Improving effector functions of antibodies for cancer treatment: Enhancing ADCC and CDC

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Abstract: As platforms for therapeutic agents, monoclonal antibodies (MAbs) have already been approved, and several MAbs have demonstrated clinical effectiveness in a variety of malignancies. However, several issues have also been emerging in antibody therapy, such as high cost and insufficient drug action. Recently, to improve MAb activity in humans, effector functions have been subjects of focus, especially antibody-dependent cell-mediated cytotoxicity (ADCC) and complement-dependent cytotoxicity (CDC). Extensive efforts have been made to enhance these effector functions of MAbs, and successful approaches have been reported by us and others, wherein the binding activity of MAbs to FcγRIIIa or C1q is increased by introducing amino acid mutations into heavy chain constant regions or through glyco-modification of Fc-linked oligosaccharides. In addition, one of the next approaches to optimizing therapeutic antibodies would be to combine multiple enhancing modifications into a single antibody platform to overcome the diverse mechanisms of clinical resistance of tumor cells. For this aim, we have recently developed a successful combination composed of ADCC-enhancing modification by the fucose depletion from Fc-linked oligosaccharides and CDC-enhancing modification by IgG1 and IgG3 isotype shuffling in heavy chains, which could be of great value for the development of third-generation antibody therapeutics.

Keywords: ADCC, CDC, effector functions, Fc oligosaccharides, IgG isotypes, nonfucosylated IgG

Introduction: Current status of therapeutic antibodies

Since the late 1990s, more than 20 monoclonal antibodies (MAbs) have been approved as therapeutic agents, and MAbs are emerging as a major new class of drugs that confer great benefits to patients. In fact, therapeutic antibodies have demonstrated improvements in overall survival and time to disease progression in a variety of malignancies, such as breast, colon, and hematological cancers.1–4 Recently, adding to antigen binding and specificity, “effector functions” have come under increasing focus as a cause of drug action by therapeutic MAbs in humans. In particular, Fc gamma receptor (FcγR) IIIa-dependent effector functions may be one of the major critical mechanisms responsible for the clinical efficacy of therapeutic MAbs; this is supported by genetic analysis of working polymorphisms of the receptor in patients.5–9 FcγRIIIa, a member of the leukocyte receptor family FcγRs, is known to be a major triggering receptor of ADCC in natural killer (NK) cells. Several therapeutic MAbs are capable of ADCC, such as anti-CD20 rituximab (Rituxan®), anti-Her2 trastuzumab (Herceptin®), anti-tumor necrosis factor-α (anti-TNF-α) infliximab (Remicade®), and anti-RhD.5,6,9–14 Complement-dependent cytotoxicity (CDC), another effector function of antibody, is also considered a possible anti-tumor mechanism of rituximab and alemtuzumab (Campath-1H).15,16 Furthermore, most therapeutic antibodies that have been licensed and developed as medical agents are of the human IgG1 isotype, which can induce strong ADCC and CDC when compared with the other heavy chain isotypes of the human antibody. These effector functions are
activated through the interactions of the Fc with either FcγRs or complements, and the interactions are affected by N-linked biantennary complex-type oligosaccharides attached to the antibody Fc region, which is heavily fucosylated in human IgG1. In addition to their multifunctional activities, therapeutic MAbs of human IgG1 isotype have long-term stability in blood via a unique neonatal Fc receptor (FcRn) mechanism, which has never been seen in small molecular drugs.17,18 Thus, the MAb platform is very suitable for developing medicines based on molecular targeting. However, we have also realized a shortcoming of the current generation of therapeutic antibodies, as exemplified by the insufficient results of numerous clinical trials, especially in antibodies recognizing surface tumor antigens, and by an economically intolerable cost with multiple high-dose administration treatment even in the case of effective therapeutic antibodies.19–22 It is time to design next-generation therapeutic antibodies that can overcome these issues.

