransferase (ZIKV MT), while developing a novel protein–complementation (PC)-based wash-free immunoassay [123], relevant for point-of-care (POC) applications and described in a separate chapter below. And recently, the second sAB partner for SARS-CoV-2 detecting assay has been identified after masking the major epitope of the SARS-CoV-2 Spike protein receptor-binding domain by the predominant sAB, that, in addition, has demonstrated an effective inhibition of the virus entry into the cell and can be used for the virus neutralization. Ability of this sAB pair to detect SARS-CoV-2 has been proven in the PC-based immunoassay, answering an urgent need for novel SARS-CoV-2 POC diagnostics during the COVID-19 pandemic [85].

Conformational sABs. It is common, that the protein used as a selection target is not conformationally uniform and can represent a dynamic or static mix of low-energy states, sometimes caused by contaminating ligand molecules or certain metal ions. Under such conditions, generation of a conformationally-selective antibody recognizing just one of the antigen states is accidental. Alternatively, a specific conformational state of an antigen could be targeted already at the design conception. As an illustration, adding or omitting maltose at the panning step in the maltose-binding protein (MBP) system resulted in the selection of a cohort of three classes of high-affinity and specificity MBP binders: endosteric (binding to the maltose-binding pocket), allosteric (opposite to the maltose-binding pocket), and peristeric (close to the maltose-binding pocket) [88, 93]. These sABs stabilize MBP in different conformational states as has been revealed in the respective crystal structures. Respective modulations of the MBP ligand-binding affinity in competitive, allosteric, or peristeric manners, were used as probes to quantify energy contributions of the ligand binding to the conformational changes in proteins.

Further development of this approach for more biological systems could in principle advance analysis of the energy landscapes of many systems including highly challenging regulatory proteins that control physiological responses to environmental changes.

sABs as tools for protein structure determination. As already mentioned, sABs represent indispensable tools for protein structure determination. They can be successfully used as fiducial markers for SP cryo-EM by adding mass (50 kDa for Fab) to the particle and assisting in its orientation [129]. Also, being easily crystallizable proteins with the known scaffold structure, sABs are widely used as crystallization chaperones that bind to a target of interest, enhance crystal packing, and provide high-quality phasing information. They have efficiently facilitated structure-determination of RNAs, RNA–protein complexes, protein complexes as well as individual proteins inherently recalcitrant to generate stable crystal lattices [83, 84, 130]. By driving a protein or its flexible parts to adopt and halt a uniform conformation, the conformation-specific sABs can enable crystallization of unstable, multi-conformational, multi-component or multi-subunit structures, membrane proteins and their complexes [81, 82, 98, 131].

Membrane proteins and nanodisc-based phage display. As it has been brought up by many investigators, membrane proteins continue to be among the most challenging targets in structural biology. The naturally programmed conformational changes in the membrane proteins are induced by external stimuli and are in the core of the signal transduction mechanism. Sophisticated interrelation between the structural and functional transformations underlying the transduction process is one of the most complicated though most exciting areas of the today’s membrane-protein science. Membrane proteins are difficult to purify in a native form, and conditions matching their natural surroundings are hard to reproduce for crystallization purposes. Until recently, most sAB selection protocols targeting membrane protein have been performed in detergents, that are poor mimics of the native lipid environment. To produce sABs while resolving this problem, a novel approach, that combines the power of phage display with the benefits of embedding membrane protein targets in the lipid-filled nanodiscs has been developed in the KossLab [91, 132]. Nanodiscs, small (5-50 nm in diameter) discoidal particles consisting of lipids enclosed by the membrane scaffold proteins [133, 134] have been widely used in investigation of functional and structural properties of the membrane proteins as a sophisticated membrane mimetic system with precise control over their size and composition. For the purpose of application in phage display, the scaffold proteins of the nanodiscs are biotinylated enabling their pull-out by the magnetic beads. The nanodisc-based native-like lipid environment and the overall configuration of the system allow the partially imbedded target protein to adopt characteristic structural transitions, and the sABs to bind to these transient conformations. The novel approach developed in the KossLab yielded a rich pool of sAB, that could be used as crystallization chaperones, fiducial markers for SP cryo-EM or energy probes for different conformational states of a number of membrane proteins, laying foundation for elucidation of the high-resolution structures of the functionally relevant protein conformational states and understanding their dynamic interplay.

