(54) Titre : ANTICORPS ANTI-GLYPICAN-3 DONT LA CINETIQUE DANS LE PLASMA EST AMELIOREE
(54) Title: ANTI-GLYPICAN-3 ANTIBODY HAVING IMPROVED KINETICS IN PLASMA

(57) Abrégé/Abstract:
[PROBLEMS] To provide: a method for controlling the half-life of an anti-glypican-3 antibody in the plasma; a pharmaceutical composition comprising an anti-glypican-3 antibody having a controlled half-life in the plasma as an active ingredient; a method for
producing the anti-glypican-3 antibody; and a method for producing a pharmaceutical composition comprising the anti-glypican-3 antibody as an active ingredient. [MEANS FOR SOLVING PROBLEMS] Disclosed are: a method for controlling the half-life of an anti-glypican-3 antibody in the plasma by modifying an amino acid residue exposed on the surface of the anti-glypican-3 antibody; an anti-glypican-3 antibody whose half-life in the plasma is controlled by the modification of an amino acid residue; a pharmaceutical composition comprising the antibody as an active ingredient; and a method for producing the pharmaceutical composition.
ABSTRACT OF THE DISCLOSURE:

A method of modulating the plasma half-life of anti-glypican 3 antibody, a pharmaceutical composition comprising as an effective component the anti-glypican 3 antibody that has a plasma half-life that has been modulated, a method of preparing the anti-glypican 3 antibody and a pharmaceutical composition comprising the anti-glypican 3 antibody as an effective component are provided. Disclosed is a method of modulating the plasma half-life of anti-glypican 3 antibody by modifying an amino acid residue that is exposed on the surface of the anti-glypican 3 antibody; and anti-glypican 3 antibody that has a plasma half-life that has been modulated by amino acid residue modification, a pharmaceutical composition comprising as an effective component the anti-glypican 3 antibody, and a method of preparing the anti-glypican 3 antibody and producing a pharmaceutical composition comprising the anti-glypican 3 antibody as an effective component.
DEMANDE OU BREVET VOLUMINEUX

LA PRÉSENTE PARTIE DE CETTE DEMANDE OU CE BREVET COMPREND PLUS D’UN TOME.

CECI EST LE TOME 1 DE 2
CONTENANT LES PAGES 1 À 137

NOTE : Pour les tomes additionnels, veuillez contacter le Bureau canadien des brevets

JUMBO APPLICATIONS/PATENTS

THIS SECTION OF THE APPLICATION/PATENT CONTAINS MORE THAN ONE VOLUME

THIS IS VOLUME 1 OF 2
CONTAINING PAGES 1 TO 137

NOTE: For additional volumes, please contact the Canadian Patent Office

NOM DU FICHIER / FILE NAME :

NOTE POUR LE TOME / VOLUME NOTE:
ANTI-GLYPCAN-3 ANTIBODY HAVING IMPROVED KINETICS IN PLASMA

BACKGROUND OF THE INVENTION

Related Applications

This application claims priority of Japanese Patent Application No. 2007-256063, filed on September 28, 2007, the contents of which are herein incorporated by reference.

Field of the Invention

The present invention relates to a method of improving the plasma (blood) kinetics of anti-glypican 3 antibodies, a pharmaceutical composition comprising an anti-glypican 3 antibody that has improved plasma kinetics as an effective component, and a method of preparing the same.

Description of the Related Art

Antibodies are stable in the blood and exhibit few side effects and for these reasons their use as drugs has been receiving attention. Among the several antibody isotypes, a large number of IgG isotype therapeutic antibodies are on the market and a large number of therapeutic antibodies are also currently under development (Janice M. Reichert, Clark J. Rosensweig, Laura B. Faden, and Matthew C. Dewitz, Monoclonal antibody successes in the clinic, Nature Biotechnology (2005) 23, 1073-8; Pavlou A. K. and Belsey M. J., The therapeutic

In addition, technologies to enhance the effector functions are being developed for producing second-generation therapeutic antibodies. For example, it is known that the antibody-dependent cellular cytotoxicity (ADCC) activity and the complement-dependent cytotoxicity (CDC) activity are enhanced by an amino acid substitution in which the amino acids constituting the Fc region of IgG isotype antibodies (referred to as IgG antibodies) are replaced by different amino acids (Kim S. J., Park Y., and Hong H. J., Antibody engineering for the development of therapeutic antibodies, Mol. Cells (2005) 20(1), 17-29). When an anti-glypican 3 antibody is produced in fucose transporter-deleted CHO cells, fucose is not attached to the sugar chains attached to the anti-glypican
3 antibody. Such an anti-glypican 3 antibody has a significantly higher ADCC activity than the anti-glypican 3 antibody that contains fucose in the branched-chain of the sugar chain, and is thought to exhibit a greater antitumor activity as a therapeutic antibody (WO 2006/067913).