**Basic structure and physiological mechanism of therapeutic antibodies**

Although five classes of immunoglobulin (IgM, IgD, IgG, IgA, and IgE) and four IgG subclasses (IgG1, IgG2, IgG3, and IgG4) are present in humans, IgG1 is primarily employed as a therapeutic agent due to its long half-life in blood (approximately 21 days) and due to its advantage in effector functions compared to those of the other Ig classes and subclasses.23 The MAbs of the human IgG1 isotype has a basic structure of 150 kDa consisting of two immunoglobulin light chains and two immunoglobulin heavy chains in covalent and noncovalent association, resulting in the formation of three independent protein moieties – two Fab regions and one Fc region – which are connected through a flexible linker designated as the hinge region (Figure 1). Fab regions in an antibody molecule are of identical structure, wherein each expresses a specific antigen-binding site, and the Fc region expresses interaction sites for ligands which can induce effector functions, including three structurally homologous cellular Fc receptor types (FcγRI, FcγRII, FcγRIII), the C1q component of the complement, and the FcRn.24,25 The physiological activities of therapeutic antibodies are mediated by two independent natural immunoglobulin mechanisms: the efficacy of therapeutic antibodies results from its specific and bivalent binding to the target antigen (eg, blockade or neutralization of target antigen or induction of apoptosis) and from effector functions that are activated only by the formation of immune complexes of the Fc and the effector ligands named above.

Heavy chains are covalently paired by disulfide bonds in hinge regions, and each heavy chain is connected to a light chain by a disulfide bond between CH1 and CL. A pair of VH and VL in Fab regions makes an antigen binding site. In the CH2 domains of Fc regions, an oligosaccharide is covalently attached to the both domains at asparagine 297 (Asn-297). The CH2 domains, an oligosaccharide is covalently attached to the both domains at asparagine 297 (Asn-297).

**Figure 1** Structure of human IgG1 antibody. The MAbs of human IgG1 isotype consists of two immunoglobulin light chains and two immunoglobulin heavy chains. Heavy chains are covalently paired by disulfide bonds in hinge regions, and each heavy chain is connected to a light chain by a disulfide bond between CH1 and CL. A pair of VH and VL in Fab regions makes an antigen binding site. In the CH2 domains of Fc regions, an oligosaccharide is covalently attached to the both domains at asparagine 297 (Asn-297).
the Fc via the multiple noncovalent interactions with the CH2 domains. Therefore, the oligosaccharide influences the Fc structure, and there are suitable oligosaccharide structures among the natural heterogeneity of the Fc-linked oligosaccharides for the binding between Fc and effector ligands. For example, human IgG1 antibodies bearing the Fc-linked oligosaccharides lacking core fucose show improved binding to FcγRIIIa and much enhanced ADCC compared to fucosylated ones.

**Importance of ADCC on the clinical efficacy of therapeutic antibodies**

There is a substantial discrepancy between the potency of therapeutic antibodies in vitro and in vivo in terms of the required doses, especially in the anticancer therapies. Cancer patients treated with therapeutic antibodies typically need to receive weekly doses of several hundred milligrams over several months to maintain an effective serum concentration of over 10 μg/mL. On the other hand, the maximal in vitro cellular cytotoxicity by ADCC of these therapeutic antibodies can be achieved at an antibody concentration of less than 10 ng/mL, which is several orders of magnitude below the targeted serum concentrations in vivo. This discrepancy — that is, the low in vivo efficacy of therapeutic antibodies in contrast to the high in vitro cytotoxicity is primarily due to competition between human serum IgG and therapeutic antibodies for binding to FcγR on effector cells; namely, endogenous human serum IgG strongly inhibits ADCC induced by therapeutic antibodies.

Although the importance of antibody effector functions of ADCC on the clinical efficacy of therapeutic antibodies has long been a matter of debate, recent clinical evidence based on genetic analysis of FcγR polymorphisms in cancer patients treated with anti-CD20 rituximab and anti-HER2 trastuzumab therapies has suggested the importance of ADCC. The superior clinical response of patients carrying the high-affinity FcγRIIIa allotype (FcγRIIIa-158Val) for anti-CD20 rituximab has been demonstrated, in contrast to that of patients carrying the low-affinity allotype (FcγRIIIa-158Phe). Breast cancer patients who responded to anti-HER2 trastuzumab with complete or partial remission have been found to have a higher capability to mediate in vitro ADCC by trastuzumab than nonresponders. A significant correlation has also been reported between the clinical responses of the therapies and the polymorphisms of low-affinity FcγR in some diseases, such as rituximab-treated patients with systemic lupus erythematosus (SLE) and Waldenström’s macroglobulinemia, Crohn’s disease patients treated with anti-TNF-α infliximab, and pregnant women with fetal hemolytic disease treated with anti-RhD. Thus, the importance of ADCC for the clinical efficacy of therapeutic antibodies is now widely recognized, and ADCC enhancement technology is expected to play a key role in the development of next-generation therapeutic antibodies with improved clinical efficacy.