sABs as “universal” fiducial markers for SP cryoEM. SP cryo-EM has emerged over the past two decades as a powerful structural biology tool for challenging macro-
molecular systems: it does not entail crystallization or phase determination problem and requires only small amounts of the experimental sample [129]. In 2017 the Nobel Prize in Chemistry was awarded jointly to Jacques Dubochet, Joachim Frank, and Richard Henderson “for developing cryo-electron microscopy for the high-resolution structure determination of biomolecules in solution” thus highlighting this technology as one of the greatest benefits to humankind [135]. Since SP cryo-EM success relies on the accurate assignment of particle location and orientation and their sufficient mass, only symmetric and heavy objects, like viruses, were appropriate for the method at the times of method conception [136, 137]. With the development of supportive new technologies, structures of many large protein complexes and oligomeric membrane proteins have been resolved to nearly atomic resolution by SP cryo-EM, the lower mass limit of the method has dropped to 50 kDa [138, 139]. By increasing the particle mass and providing fiducial markers for particle alignments, sAbs significantly enable cryo-EM analysis and protein-structure determination [140, 141].

The antibody-based fiducial markers have considerably aided the membrane-protein structural studies in the field of G-protein coupled receptor (GPCRs) [142-144]. There are hundreds of different GPCRs belonging to 3 major classes encoded in human genome. They play a central physiological role in the regulation of cellular responses to a wide variety of stimuli in both health and disease and thus represent one of the largest types of surface receptors targeted by drugs. In the KossLab, sAbs have been generated to the two major G-protein subclasses binding GPCRs: trimeric G, and Gs, as well as mini-Gs, with the goal to overcome specificity limitations of the fiducials tailored for a particular GPCR. Epitope binding revealed that multiple distinct epitopes were targeted for each of trimeric G-protein, while some of the sAbs cross-react between the trimeric G, and Gs, suggesting their universal application across the subclasses. Similarly, sAbs were generated to a representative of another class of GPCR signaling partner, G-protein receptor kinase 1, supporting the generalizability of the fiducial marker approach. EM data confirmed efficacy of the sAbs as single and dual fiducials for multiple GPCR signaling complexes [87].

In addition, a universal sAB-targeted tag system that eliminates the need for a specific antibody for each individual membrane protein under structural study has been developed. To that end, an engineered variant of the apocytochrome b562, 12 kDa protein BRIL has been chosen as such a tag since it contains terminal helical extensions easily adjustable for the distortion-free seamless connection to α-helices present either in the loops [145] or at the termini [146, 147] of the membrane proteins. Several sAbs generated and matured by the customized phage display selections against BRIL gained sub-nanomolar affinities and unhindered BRIL binding in multiple fusion systems. Negative-stain TEM and cryoEM structures of several examples of BRIL-membrane protein chimera demonstrated effectiveness of the sAB–BRIL systems as universal fiducial markers [86]. Thus, the technology is advancing towards making the “universal” sAbs for the whole classes of proteins enabling their crystallographic and SP-cryoEM studies and potentially forming the “off the shelf” line of enhancement reagents. These would eliminate the need to incorporate the challenge of an antibody-engineering into the workflow of protein-structure determination projects.

Tools like sAB-based crystallization chaperones and fiducial markers have drastically improved the level and quality of structural analyses. To further improve effectiveness of the sAbs as structure-determination chaperons, a special phage display engineering strategy, including a heat-stress step for the selected clones, has been used to generate Fab scaffold variants, that significantly reduces inherent flexibility of the “elbow” regions, which link the constant and variable domains of the Fab, disordering its structure. The strategy was validated using the previously recalcitrant Fab–antigen complexes: introduction of the engineered elbow region remarkably enhanced crystallization and diffraction resolution, while their high affinity and stability retained [148]. Also, sAB solubility, was addressed in the design of shotgun scanning libraries introducing aspartate as a negative design element at the antigen-binding site. The selected high-affinity variants featured a polar ring surrounding the paratope. This significantly enhanced sAB specificity and solubility, crucial for easily aggregating antibodies, while maintained its affinity to the antigen [149]. These universal improvements of the overall sAB rigidity and solubility have further shaped them for structural determination studies.