In addition to such technologies for enhancing the effector functions, other technologies are also known in which the plasma half-life of an antibody is increased or decreased by amino acid substitution on the amino acids constituting the Fc region of the antibody (Hinton P. R., Xiong J. M., Johlfis M. G., Tang M. T., Keller S., and Tsurushita N., An engineered human IgGl antibody with longer serum half-life, J. Immunol. (2006) 176(1), 346-56; and Ghetie V., Popov S., Borvak J., Radu C., Matesoi D., Medesan C., Ober R. J., and Ward E. S., Increasing the serum persistence of an IgG fragment by random mutagenesis, Nat. Biotechnol. (1997) 15(7), 637-40). If a technology that prolongs the plasma half-life of antibodies is applied to therapeutic antibodies, it is expected that the dose of the administered therapeutic antibody is reduced and its interval of administration is extended, which will enable the provision of less expensive therapeutic antibodies with a high convenience factor.

In specific terms, the plasma half-life can be extended by substituting an amino acid of the Fc region of an IgG antibody with another amino acid resulting in improving the IgG antibody’s affinity for the neonatal Fc receptor, which is
known to be a salvage receptor for the IgG antibody. In addition, it is also known that the plasma half-life is increased by shuffling the individual domains (CH1, CH2, CH3) constituting the constant region of the antibody (Zuckier L. S., Chang C. J., Scharff M. D., and Morrison S. L., Chimeric human-mouse IgG antibodies with shuffled constant region exons demonstrate that multiple domains contribute to in vivo half-life, Cancer Res. (1998) 58(17), 3905-8). However, since the amino acid sequence of the constant region of the IgG antibody is conserved in humans, an antibody having an artificial amino acid substitution as described above in the amino acids constituting the constant region may cause side effects by exhibiting immunogenicity in the human body. It is therefore preferred that only a small number of amino acids be substituted.

Technologies involving amino acid substitution in the variable region (also referred to as V region) of IgG antibodies reported to date include humanization technology (Tsurushita N., Hinton P. R., and Kumar S., Design of humanized antibodies: from anti-Tac to Zenapax, Methods (2005) 36(1), 69-83), affinity maturation where amino acids in the complementarity-determining region (CDR) is substituted in order to increase the binding activity (Rajpal A., Beyaz N., Haber L., Cappuccilli G., Yee H., Bhatt R. R., Takeuchi T., Lerner R. A., and Crea R., A general method for greatly improving the affinity of antibodies by using combinatorial
libraries, *Proc. Natl. Acad. Sci. USA* (2005) 102(24), 8466-7) and amino acid substitution in the amino acids constituting the framework region (FR) for improving the physicochemical stability (Ewert S., Honegger A., and Pluckthun A., Stability improvement of antibodies for extracellular and intracellular applications: CDR grafting to stable frameworks and structure-based framework engineering, *Methods* (2004) 34(2), 184-99). Unlike the case with amino acid substitution in the constant region (also referred to as C region), amino acid substitution in the variable region is generally used for improving the characteristics (e.g., stability) and enhancing the function (e.g., antigen binding activity) of antibodies. Since the amino acid sequence constituting the CDR of humanized antibodies is derived from the amino acid sequence of a nonhuman animal species, the risk of generating immunogenicity by introducing an artificial amino acid substitution in this sequence is thought to be lower than amino acid substitutions in a sequence in other regions. Moreover, with regard to an artificial amino acid substitution in the amino acid sequence constituting the FR of humanized antibodies, it is thought that such a substitution poses little risk of generating immunogenicity if the FR amino acid sequence obtained as a consequence of substitution is the same as any of the plurality of human antibody FR amino acid sequences that are published in, for example, the Kabat database (http://ftp.ebi.ac.uk/pub/databases/kabat/), the IMGT database
(http://imgt.cines.fr/), and so forth. Furthermore, the immunogenicity can be reduced by reselecting a human antibody sequence that is very similar to the FR amino acid sequence obtained as a consequence of substitution, from the plurality of human antibody FR amino sequences that are published in the Kabat database, the IMGT database, and so forth (WO 1999/018212).

In contrast, the only methods known for improving the plasma half-life of IgG antibodies are, as described above, amino acid substitution of amino acids constituting the Fc region, which is a part of the constant region, and no methods have been reported to date that bring about an improvement in the plasma half-life of IgG antibodies by amino acid substitution of the amino acids constituting the variable region, which is believed to carry little risk of invoking immunogenicity. The reason for this is, in part, that the plasma half-life of IgG antibodies are believed to largely depend on antigen-dependent depletion and binding to the neonatal Fc receptor, a salvage receptor for IgG antibodies (Lobo E. D., Hansen R. J., and Balthasar J. P., Antibody pharmacokinetics and pharmacodynamics, J. Pharm. Sci. (2004) 93(11), 2645-68), and that the functions and properties of the variable region may not have a significant influence on the plasma half-life.