**ADCC-enhancing technology**

**Artificial amino acid modifications in the Fc**

The effector functions of an antibody can be manipulated by engineering amino acid sequences of the constant regions. In particular, facilitating FcγRIIIa binding by mutated antibody sequences may be of therapeutic value. Thus, amino acid alterations in the Fc that possess improved FcγRIIIa binding and resultant ADCC-enhancing capacity have been extensively studied, mainly by biotech companies, using random or rational designing approaches with high-throughput protein expression systems. A group at Genentech has discovered variant Fcs with enhanced ADCC-inducing capacity having up to three mutations, based on a comprehensive alanine scanning method on the accessible surface of the Fc. The Xencor group has employed a computational design algorithm and found a set of Fc variants with extremely strong FcγRIIIa binding (up to >100-fold enhancement in KD), which showed a strong cytotoxic activity in cynomolgus monkeys. The Macrogenics group has also described Fc mutants with enhanced ADCC by using a unique yeast display system.
Removal of fucose from Fc-linked oligosaccharides

It is widely recognized that the core fucose of Fc-linked oligosaccharides greatly affects the ADCC of therapeutic antibodies and the removal of the fucose markedly enhances the ADCC via improved FcγRIIIa binding without altering antigen binding or CDC. ADCC enhancement by fucose removal is now considered a very important technique for the development of therapeutic antibodies mediating ADCC.21,37,41,42,45,51–63 The MAbs of IgG1 bearing the biantennary-complex type of Fc-linked oligosaccharides lacking core fucosylation exerts the strongest ADCC in a comparison of glycoforms.51 In human IgG, Fc-linked oligosaccharides lacking core fucose exist as portions of naturally occurring heterogeneities, and therefore there is little concern regarding its intrinsic immunogenicity.31,32 Importantly, the enhancement of ADCC by the core fucose removal is thought to leach the saturated level, according to a study wherein no significant difference in ADCC was detected between the ADCC-enhanced IgG1 by either the core fucose removal or amino acid mutations S239D/S298A/I322E, which have higher FcγRIIIa binding affinity.60 Moreover, no additive effect is shown on the B-cell depletion activity of anti-CD20 IgG1 rituximab in human blood by the combination of these techniques, although an additive increase in FcγRIIIa binding is observed in a BIAcore biosensor analysis. Thus, the enhanced FcγRIIIa binding of nonfucosylated antibodies appears to be sufficient to maximize the ADCC of therapeutic antibodies without introducing artificial amino acid mutations. Furthermore, the nonfucosylated antibodies have achieved high effector activity at low doses, against tumor cells expressing low levels of antigen, or even in NK cells harboring the low-affinity FcγRIIIa allotype (FcγRIIIa-158Phe).41,53 Strong ADCC enhancement of the nonfucosylated therapeutic antibodies has also been demonstrated in the analysis using NK cells derived from breast cancer patients.54 These results showing ADCC enhancement are well reflected by in vivo analysis. The superior in vivo efficacy of nonfucosylated antibodies has also been demonstrated in the mouse model of engrafted human peripheral blood mononuclear cells (PBMC), and nonfucosylated antibodies have shown improved B-cell depletion activity in human blood analysis in vitro despite the strong inhibitory effects of human serum IgG molecules.21,42,45,51 In spite of the great benefits of nonfucosylated antibodies, currently licensed therapeutic antibodies are composed of a mixture of fucosylated and nonfucosylated immunoglobulin molecules (over 90% of the Fc-linked oligosaccharides are fucosylated) and therefore unfortunately fail to achieve optimized ADCC due to the competition between the two forms for the antigens on the target cells.21,45,51 This is the case because the construction of a robust process for the production of therapeutic antibodies lacking core fucosylation has proven to be a challenge.64,65 However, the manufacture of recombinant therapeutic antibodies fully lacking core fucosylation with a fixed quality has been achieved through gene engineering of the fucosylation pathway in the cells for antibody production.52,66–68 The application of nonfucosylated antibodies is expected to be among the most powerful and elegant approaches to improving the efficacy of therapeutic antibodies, and the use of cells with α-1,6-fucosyltransferase gene (FUT8) knockout is one of the most successful approaches for the manufacture of nonfucosylated antibodies (Potelligent technology). Nonfucosylated products manufactured using this approach have already been entered into multiple clinical trials.