**ENGINEERED PROTEIN G AND ANTIBODY-SCAFFOLD VARIANTS**

**Ig-binding proteins.** Fortunately, nature has provided protein engineers not only with the IgG molecule as a framework for sAbs but also with Ig-binding proteins enabling their isolation, detection, and assembly. Several species of infectious bacteria express multivalent Ig-binding proteins on their surface. Single domains from a number of almost identical entities comprising multidomain bacterial protein A (Staphylococcus aureus), protein G (Streptococcus species C and G), and protein L (Peptostreptococcus magnus) have been characterized in detail both biochemically and structurally [150]. Each of these proteins has a different antibody binding profile in terms of the portion of the antibody that is recognized and the species and type of antibodies it binds. Protein A has greater affinity for the rabbit, pig, dog, and cat IgG.
whereas protein G has greater affinity for the mouse and human IgG. In addition to the predominant binding of both proteins A and G to the interface of CH2-CH3 heavy chains in the Fc (fragment crystallizable), they also bind to the Fab: Protein A — to VH, while protein G binds simultaneously, but very weakly, to CH1, a highly conserved domain across many isotypes and species, and CL kappa. Protein L only binds to VL kappa of the Fab part and exclusively [151, 152].

Proteins A, G, and L have a scope of biochemical, biotechnological, and medical applications and are widely used in antibody purification and test methods like immunoprecipitation, ELISA, and Western blotting [153]. The “universal” recombinant fusion protein A/G comprised of four domains of protein A and two domains of protein G demonstrates additive properties of the two proteins and captures antibodies from the combined spectra of species and isotypes [154, 155].

Recent engineering efforts in applied immunological and biochemical research have led to production of the synthetic ligands mimicking protein A and L (peptides, engineered protein domains, and designed artificial molecules). These mimetics can replace the originals for purification of antibodies, while surmounting some drawbacks such as high cost, low binding capacity, limited life cycles and so on [156].

An engineered GA1 variant. Protein G is used for IgG purification by the virtue of its affinity to the Fc portion of the molecule (KD \( \sim 10 \) nM), while its binding to the Fab CH and CL is pretty weak (KD \( \sim 3 \) \( \mu \)M). The 4D5 Herceptin Fab scaffold (Fab5) of the KossLab pipeline containing E123S mutation in the Fab CL has been used for affinity maturation of protein G (a 65 aa-long C2 domain) [157]. The affinity maturation design involved soft randomization of the two points of contact with the Fab5 scaffold: one, residues 15-24, involved in the formation of antiparallel \( \beta \)-strands with Fab CH and second, residues 37-43, interacting with the short non-conserved \( \beta \)-helical region connecting \( \beta \)-strands in CL. The matured variant GA1 demonstrated the highest improvement in the Fab5 binding affinity, mostly due to substitutions of NDNG in the positions 40-43 for YVHE. In the crystal structure of GA1–Fab5 complex, these residues form a structural helical cap in the GA1 interdigitating with the CL \( \alpha \)-helical residues SQLKS improving the interface complementarity.

An engineered FAB\textsuperscript{LRT} scaffold variant. Compared to the wild-type protein G, GA1 demonstrates a significant affinity boost in binding Fab5 (KD \( \sim 50 \) nM) — adequate for applications that involve genetically linked GA1 strings to make multi-valent Fab assemblages [157]. However, binding is still characterized by fast dissociation kinetics that is not optimal for the desired non-equilibrium applications. To further improve affinity and dissociation kinetics of the GA1–Fab complex, we undertook a reciprocal phage display approach: Fab5 scaffold was affinity matured against GA1 [123]. The library design was focusing on the residues 123-127 (SQLKS) of CL. Kunkle mutagenesis using NNK/NNT randomization yielded \( \sim 10^{10} \) clones with theoretical diversity of \( \sim 1.7 \times 10^7 \) unique codon variants (Fig. 3). Several selected variants demonstrated significantly increased binding affinity and