It has also been reported that the isoelectric point (pI) of IgG antibody is lowered by arionization of IgG antibody by
succinylation (Yamasaki Y., Sumimoto K., Nishikawa M., Yamashita F., Yamaoka K., Hashida M., and Takakura Y., Pharmacokinetic analysis of in vivo disposition of succinylated proteins targeted to liver nonparenchymal cells via scavenger receptors: importance of molecular size and negative charge density for in vivo recognition by receptors, Pharmacol. Exp. Ther. (2002) 301(2), 467-77); and that the pI of IgG antibody is raised by cationization of the IgG antibody by modification with polyamine (Poduslo J. F. and Curran G. L., Polyamine modification increases the permeability of proteins at the blood-nerve and blood-brain barriers, Neurochem. (1996) 66(4), 1599-609). However, in both cases there was no increase in the plasma half-life of the modified IgG antibody, but rather the plasma half-life was decreased. Thus, an increase in the plasma half-life of IgG antibodies cannot be realized by modification of the pI of the IgG antibody by the above-described chemical modification of the IgG antibody.

SUMMARY OF THE INVENTION

The present invention was pursued in view of the circumstances described above. An object of the present invention is to provide a method of modulating the plasma (blood) half-life of anti-glypican 3 antibody, an anti-glypican 3 antibody having a modulated plasma half-life and a pharmaceutical composition comprising the antibody as an effective component, as well as a method of preparing the
anti-glypican 3 antibody and the pharmaceutical composition. Another object of the present invention is to provide a method for modulating cytotoxicity of an antibody by modulating the plasma half-life of the antibody having cytotoxicity, an antibody with modulated cytotoxicity and a pharmaceutical composition, comprising the antibody, as well as a method of preparing the antibody and the pharmaceutical composition.

The present inventors carried out focused investigations into methods for modulating the plasma half-life of an antibody (e.g. anti-glypican 3 antibody). As a result, the present inventors discovered that the plasma half-life of an antibody (e.g. anti-glypican 3 antibody) can be modulated by modifying - among the amino acid residues constituting the variable region and the constant region of an antibody (e.g. anti-glypican 3 antibody) - amino acid residues exposed on the surface of this antibody molecule and thereby controlling the surface charge of the antibody molecule. Specifically, among the amino acid residues in the amino acid sequence constituting the variable region and the constant region of an antibody (e.g. anti-glypican 3 antibody), particular amino acid residues were identified that can modulate the plasma half-life of the antibody (e.g. anti-glypican 3 antibody) through modifying the surface charge on the antibody molecule without affecting the structure or function of the antibody, e.g., the antigen binding activity. The present inventors also confirmed that an antibody (e.g. anti-glypican 3 antibody)
having a half-life modulated in this manner in fact retains its antigen binding activity. The present inventors also found that modulation of the plasma half-life of an antibody (e.g. anti-glypican 3 antibody) increases the tumor proliferation inhibiting activity on cancer cells exhibited by cytotoxic antibodies, such as the anti-glypican 3 antibody.

The present invention relates to a method of modulating the plasma half-life of an antibody (e.g. anti-glypican 3 antibody) by modifying an amino acid residue that is exposed on the surface of the antibody, an antibody (e.g. anti-glypican 3 antibody) that has a modulated plasma half-life by amino acid residue modification, a pharmaceutical composition comprising the antibody as an effective component, and a method of preparing such a pharmaceutical composition. More specifically, the present invention provides the following:

[1] A method for preparing an anti-glypican 3 antibody with modulated plasma kinetics, said method comprising the steps of:
(a) culturing a host cell bearing a nucleic acid that encodes the anti-glypican 3 antibody under conditions allowing for expression of the nucleic acid, wherein the anti-glypican 3 antibody has an amino acid sequence altered to causes a modification in the charge of at least one amino acid residue that can be exposed on the surface of the antibody; and
(b) recovering the anti-glypican 3 antibody from the host cell culture;
[2] The method according to [1], wherein the modulation of the plasma kinetics is increase or decrease in a parameter selected from the plasma half life, the mean plasma residence time, and the plasma clearance;

[3] The method according to [1], wherein the modification in the charge of the amino acid residue is achieved by amino acid substitution;

[4] The method according to [1], wherein the amino acid residue that can be exposed on the surface of the anti-glypican 3 antibody is located in a region in the anti-glypican 3 antibody other than the FcRn binding region;

[5] The method according to [4], wherein the FcRn binding region comprises the Fc region;

[6] The method according to [4], wherein the FcRn binding region comprises the amino acid residues of the EU numbers 250, 253, 310, 311, 314, 428, 435, 436 according to the Kabat numbering;

[7] The method according to [1], wherein the anti-glypican 3 antibody is an IgG antibody;

[8] The method according to [1]-[7], wherein the amino acid residue whose charge is modified is an amino acid residue present in the heavy chain variable region or the light chain variable region;

[9] The method according to [8], wherein the anti-glypican 3 antibody comprises a complementarity-determining region (CDR) derived from a non-human animal, a framework region (FR)
derived from human, and a constant region derived from human, and wherein the modification in the charge of the amino acid residue is achieved by substitution of at least one amino acid residue that can be exposed on the antibody surface in the CDR or FR of the antibody with an amino acid residue that has a charge different from that of the amino acid residue;