CDC as an effector function of therapeutic MAbs

CDC is a cytolitic cascade mediated by a series of complement proteins abundantly present in serum. It is triggered by the binding of C1q to the constant region of cell-bound antibody molecules (Figure 3). For the induction of strong CDC activity, various biological and structural features of antigen molecules are required, such as relatively high expression, the presence of small or folded extracellular portions, or epitopes that retarget antigens into lipid rafts in the case of anti-CD20 MAbs.69,70 In addition, CDC is negatively regulated by complement-regulatory proteins (CRPs: CD46, CD55, and CD59) expressing on the cell surface.71,72 The relationship between complement activation and therapeutic activity is also suggested in several reports. For example, Kennedy and colleagues have reported rapid complement consumption after rituximab administration, and Treon and colleagues have reported upregulation of CRPs in returning tumor cells after rituximab treatment.73,74

CDC-enhancing technologies

The binding of the C1q component to the Fc, the initial step of the complement cascade, affects the intensity of the following complement activation, and several approaches have succeeded in enhancing CDC by facilitating the binding of the antibody constant region to C1q. As a result of engineered amino acid mutations inserted into either Fc or the
hinge region, designed antibody constant regions possessing improved C1q binding have been achieved.75,76

Another example of CDC enhancement is the engineering of the heavy chain by shuffling IgG1 and IgG3 sequences within a heavy chain constant region, as recently reported by our group; this is called Complegent technology (Figure 4).77 Apart from amino acid and structural idiosyncrasies, the four isotypes of human IgG differ from each other in the potencies of effector functions and other activities.41,78–82 In general, the rank orders of potency are IgG1 ≅ IgG3 ≫ IgG4 ≅ IgG2 for ADCC and IgG3 ≅ IgG1 ≫ IgG2 = IgG4 for CDC.6,56,78–81 As a result of the comparison of a comprehensive set of structural shuffling between IgG1 and IgG3, the mixed isotype heavy chain variant, wherein a part of the Fc portion of human IgG1 heavy chain is converted into the corresponding part of human IgG3 (Complegent), showed unexpectedly strong C1q binding and CDC that was even better than those of parental IgG1 and IgG3. ADCC of the Complegent antibody was unchanged from that of parent IgG1 antibody. Furthermore, an anti-CD20 antibody with Complegent-type constant regions have shown improved B-cell depletion in cynomolgus monkeys compared to anti-CD20 IgG1.77 As a result of single intravenous injection, significant enhancement in the levels of B-cell depletion was observed for Complegent-type antibody, which was manifested in a significant delay in time to recovery of B cells. This approach succeeded in enhancing CDC also in the therapeutic anti-CD52 antibody alemtuzumab, and thus may be applicable to a wide range of target molecules.

**Enhancement of multiple anti-tumor functions: The next target of antibody optimization?**
As discussed above, antibody effector functions can be manipulated by engineering amino acid sequences of the constant regions. Amino acid alterations in the Fc that possess improved FcyRIIIa binding and resultant ADCC-enhancing capacity have been extensively studied and successfully obtained by mutational or computational analyses on the Fc: FcyRIIIa contact site or by direct evolitional methods using a comprehensive display library.47,48,50,83 Similarly, binding to C1q or the neonatal Fc receptor, which is involved in IgG clearance, can also be improved by introducing amino acid mutations into the Fc or the hinge.75,76,84,85 Typically, these variant Fcs require at least 2 to 5 amino acid alterations from unmutated wild-type sequences.

