Subtleties and Dynamics of Antibody Function: Lessons from Bovine Antibody Fragments

Ashish Sachan and Azad K. Kaushik

Department of Molecular and Cellular Biology, University of Guelph, Guelph Ontario N1G2W1, Canada

Corresponding author: Azad K. Kaushik, Department of Molecular and Cellular Biology, University of Guelph, Guelph Ontario N1G2W1, Canada, Tel: 424854389; Email: akaushik@uoguelph.ca

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Abstract

The global market for antibody-based drugs is expected to double within a decade given their clinical success against diseases where conventional therapies fail. Our studies of antibody structure and function have revealed subtle differences in the role of $V_H$ and $V_L$ domains in antigen recognition and virus neutralization functions. Studies of bovine antibody fragments, such as, scFv (single chain fragment variable) highlight how a single substitution mutation in the framework-3 may influence viral neutralization potency. The viral neutralization potency of bivalent diabody, i.e., dimerized scFv, is almost doubly enhanced when compared to its monomeric form as scFv. Further scFv multimerization as tri- and tetrabody does not necessarily enhance virus neutralization function. Future design of antibody-based drugs should take into consideration subtle structural modifications in the framework regions that may influence the desired function.

Keywords: Polyspecificity; Monospecificity; Heavy chain; Light chain; Antibody fragment; scFv; Fd; $V_H$; $V_L$

Introduction

Antibody, nature’s most magnificent molecule, has come of age moving forward from an era of polyclonal and monoclonal antibodies to its applications as functional antibody fragments or modifications thereof for developing new drugs against vexing diseases, such as, cancer and viral infections [1-4]. The conventional therapies against these diseases have not been successful resulting in a shift towards development of novel antibody-based therapeutics. The clinical success of antibodies can be assessed from their major commercial impact as evidenced by the market for antibody drugs being over $60 billion in 2013, which is expected to double globally within the decade.

Role of $V_H$ and $V_L$ in Antibody Functions

Current knowledge of the structure and function of bovine antibodies [5,6] provides important clues to strategically design antibody-based drugs by optimized engineering [7-12]. Analysis of linked $V_H$+$V_L$ pair (single chain fragment variable; scFv) and individual fragment variable heavy (Fd$V_H$) or fragment variable light (Fd$V_L$) domain of multispecific IgM and monospecific IgG antibodies as compared to whole antibody molecule show subtle differences in their antigen recognition and viral neutralization functions [13]. In some instances, including multispecific IgM and monospecific IgG antibodies, it is mainly the Fd$V_H$ that recognizes the antigen [13]. There exist subtle differences in antigen recognition pattern among Fd$V_H$, scFv and whole antibody molecule. In some instances, the presence of constant region present in the whole IgM antibody enhanced antigen recognition function while in others it did not [13]. Similarly, subtle differences existed between Fd$V_H$ and scFv where the absence of light chain variable region led to loss of antigen recognition function [13]. Thus, the role of Fd$V_L$ in providing structural support to antigen recognition is not excluded. In case of monospecific IgG antibody, both Fd$V_H$ and Fd$V_L$ in scFv format are required for antigen recognition as well as viral neutralization functions. The phenomenon of antibody degeneracy is clearly evident in an induced monospecific IgG antibodies [13].