**Fig. 3.** Construction of the phage-display library for Fab scaffold GA1-binding affinity maturation. Schematic representation of the degenerate-codon based randomization by Kunkel mutagenesis: a random Fab5-encoding phagemid containing dU substitutions for T was isolated from CJ236 E. coli strain (1), annealed with a synthetic DNA oligonucleotide comprised of a randomization sequence: \( \text{LC}^{123}(\text{NNK})_4(\text{NNT})(\text{NNK})_4\text{LC}^{127} \), flanked by 15 nt of the wild-type overhangs (2) used to prime the DNA synthesis by T4 DNA polymerase; the synthesized dU/T duplex phagemid molecule was transformed into the TG1 E. coli strain aiming for destruction of the wild-type dU-containing strand and efficient replication of the mutant T-containing strand (4). The following superinfection with M13 K07 helper phage resulted in the mutant phage propagation (5).
reduced dissociation rate. The best selected Lc variant with ~500-fold improved affinity to GAI contained a serendipitous two amino acid deletion, which may have occurred during the synthesis of randomizing DNA oligonucleotides. Replacement of the original SQLKS sequence of Fab with ΔΔLRT (FabLRT) resulted in the \( K_d \) of ~100 \( \mu M \), slow dissociation rate (~2.4 \times 10^{-4} \text{ s}^{-1} \) and did not affect Fab stability or expression.

The key structural basis underlying this ultra-high affinity of the FabLRT was revealed in a crystal structure solved for the GAI–FabLRT complex: a significant deletion-induced rearrangement and extension of the interface has packed the guanidinium group of Arg124 (replacing Lys126 in Fab) against the aromatic ring of Tyr40 of GAI in a cation–π interaction, while providing H-bonding with the Tyr40 carbonyl via the secondary amine in \( \varepsilon \) position of Arg124 [123].

A NOVEL PLUG-AND-PLAY PLATFORM

The ultra-high GAI binding affinity and extremely low dissociation rate of the FabLRT encouraged us to develop a multivalent and multi-specific platform based on the GAI module that could be easily modified just by exchange of the Fab component. Although the FabLRT–GAI interaction is not covalent, no exchange between the partners, which could compromise the performance of such a platform, has been detected within the timeframe of experiments.

There is a great variety of possible functionally relevant GAI genetic fusions, allowing the researcher to design highly customized affinity reagents that can be assembled in a wide scope of different formats in a straightforward way (Fig. 4a). The simplest example is being a string of genetically linked GAI modules (Fig. 4b) for binding multiple, same or different-specificity, FabLRT molecules for the avidity boost or “randomized” multi-specificity, respectively.

GAI could be genetically coupled with the enzymes and protein tags for antigen detection by means of FabLRT (Fig. 4c): we have successfully applied GAI fusions to β-lactamase (with a fluorogenic substrate) or SNAP-tag (modified with a fluorescent label) to be paired with the interchangeable FabLRT of a variety of specificities for routine antigen detection (ELISA, CFC) and imaging purposes. A novel wash-free β-lactamase complementation-based detection assay (Fig. 4d) has been developed by fusing GAI to two split β-lactamase complementation fragments [158] (details of this work are discussed in the next chapter). Also, linking of GAI to the IgG Fc fragment replacing Fab, resulted in the IgG mimic (Fig. 4e) with the interchangeable FabLRT specificity that could be utilized in place of multiple IgGs in the Fc-mediated processes. In addition, the GAI fusions to the transmembrane domains (TMDs) of cell receptors made in the endoplasmic reticulum, anchors the exchangeable Fab component to the eukaryotic cell surface, thus, enabling investigation of the cell–cell signaling and interaction, and many other applications.

It turned out that the Fab-binding affinity maturation has been beneficial for GAI in other aspects as well: it significantly reduced Fc binding and practically abolished GAI affinity to all natural Fabs tested, including human kappa (FabH) – the parental to Fab[5] [123]. This made the GAI–FabLRT interactions insensitive to the presence of endogenous IgGs, which are very likely in many samples as a background – an essential detail for an experimental platform. High specificity and orthogonal nature of the GAI–FabLRT binding also allowed us to genetically connect a second specificity to the GAI module in a form of FabH, or scFv, without being concerned of their possible interaction and interference with the FabLRT binding. The resulting bi-Fabs, functionally mimicking the bi-specific T-cell engagers (BiTEs), very efficient immunotherapeutics, have the advantage of one easily interchangeable specificity and are discussed in more details below. FabLRT, FabH, and scFvs containing and proximity-sensitive FabLRT and split-scFv GAI-complexes represent two of many possible tri-specific formats (Fig. 4f). In addition, it is important to note, that the control over the linker length in the GAI fusions could be an essential variable for applications in the systems with specific dimensional or steric demands.