One of the next directions for the optimization of therapeutic antibodies would be to combine multiple enhancing modifications into a single antibody platform. In theory, different Fc modifications can be “piled up” to create a variant sequence having accumulated amino acid mutations that possess multiple enhanced functions. Stavenhagen and colleagues have reported a successful example of an engineered Fc sequence having five amino acid mutations and
possessing enhanced binding capacity to both FcγRIIa and FcγRIIIa.50 Nevertheless, the effect of combining multiple sets of amino acid alterations would be difficult to predict, probably because the conformational change of overall antibody molecules could lead to unexpected changes in some biological activities. Actually, artificial Fc mutations have often been reported to induce unpredictable functional impairments; for example, a reduction in FcγRIIIa-binding activity is often seen for the Fc mutations that improve antibody binding to C1q, FcγRI, or FcRn.47,75,85 Notably, these unpredictable phenomena did occur even though the FcγRIIIa contact sites of IgG did not always overlap with other contact sites, especially for those of FcRn (located in the CH2:CH3 cleft), which seem to be far apart from the FcγRIIIa binding sites (located between the two lower hinges.35,86,87

Despite such technical limitations, many companies have made extensive progress in developing the next-generation anti-CD20 antibodies since the successful launch of rituximab. Now multiple new anti-CD20 antibodies with diverse effector functions are beginning to enter clinical trials (Table 1).

Many of the new anti-CD20 antibodies are designed to improve the ADCC of rituximab. Some of them might be attributable to the inherent properties of parent antibodies different from those of rituximab (clone 2B8). Interestingly, some antibodies possess engineered constant regions with increased ADCC-inducing capacity, either by mutating amino acid sequences of the Fc (AME-133) or by engineering the glycoform of the Fc-linked oligosaccharides (GA101).

**AccretaMAb platform: A universal platform technology for the next generation of therapeutic antibodies with enhanced ADCC and CDC**

An important feature of Potelligent technology is that it can be applied to a broad spectrum of antibody backbone structures. This ADCC-enhancing effect of defucosylation was initially found for human IgG1 background, and was then expanded to include various types of immunoglobulins (other human IgG subclasses or murine IgG2) and antibody-like engineered molecules (Fc fusion proteins).37,55–57,59,92 In every case examined, fucose removal did not seem to influence any biological activities other than FcγRIIIa-related functions. Although fucose removal from antibody oligosaccharides is associated with a significant thermodynamic change in the FcγRIIIa-binding state of IgG1,62 it does not significantly influence the overall IgG1 conformation except for a small change in the hydration mode around the fucose residue as revealed by crystal structure and NMR analyses.63 It is likely that the minimal conformational change allows this phenomenon to be easily applied to a wide range of antibody species.

More recently, we have shown that fucose removal from Complegent antibodies (IgG1/IgG3 chimeric isotype antibodies), in what we call the AccretaMAb platform, results in the full retention of potent ADCC and CDC, to levels that are comparable to those seen for Potelligent

<table>
<thead>
<tr>
<th>Antibody</th>
<th>Developer</th>
<th>Type</th>
<th>Parent MAb</th>
<th>Effector functions in comparison with rituximab</th>
<th>Reference</th>
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<tbody>
<tr>
<td>Ofatumumab</td>
<td>GSK/Genmab</td>
<td>IgG1</td>
<td>2F2</td>
<td>Potent CDC</td>
<td>88</td>
</tr>
<tr>
<td>Ocrelizumab</td>
<td>Genentech</td>
<td>IgG1</td>
<td>2H7</td>
<td>Potent ADCC and decreased CDC</td>
<td>89</td>
</tr>
<tr>
<td>Veltuzumab</td>
<td>Immunomedics</td>
<td>IgG1</td>
<td>2B8</td>
<td>Comparable to rituximab</td>
<td>90</td>
</tr>
<tr>
<td>AME-133</td>
<td>AME</td>
<td>Mutated IgG1</td>
<td>Synthetic</td>
<td>Potent ADCC</td>
<td>83</td>
</tr>
<tr>
<td>GA101</td>
<td>Roche/GlycArt</td>
<td>Glycoengineered IgG1</td>
<td>B1</td>
<td>Potent ADCC/direct apoptosis, no CDC</td>
<td>91</td>
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**Table 1 Effector functions of selected next-generation anti-CD20 MAbs**
or Complement alone, respectively. The successful combination of ADCC- and CDC-enhancing modifications in the AccretaMAb platform is constituted by two essential structural factors: (i) IgG1/IgG3 mixed constant regions that enhance C1q binding and CDC activity, and (ii) nonfucosylated oligosaccharides linked to the Fc of the mixed variant constant regions that maximize its ADCC.