[10] The method according to [9], wherein the modification in the charge of the amino acid residue is achieved by:
(1) at least one substitution in the heavy chain variable region shown in SEQ ID NO: 1 selected from:
(a) substitution of Q that is the 43rd amino acid residue with K,
(b) substitution of D that is the 52nd amino acid residue with N, and
(c) substitution of Q that is the 107th amino acid residue with R;
and/or
(2) at least one substitution in the light chain variable region shown in SEQ ID NO: 7 selected from:
(d) substitution of E that is the 17th amino acid residue with Q,
(e) substitution of Q that is the 27th amino acid residue with R, and
(f) substitution of Q that is the 105th amino acid residue with R;

[11] The method according to [9], wherein the modification in
the charge of the amino acid residue is achieved by:

(1) at least one substitution in the heavy chain variable region shown in SEQ ID NO: 1 selected from:
(a) substitution of K that is the 19th amino acid residue with T,
(b) substitution of Q that is the 43rd amino acid residue with E,
(c) substitution of Q that is the 62nd amino acid residue with E,
(d) substitution of K that is the 63rd amino acid residue with S,
(e) substitution of K that is the 65th amino acid residue with Q, and
(f) substitution of G that is the 66th amino acid residue with D;
and/or

(2) at least one substitution in the light chain variable region shown in SEQ ID NO: 7 selected from:
(g) substitution of R that is the 24th amino acid residue with Q,
(h) substitution of Q that is the 27th amino acid residue with E,
(i) substitution of K that is the 79th amino acid residue with T,
(j) substitution of R that is the 82nd amino acid residue with S, and
(k) substitution of K that is the 112nd amino acid residue with E;

[12] The method according to [11], further comprising at least one modification in the heavy chain constant region shown in SEQ ID NO: 31 selected from:

(a) substitution of H that is the 151st amino acid residue with Q,

(b) substitution of K that is the 157th amino acid residue with Q,

(c) substitution of R that is the 238th amino acid residue with Q,

(d) substitution of D that is the 239th amino acid residue with E,

(e) substitution of L that is the 241st amino acid residue with M, and

(f) substitution of Q that is the 302nd amino acid residue with E;

[13] The method according to [9]-[12], wherein the anti-glypican 3 antibody has a reduced content of fucose attached to the Fc region of the antibody;

[14] An anti-glypican 3 antibody prepared by the method according to [1]-[13];

[15] A method for preparing an antibody with modulated plasma kinetics, said method comprising the steps of:

(a) culturing a host cell bearing a nucleic acid that encodes the antibody under conditions allowing for expression of the
nucleic acid, wherein the antibody has an amino acid sequence altered to causes a modification in the charge of at least one amino acid residue in the constant region in the antibody other than the FcRn binding region; and
(b) recovering the antibody from the host cell culture;

[16] The method according to [15], wherein the modulation of the plasma kinetics is increase or decrease in a parameter selected from the plasma half life, the mean plasma residence time, and the plasma clearance;

[17] The method according to [15], wherein the modification in the charge of the amino acid residue is achieved by amino acid substitution;

[18] The method according to [17], wherein the antibody is an IgG antibody;

[19] The method according to [18], wherein the antibody is an IgG1 antibody;

[20] The method according to [17], wherein the modification in the charge of the amino acid residue is achieved by substitution of at least one amino acid residue of an IgG1 antibody with a corresponding amino acid residue of an IgG4 antibody;

[21] The method according to [15]-[20], wherein the FcRn binding region comprises the amino acid residues of the EU numbers 250, 253, 310, 311, 314, 428, 435, and 436 according to the Kabat numbering;
[22] The method according to [20], wherein the modification in the charge of the amino acid residue is achieved by at least one substitution in the heavy chain constant region shown in SEQ ID NO: 31 selected from:

(a) substitution of H that is the 151st amino acid residue with Q,
(b) substitution of K that is the 157th amino acid residue with Q,
(c) substitution of R that is the 238th amino acid residue with Q,
(d) substitution of D that is the 239th amino acid residue with E,
(e) substitution of L that is the 241st amino acid residue with M, and
(f) substitution of Q that is the 302nd amino acid residue with E;

[23] The method according to [15]-[22], wherein the antibody is an anti-glypican 3 antibody;

[24] A method of stabilizing an anti-glypican 3 antibody that comprises a complementarity-determining region (CDR) derived from a non-human animal, a framework region (FR) derived from human, and a constant region derived from human, said method comprising the steps of:

(a) culturing a host cell bearing a nucleic acid that encodes the anti-glypican 3 antibody under conditions allowing for expression of the nucleic acid, wherein the anti-glypican 3
antibody has an amino acid sequence altered to increase in the Tm value of the antibody by a modification of at least one amino acid residue; and

(b) recovering the antibody from the host cell culture;

[25] The method according to [24], wherein the amino acid residue is present in the FR1 region and/or the FR2 region of the heavy chain or the light chain;

[26] The method according to [25], wherein an amino acid residue in the FR2 region of the heavy chain is substituted with an amino acid residue of the FR2 region of the VH4 subclass;