A novel β-lactamase complementation-based assay.

As mentioned above, we have developed a wash-free antigen-detection system based on the GAI module fusions in a protein-fragment complementation format, widely used for identification and studying of protein–protein interactions. It is comprised of two fusions of GAI to two complementary β-lactamase fragments (BLFs) individually pre-bound to the two FabLRT component targeting non-redundant epitopes of the antigen of choice. When the fusions bind independently to the non-overlapping epitopes on the antigen by means of FabLRT component, proximity of the BLFs attained within the immune complex should allow them to reassociate into the active enzyme, detectable with a fluorogenic substrate (Fig. 4d). Success of complementation and enzyme reactivation depend to a great extent on the appropriateness between the length of fusion linkers and the distance between the antigen epitopes targeted by FabS.

The recent world threats caused by severe viral infections motivated us to choose Ebola and Zika virus proteins (EBOV NTCT and MT ZIKV, respectively) as antigens to be detected in the system and used initially for generation of the high-affinity sABs by phage-display selections reformatted next into the FabLRT scaffold. The independent mode of antigen binding by the Fab pairs was confirmed by SPR. The 30 aa-long linker (~100 Å stretched) used in all GAI fusions turned out to be sufficient for successful complementation; β-lactamase activ-
ity was readily detected by a fluorescent signal in the reactions containing viral antigens with antigen-appropriate FabLRT pairs and complementary GA1-BLF [123].

This wash-free sandwich antigen-detection system has been also applied for SARS-CoV-2 detection using a pair of FabLRT targeting two non-overlapping epitopes of the Spike RBD. The detection limit of the systems (∼10 nM) falls within the standard range for a sandwich antigen-detection immunoassay, like ELISA. Together with the simple and fast non-wash procedure, reproducible quantitative results, and detected-antigen adjustability, easily achievable by FabLRT pair exchange, this makes the protein-complementation GA1-based assay a suitable candidate for POC applications; in addition, it was shown, that its components can be freeze-dried and fully reactivated after rehydration [85].

The strong assay performance confirmed high potential of GA1 as an effective non-covalent bond between the BLFs and FabLRT of any specificity, thus, laying the basis for numerous plug-and-play opportunities.

**A novel plug-and-play BI-Fab format.** Our next endeavor undertaken was to apply the ultra-high affinity GA1-FabLRT platform for building bi-Fabs complexes mimicking structurally and, hopefully, functionally the bi-specific T-cell engagers (BiTEs), with an advantage of one varying specificity providing by the interchangeable Fab component in a plug-and-play manner.

Classical BiTEs represent engineered fusion molecules, consisting of two different scFvs of particular specificities—one, binding to T-cells via the CD3 component of the T-cell receptor complex, and the other—to a cancer cell via a tumor-specific molecule [27]. This engage-
ment leads to the T-cell dependent death of the cancer cell.

The principle of two-molecule design of bi-Fabs exploits the GA1 module as a reliable non-covalent fixative for Fab\textsubscript{LRT} already experimentally approved in this capacity in the plug-and-play detection assay discussed above. Since GA1 orthogonal binding demonstrates no measurable binding to Fab\textsubscript{H} scaffold, it allows to genetically fuse Fab\textsubscript{H} to GA1 through a flexible linker without affecting the GA1-binding when Fab\textsubscript{LRT} is added to the fusion, forming a bi-Fab—a stable association of two distinct antigen-binding specificities [123].