Framework Residues can Influence Antibody Function

Our laboratory was the first to develop bovine single chain fragment variable (scFv) using $V_H$ and $V_L$ from an IgG1 antibody against bovine herpes virus type 1 (BoHV-1) [14,15]. The $V_H$ and $V_L$ linked with a flexible 18 amino acid linker (GGQSRG3S3G3S) in $V_H$-$V_L$ orientation allowed proper $V_H$+$V_L$ pairing to form a monomeric single chain fragment variable (scFv-18L; Figure 1) with a single antigen-binding site. Indeed, scFv-18L recognized and neutralized BoHV-1 in vitro at molar concentrations as low as 0.18 M. The structural and functional dynamics of the framework regions of the bovine antibody are influenced by the 3rd framework region (FR3) as it has high AGPY: TNC (mutational hot spots) as well as high (A+T): (G+C) ratio [16]. For these reasons, we tested in parallel the antigen recognition and viral neutralization dynamics of a scFv-18L with a substitution mutation at position 98 of FR3 (Asp to Gly; prior to anchor residue), designated as scFv4m-18L. Both wild type scFv-18L and mutant scFv4m-18L recognized the target epitope on the envelope of BoHV-1 in an immunofluorescence assay. Hence, the qualitative antigen recognition function was not affected by the single substitution mutation in the FR3 of mutant scFv. Importantly, the single substitution mutation in the FR3 affected the virus neutralization potency by 2.7 folds as compared to the scFv4m-18L [15].

The single substitution mutation in FR3 would not have been expected to influence viral neutralization potency to such a large extent as both the wild type and mutant scFv had identical CDRs, including the CDR3H that mainly makes contact with an epitope. Such a remarkable decrease in viral neutralization potency as a result of single substitution mutation in the FR3 suggests subtle influence of framework regions on the configuration of the antigen-binding site.
Viral Neutralization Function of Bivalent Diabodies is Comparable to Tri- And Tetra-Bodies

In attempts to enhance viral neutralization potency of the scFvs, our laboratory developed scFvs capable of neutralizing BoHV-1 with 7 (GQS2RS2) and 2 (GS) amino acid linkers [14,17]. A shortening of linker size to ≤ 7 amino acids does not permit monomeric configuration but results in complementary VH+VL pairing of two fragments with two antigen-binding sites, called bivalent diabody (Figure 1) [18-20]. Theoretically, it would be expected that half a molar concentration of bivalent diabody would be required to achieve viral neutralization as compared to monovalent scFv-18L. The actual experimental observations indeed proved it to be nearly so, as 0.1 µM concentration of diabody (scFv1-7L) is required to achieve viral neutralization as compared to 0.18 µM diabody (scFv-18L) [14,15]. Clearly, scFv dimerization helps proportionately enhance viral neutralization potency of scFv. Further attempts to enhance viral neutralization potency were made by reducing the linker size to two amino acids that led to generation of either trivalent tribody or tetravalent tetrabody (Figure 1) [17]. A comparison of monomeric scFv-18L with tri- or tetrabody revealed only two-fold difference in viral neutralization potency. Thus, there is no significant enhancement of viral neutralization potency of tri- or tetrabody when compared to bivalent diabody [17].

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Glycosylation does not Affect Antibody Functions

The construction and expression of various antibody fragments using different microbial and mammalian systems is discussed elsewhere [21-26], each with its own advantages or disadvantages. Certain protein expression systems, such as, P. pastoris, cause extensive glycosylation of the expressed recombinant antibody fragments. We did not observe that glycosylation introduced by P. pastoris affected antigen recognition and viral neutralization functions of recombinant scFvs or single domains. There is a caveat with regard to the Ni chelating affinity purification, however, as diabodies, tri- and tetrabodies are relatively difficult to purify because of commonly experienced protein aggregation and cleavage or degradation of histag.

Future Directions

Based on studies with the bovine antibody fragments in our laboratory, it is appropriate to conclude that future design of antibody-based drugs should take into consideration subtle structural modifications that could affect antigen recognition and viral neutralization functions. At the same time, it is possible to optimize functions of an antibody or its fragments by optimizing framework residues or including the constant domain. In addition, there exist limits to enhancing scFv viral neutralization potency via multimerization since no significant differences exist in terms of molar concentration beyond bivalent diabody when compared to tri- and tetrabodies. Such fine functional subtleties of scFvs and their multimers would be obvious once their 3-dimensional structural properties, including conformational flexibilities, are fully defined.

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References