As for the structural basis of AccretaMAb, how the IgG1/IgG3 chimeric isotype sequence greatly enhances CDC activity remains elusive. Interestingly, in addition to the hinge derived from IgG1 and the CH2 from IgG3, the other constant domains (the CH1 from IgG1 and the CH3 from IgG3) are also required for its optimized CDC activity. The CH1 and the CH3 domains are not considered direct contact sites between IgG:C1q, as revealed by a comprehensive structural shuffling analysis between IgG1 and IgG3. Thus it appears likely that the specific combinations of multiple IgG domains of different subclasses render the overall variant antibody conformation capable of binding C1q with unexpectedly strong affinity. Importantly, the FcγRIIIa binding and the ADCC of this novel chimeric constant region are as strong as those of IgG1, and can be further maximized by the removal of fucose from Asn297-linked oligosaccharides.

**Clinical implication of engineered antibodies with multiple-enhanced functions**

It has been suggested that tumor cells avoid attack from therapeutic antibodies by using various mechanisms, which might lead to insufficient efficacy in the clinic (Table 2).

Thus, in order to overcome the diverse mechanisms underlying clinical resistance by tumor cells, the enhancement of multiple functions of therapeutic antibodies might be a promising approach. Regarding AccretaMAb, for example, the dual enhancement of effector functions (ADCC and CDC) is expected to bring some additional advantages; these two functions might act complementarily. For instance, tumor cells that are highly resistant to attack by complementary regulatory proteins would be destroyed more effectively by potent ADCC due to defucosylated Fc-linked oligosaccharides. Conversely, in circumstances where the accessibility of ADCC effector cells is severely limited by tight solid tumor tissues, the potent CDC of AccretaMAb, which does not require cellular factors, would be a powerful weapon for eliminating tumor cells.

Furthermore, the increased complement fixing and enhanced ADCC activities might have a synergistic effect through an interaction between complement opsonized on the target cells and complement receptors (eg, CR3) present on effector cells, which would increase ADCC of antibodies. However, the clinical importance of this mechanism is far less validated and remains to be elucidated.

One of the potential issues in future therapy with artificially engineered antibodies is immunogenicity; namely, the use of mutated sequences may increase the risk of newly generated non-self peptides that bind to MHC class II molecules, especially for those having a higher number of accumulated amino acid alterations. Interestingly, the AccretaMAb platform is fully composed of natural structural components: IgG1 and IgG3 sequences and nonfucosylated N-linked oligosaccharides (constitutes ~10% of human serum IgG oligosaccharides). Future clinical trials of AccretaMAb will also confirm the hypothesis that the variant constant regions consisting of fully natural components would render the antibodies potentially less immunogenic compared to artificial mutation approaches.

**Perspectives**

After a number of successes and failures in the lab, engineered antibodies with optimized effector functions now

**Table 2 Proposed mechanisms of resistance by tumor cells to antibody therapy**

<table>
<thead>
<tr>
<th>(i) Tumor factors</th>
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<tr>
<td>- Poor penetration of effector cells of ADCC into solid tumor tissues</td>
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<tr>
<td>- Complement-regulatory proteins (CD46, CD55, and CD59) on the tumor cells</td>
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<tr>
<td>- Soluble antigens</td>
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<td>- Altered signaling (eg, bcl-2 elevation in rituximab therapy)</td>
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<th>(ii) Host factors</th>
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<tr>
<td>- Heterogeneity in ADCC according to FcγRIIIa functional polymorphism</td>
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<tr>
<td>- Inactivation of effector cells in patients (eg, downregulation of ζ chain, NKG2D, NKP30, NKP44, or NKP46 in NK cells)</td>
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<th>(iii) Therapy factors</th>
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<tr>
<td>- Suppression of effector cells by chemotherapy</td>
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It is obvious that the choice of target functions of an antibody to be enhanced should be determined carefully based on a biological and mechanistic understanding of the target tumors and antigens, since various cancer subtypes of different origins display diverse biological profiles and resultant mechanisms of clinical resistance. Yet the major issue of current antibody therapy remains insufficient efficacy. Therefore, the expanding technologies that enhance multiple anti-tumor mechanisms of therapeutic antibodies would be of great value for the development of third-generation antibody therapeutics as long as they are used in the right patients, with the right dosage and on the right schedule, and perhaps monitored with appropriate biomarkers.

Disclosure
The authors report no conflicts of interest in this work.

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