[27] The method according to [25], wherein an amino acid residue in the FR2 region of the light chain is substituted with an amino acid residue of the FR2 region of the VK3 subclass;

[28] The method according to [24]-[27], wherein the substitution of the amino acid residue is achieved by:

(1) at least one substitution in the heavy chain variable region shown in SEQ ID NO: 1 selected from:

(a) substitution of V that is the 37th amino acid residue with I,

(b) substitution of A that is the 40th amino acid residue with P,

(c) substitution of M that is the 48th amino acid residue with I, and
(d) substitution of L that is the 51st amino acid residue with I;

and/or

(2) at least one substitution in the light chain variable region shown in SEQ ID NO: 7 selected from:

(e) substitution of L that is the 42nd amino acid residue with Q,

(f) substitution of S that is the 48th amino acid residue with A, and

(g) substitution of Q that is the 50th amino acid residue with R;

[29] A method for preparing an antibody with modulated cytotoxicity, comprising the steps of:

(a) culturing a host cell bearing a nucleic acid that encodes the antibody under conditions allowing for expression of the nucleic acid, wherein the antibody has an amino acid sequence altered to causes a modification in the charge of at least one amino acid residue that can be exposed on the surface of a cytotoxic antibody; and

(b) recovering the antibody from the host cell culture;

[30] The method according to [29], wherein the modification in the charge of the amino acid residue is achieved by amino acid substitution;

[31] The method according to [29], wherein the amino acid residue that can be exposed on the surface of the antibody is located in a region in the antibody other than the FcRn
binding region;

[32] The method according to [31], wherein the FcRn binding region comprises the Fc region;

[33] The method according to [31], wherein the FcRn binding region comprises the amino acid residues of the EU numbers 250, 253, 310, 311, 314, 428, 435, 436 according to the Kabat numbering;

[34] The method according to [29], wherein the antibody is an IgG antibody;

[35] The method according to [29]-[34], wherein the amino acid residue whose charge is modified is an amino acid residue present in the constant region of the antibody;

[36] The method according to [29]-[34], wherein the amino acid residue whose charge is modified is an amino acid residue present in the heavy chain variable region or the light chain variable region of the antibody;

[37] The method according to [36], wherein the antibody is an antibody that comprises a complementarity-determining region (CDR) derived from a non-human animal, a framework region (FR) derived from human, and a constant region derived from human, and wherein the modification in the charge of the amino acid residue is achieved by substitution of at least one amino acid residue that can be exposed on the antibody surface in the CDR or FR of the antibody with an amino acid residue that has a charge different from that of the amino acid residue;
[38] The method according to [37], wherein the modification in the charge of the amino acid residue is achieved by:

(1) at least one substitution in the heavy chain variable region shown in SEQ ID NO: 1 selected from:

(a) substitution of K that is the 19th amino acid residue with T,
(b) substitution of Q that is the 43rd amino acid residue with E,
(c) substitution of Q that is the 62nd amino acid residue with E,
(d) substitution of K that is the 63rd amino acid residue with S,
(e) substitution of K that is the 65th amino acid residue with Q, and
(f) substitution of G that is the 66th amino acid residue with D;

and/or,

(2) at least one substitution in the light chain variable region shown in SEQ ID NO: 7 selected from:

(g) substitution of R that is the 24th amino acid residue with Q,
(h) substitution of Q that is the 27th amino acid residue with E,
(i) substitution of K that is the 79th amino acid residue with T,
(j) substitution of R that is the 82nd amino acid residue with S, and

(k) substitution of K that is the 112nd amino acid residue with E;

[39] The method according to [38], further comprising at least one substitution in the heavy chain constant region shown in SEQ ID NO: 31 selected from:

(a) substitution of H that is the 151st amino acid residue with Q,

(b) substitution of K that is the 157th amino acid residue with Q,

(c) substitution of R that is the 238th amino acid residue with Q,

(d) substitution of D that is the 239th amino acid residue with E,

(e) substitution of L that is the 241st amino acid residue with M, and

(f) substitution of Q that is the 302nd amino acid residue with E;

[40] The method according to [36], wherein the antibody comprises a complementarity-determining region (CDR) derived from a non-human animal, a framework region (FR) derived from human, and a constant region derived from human, and wherein the modification in the charge of the amino acid residue is achieved by substitution of at least one amino acid residue that can be exposed on the antibody surface in the constant
region of the antibody with an amino acid residue that has a charge different from that of the amino acid residue;

[41] The method according to [40], wherein the substitution is at least one substitution in the heavy chain constant region shown in SEQ ID NO: 31 selected from:

(a) substitution of H that is the 151st amino acid residue with Q,

(b) substitution of K that is the 157th amino acid residue with Q,

(c) substitution of R that is the 238th amino acid residue with Q,

(d) substitution of D that is the 239th amino acid residue with E,

(e) substitution of L that is the 241st amino acid residue with M, and

(f) substitution of Q that is the 302nd amino acid residue with E;

[42] The method according to [37]-[41], wherein the antibody has a reduced content of fucose attached to the Fc region of the antibody;

[43] An antibody prepared by the method according to [29]-[42];