To mimic the BiTE design in the bi-Fab format, we choose two antigen targets: (i) T-cell specific CD3 receptor commonly used in the BiTE therapy and as a specific cell surface marker, and (ii) HER2 receptor, highly overexpressed on the cell surface in many cancer cells such as breast-cancer cell lines, including SKBR3. To that end, three well-characterized antibodies were chosen: the already tested in the BiTE format humanized \(\alpha\)-CD3 antibodies, OKT3 or UCHT1; and \(\alpha\)-HER2 trastuzumab, highly therapeutic anti-cancer antibody. To construct bi-Fabs in either “polarities”, each antibody was reformatted in both Fab\textsubscript{H} and Fab\textsubscript{LRT} scaffolds and all three resultant Fab\textsubscript{H} variants were fused to GA1 at their HC C-termini via a 13 aa-long GS linker. The bi-Fabs were assembled combining the CD3 or HER2 specificities reciprocally as either Fab\textsubscript{H}–GA1 fusion or Fab\textsubscript{LRT} (Fig. 5).

All the bi-Fab combinations showed efficient T-cell engagement activity, similar to the activity of the covalently linked BiTE, as measured in the PBMC-SKBR3 co-cultures by three readouts: (i) activity of the cytoplasmic enzyme, lactate dehydrogenase, released into the medium upon cell lysis caused by activation of cytotoxic T-cells; (ii) interleukin IL2, and (iii) interferon \(\gamma\) produced by T-helper cells. As expected, functionality of the bi-Fabs was absolutely dependent upon the genetic fusion of GA1 to Fab\textsubscript{H}, as no detectable activity was observed when all the individual components (Fab\textsubscript{H}, Fab\textsubscript{LRT}, GA1) were added as three separate unlinked entities [123].

Given the ease of interchanging Fab\textsubscript{LRT} in the bi-Fabs, we envision the following possible advances of the GA1-based plug-and-play BiTE mimetics. A high-throughput platform can be developed to profile bi-Fabs

Fig. 5. Schematic representation of T-cell engagement in cytolytic synapse with cancer cell expressing HER2 receptor by means of bi-Fab inter-cellular bridging: it binds HER2 receptor with Fab\textsubscript{H}, genetically fused to GA1, while Fab\textsubscript{LRT}, non-covalently attached to GA1, binds to CD3\delta\epsilon co-receptor of T-cell specific receptor (TCR). Activation of the CD8 cytotoxic T-cell involves cytolytic granule fusion and release directed toward the cancer cell, while the activated CD4 helper T-cells increase production and secretion of cytokines interferon \(\gamma\) and IL2, activating cytokine production, cytolytic effect, and proliferation of peripheral blood immune cells. Lysis of the attached cancer cell is caused by the membrane perforation by perforin, and subsequent programmed cell death induced by granzymes, entering through the perforin pores into the cancer cell cytoplasm. Compilation of the T-cell activation effects is shown. Bi-Fab is colored as following: GA1 is shown in red; fused to it, Fab\textsubscript{H} (specific for HER2 receptor of cancer cell) — in green, and Fab\textsubscript{LRT}, specific to CD3 — in blue. Two different CD3-specific Fab\textsubscript{LRT}, derivatives of OKT3 and UCHT1 antibodies, were interchangeably used with trastuzumab-derived Fab\textsubscript{H} fused to GA1.
targeting many different cancer-specific cell-surface markers through the FabLRT assortment. Then, the FabLRT variety could target any number of surface antigens on a certain type of cancer cells, amplifying therapeutic effect. In addition, as in other GAL-fusion applications, the linker length could be easily adjusted tailoring the bi-Fab for the particular antigen or even antigen epitope. Multiple GAL modalities could be also included into the design to exploit possible avidity effects in the resulting multi-valent bi-specific formats.

**PERSPECTIVES**

Protein engineering is now a mature field of protein science. While the ability to clone, express, and purify proteins is well established and is incorporated into the most standard operating procedures in the research laboratories, the field will continue to evolve as new technologies that build on those described here are brought into practice. One might argue that protein engineering has been practiced for hundreds of years through genetic selections. However, attempts to understand molecular basis of the phenotypic alterations and to rationally design them have only become possible through the relatively recent process involving integration of the protein structure information. An important aspect of protein engineering is that it has both basic and applied research applications. Thus, we can work on projects that build our basic understanding of the molecular basis of biological systems, while at the same time, produce molecules that are beneficial to health care and to society in general.

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