[44] The antibody according to [43], wherein the antibody is an anti-glypican 3 antibody;

[45] An antibody comprising:

(1) a heavy chain variable region shown in SEQ ID NO: 1 in
which the amino acid sequence comprises at least one
substitution selected from:

(a) substitution of K that is the 19th amino acid residue with T,
(b) substitution of Q that is the 43rd amino acid residue with E,
(c) substitution of Q that is the 62nd amino acid residue with E,
(d) substitution of K that is the 63rd amino acid residue with S,
(e) substitution of K that is the 65th amino acid residue with Q, and
(f) substitution of G that is the 66th amino acid residue with D;

and/or

(2) a light chain variable region shown in SEQ ID NO: 7 in
which the amino acid sequence comprises at least one
substitution selected from:

(g) substitution of R that is the 24th amino acid residue with Q,
(h) substitution of Q that is the 27th amino acid residue with E,
(i) substitution of K that is the 79th amino acid residue with T,
(j) substitution of R that is the 82nd amino acid residue with S, and
(k) substitution of K that is the 112nd amino acid residue with E;

[46] The antibody according to [45], comprising the heavy chain shown in SEQ ID NO: 3 and the light chain shown in SEQ ID NO: 9;

[47] The antibody according to [45], comprising the heavy chain shown in SEQ ID NO: 5 and the light chain shown in SEQ ID NO: 11;

[48] The antibody according to [45] which comprises a heavy chain variable region shown in SEQ ID NO: 27 and a light chain variable region shown in SEQ ID NO: 28;

[49] The antibody according to [45] which comprises a heavy chain variable region shown in SEQ ID NO: 27 and a light chain variable region shown in SEQ ID NO: 29;

[50] The antibody according to [45]-[49] comprising a constant region of a human antibody;

[51] The antibody according to [50], wherein the constant region comprises a sequence shown in SEQ ID NO: 32 or SEQ ID NO: 33;

[52] An antibody comprising:

(1) a heavy chain variable region shown in SEQ ID NO: 1 in which the amino acid sequence comprises at least one substitution selected from:

(a) substitution of Q that is the 43rd amino acid residue with K,

(b) substitution of D that is the 52nd amino acid residue with
(c) substitution of Q that is the 107th amino acid residue with R;

and

(2) a light chain variable region shown in SEQ ID NO: 7 in which the amino acid sequence comprises at least one substitution selected from:

(d) substitution of E that is the 17th amino acid residue with Q,

(e) substitution of Q that is the 27th amino acid residue with R, and

(f) substitution of Q that is the 105th amino acid residue with R;

[53] The antibody according to [52], comprising the heavy chain variable region shown in SEQ ID NO: 4 and the light chain variable region shown in SEQ ID NO: 10;

[54] The antibody according to [52], comprising the heavy chain variable region shown in SEQ ID NO: 6 and the light chain variable region shown in SEQ ID NO: 12;

[55] The antibody according to [52]-[54] comprising a constant region of a human antibody;

[56] An antibody comprising at least one substitution in the amino acid sequence of the heavy chain constant region shown in SEQ ID NO: 31 selected from:

(a) substitution of H that is the 151st amino acid residue with Q,
(b) substitution of K that is the 157th amino acid residue with Q,
(c) substitution of R that is the 238th amino acid residue with Q,
(d) substitution of D that is the 239th amino acid residue with E,
(e) substitution of L that is the 241st amino acid residue with M, and
(f) substitution of Q that is the 302nd amino acid residue with E;

[57] An antibody comprising a heavy chain constant region shown in SEQ ID NO: 33;

[58] The antibody according to [45]-[57], wherein the antibody has a reduced content of fucose attached to the Fc region of the antibody;

[59] A composition comprising the antibody according to [45]-[58] and a pharmaceutically acceptable carrier.

[60] An anticancer agent comprising as an effective component the antibody according to [45]-[58];

[61] The anticancer agent according to [60], wherein the cancer is liver cancer;

[62] A nucleic acid that encodes a polypeptide of the antibody according to [45]-[58];

[63] A host cell comprising the nucleic acid according to [62];

[64] The host cell according to [63], wherein the host cell is
a fucose transporter-deficient animal cell, a
fucosyltransferase-deleted animal cell, or an animal cell in
which a complex branched sugar chain modification is modified;
[65] A method for preparing an antibody comprising culturing
the host cell according to [63] or [64] and recovering a
polypeptide from the cell culture.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 is a chart obtained from the differential
scanning calorimetric (DSC) measurement of the Hspu2.2Lspu2.2
(Hu2.2Lu2.2) antibody;

Figure 2 is an electrophoretogram of HOLO antibody and
Hspu2.2Lspu2.2 (Hu2.2Lu2.2) antibody in high pI isoelectric
electrophoresis, wherein lanes 1 and 4 show pI markers, lane 2
shows HOLO antibody, and lane 3 shows Hspu2.2Lspu2.2
(Hu2.2Lu2.2) antibody, where the numerical values show the pI
values of the pI marker molecules and the arrows show the
electrophoretic mobilities of the corresponding pI marker
molecules;

Figure 3 is an electrophoretogram of HOLO antibody and
Hspd1.8Lspd1.6 (Hd1.8Ld1.6) antibody in low pI isoelectric
electrophoresis, wherein lanes 1 and 4 show pI markers, lane 2
shows HOLO antibody, and lane 3 shows Hspd1.8Lspd1.6
(Hd1.8Ld1.6) antibody, where the numerical values show the pI
values of the pI marker molecules and the arrows show the
electrophoretic mobilities of the corresponding pI marker
molecules;

Figure 4 is a diagram that shows the binding affinity of H15L4 antibody and HOLO antibody for glypican 3 antigen in competitive ELISA, wherein the black diamond refers to the binding affinity of the HOLO antibody and the grey square refers to the binding affinity of the H15L4 antibody;

Figure 5 is a diagram that shows the binding affinity of Hspu2.2Lspu2.2 (Hu2.2Lu2.2) antibody and HOLO antibody for glypican 3 antigen in competitive ELISA, wherein the black diamond refers to the binding affinity of the HOLO antibody and the grey square refers to the binding affinity of the Hspu2.2Lspu2.2 (Hu2.2Lu2.2) antibody;

Figure 6 is a diagram that shows the binding affinity of Hspd1.8Lspd1.6 (Hd1.8Ld1.6) antibody and HOLO antibody for glypican 3 antigen in competitive ELISA, wherein the black diamond refers to the binding affinity of the HOLO antibody and the grey square refers to the binding affinity of the Hspd1.8Lspd1.6 (Hd1.8Ld1.6) antibody;

Figure 7 shows the antitumor activity in a human liver cancer-transplant mouse model of the HOLO antibody, Hspu2.2Lspu2.2 (Hu2.2Lu2.2) antibody, and Hspd1.8Lspd1.6 (Hd1.8Ld1.6) antibody;

Figure 7A shows the antitumor activity in a human liver cancer-transplant mouse model of the HOLO antibody, Hspu2.2Lspu2.2 (Hu2.2Lu2.2) antibody, and Hspd1.8Lspd1.6 (Hd1.8Ld1.6) antibody when each test antibody was administered
to the model at a dose of 5 mg/kg, wherein the black diamond shows the activity for the administration of vehicle, the black triangle shows the effect of the administration of the Hspd1.8Lspd1.6 (Hdl.8Ld1.6) antibody, the white circle shows the effect of the administration of the Hspu2.2Lspu2.2 (Hu2.2Lu2.2) antibody, and the black square shows the effect of the administration of the HOLO antibody;

Figure 7B shows the antitumor activity in a human liver cancer-transplant mouse model of the HOLO antibody, Hspu2.2Lspu2.2 (Hu2.2Lu2.2) antibody, and Hspd1.8Lspd1.6 (Hdl.8Ld1.6) antibody when each test antibody was administered to the model at a dose of 1 mg/kg, wherein the black diamond shows the activity for the administration of vehicle, the black triangle shows the effect of the administration of the Hspd1.8Lspd1.6 (Hdl.8Ld1.6) antibody, the white circle shows the effect of the administration of the Hspu2.2Lspu2.2 (Hu2.2Lu2.2) antibody, and the black square shows the effect of the administration of the HOLO antibody;

Figure 8 shows the blood antibody concentrations in a human liver cancer-transplant mouse model of the HOLO antibody, Hspu2.2Lspu2.2 (Hu2.2Lu2.2) antibody, and Hspd1.8Lspd1.6 (Hdl.8Ld1.6) antibody;

Figure 8A shows the blood concentration of the antibody administered to a human liver cancer-transplant mouse model of the HOLO antibody, Hspu2.2Lspu2.2 (Hu2.2Lu2.2) antibody, and Hspd1.8Lspd1.6 (Hdl.8Ld1.6) antibody when each test antibody
was administered to the model at a dose of 5 mg/kg, wherein the black triangle shows the blood concentration of the Hspd1.8Lspd1.6 (Hd1.8Ld1.6) antibody, the white circle shows the blood concentration of the Hspu2.2Lspu2.2 (Hu2.2Lu2.2) antibody, and the black square shows the blood concentration of the H0L0 antibody;

Figure 8B shows the blood antibody concentration in a human liver cancer-transplant mouse model, of the H0L0 antibody, Hspu2.2Lspu2.2 (Hu2.2Lu2.2) antibody, and Hspd1.8Lspd1.6 (Hd1.8Ld1.6) antibody when each test antibody was administered to the model at a dose of 1 mg/kg, wherein the black triangle shows the blood concentration of the Hspd1.8Lspd1.6 (Hd1.8Ld1.6) antibody, the white circle shows the blood concentration of the Hspu2.2Lspu2.2 (Hu2.2Lu2.2) antibody, and the black square shows the blood concentration of the H0L0 antibody; and

Figure 9 shows the ADCC activity of test antibodies against HepG2 cells, a human liver cancer cell line, wherein the black triangle shows the ADCC activity by the Hspd1.8Lspd1.6 (Hd1.8Ld1.6) antibody, the white circle shows the ADCC activity by the Hspu2.2Lspu2.2 (Hu2.2Lu2.2) antibody, and the black square shows the ADCC activity by the H0L0 antibody.

Figure 10 shows the binding affinity for the antigen glypican 3 measured by a competitive ELISA of the H0L0 antibody, Hd1.8Ld1.6 antibody, pH7pL14 antibody and pH7pL16
antibody, wherein the black circle shows the binding activity of the H0L0 antibody, the white circle shows the binding activity of the Hdl.8Ld1.6 antibody, the black square shows the binding activity of the pH7pL14 antibody, and the white square shows the binding activity of the pH7pL16 antibody.

Figure 11 shows the antitumor activity of the H0L0 antibody, pH7pL14 antibody and pH7pL16 antibody in a mouse model implanted with human hepatic cancer, wherein * shows the antitumor activity of the H0L0 antibody, the white circle shows the antitumor activity of the Hdl.8Ld1.6 antibody, the black square shows the antitumor activity of the pH7pL14 antibody, and the white square shows the antitumor activity of the pH7pL16 antibody.

Figure 12 shows the blood concentration of the antibody in mice of the H0L0 antibody, Hdl.8Ld1.6 antibody, pH7pL14 antibody, pH7pL16 antibody and pH7M85pL16, wherein * shows the blood concentration of the H0L0 antibody, the white circle shows the blood concentration of the Hdl.8Ld1.6 antibody, the black square shows the blood concentration of the pH7pL14 antibody, the white square shows the blood concentration of the pH7pL16 antibody, the black triangle shows the blood concentration of the pH7M85pL16 in mouse.

Figure 13 shows the ADCC activity by H0L0 antibody, Hdl.8Ld1.6 antibody, pH7pL14 antibody, and pH7pL16 antibody against HepG2 cells, a human liver cancer cell line, wherein the black circle shows the ADCC activity by the H0L0 antibody,
the white circle the ADCC activity by the Hd1.8Ld1.6 antibody, the black square the ADCC activity by the pH7pL14 antibody, and the white square the ADCC activity by the pH7pL16 antibody.

Figure 14 shows the binding affinity for the antigen glypican 3 measured by a competitive ELISA of the HOLO antibody, HOM85L0 antibody, pH7pL16 antibody and pH7M85pL16 antibody, wherein the black triangle shows the binding activity of the HOLO antibody, the black square shows the binding activity of the HOM85L0 antibody, the asterisk shows the binding activity of the pH7pL16 antibody, and the white diamond shows the binding activity of the pH7M85pL16 antibody.

Figure 15 shows the ADCC activity by the pH7pL16 antibody and pH7M85pL16 antibody against HepG2 cells, a human liver cancer cell line, wherein the white square shows the ADCC activity by the pH7pL16 antibody, and the black triangle shows the ADCC activity by the pH7M85pL16 antibody.

DESCRIPTION OF THE PREFERRED EMBODIMENTS

The present invention provides a method of modulating the plasma kinetics of an antibody (e.g. anti-glypican 3 antibody). In a preferred embodiment of the present invention, the method comprises the modification in the charge of at least one amino acid residue that can be exposed on the surface of the antibody (e.g. anti-glypican 3 antibody). That is, the plasma kinetics of an antibody (e.g. anti-glypican 3 antibody) can be modulated by modifying the charge of an amino acid residue in
the antibody to cause a change in the isoelectric point (pI) thereof. The antibody (e.g. anti-glypican 3 antibody) having modulated plasma kinetics is able to exhibit an antitumor activity on cancer cells that is superior to that of the unmodulated antibody.

Among the several antibody isotypes, the principle metabolic pathways of the IgG antibody do not proceed via renal excretion due to the sufficiently high molecular weight of the IgG antibody. The IgG antibody, which contains the Fc region as a part of its molecule, is known to have a long in vivo half-life due to recycling by a salvage pathway mediated by the neonatal Fc receptor (FcRn), which is expressed by endothelial cells in, for example, the vascular system. It is thought that the IgG antibody is metabolized mainly by metabolic pathways in endothelial cells (He X. Y., Xu Z., Melrose J., Mullowney A., Vasquez M., Queen C., Vexler V., Klingbeil C., Co M. S., and Berg E. L., Humanization and pharmacokinetics of a monoclonal antibody with specificity for both E- and P-selectin, J. Immunol. (1998), 160(2), 1029-35). That is, it is thought that the IgG antibody is recycled through binding to the FcRn of IgG antibody nonspecifically taken up by the endothelial cell, while the IgG antibody that cannot be bound is metabolized. IgG antibody having the Fc region modified to lower the binding activity to FcRn exhibits a shorter half-life in the blood. In contrast, the plasma half-life of the IgG antibody can be increased by modifying
amino acid residues constituting the Fc region of the IgG antibody so as to increase the binding activity to FcRn (He X. Y., Xu Z., Melrose J., Mullowney A., Vasquez M., Queen C., Vexler V., Klingbeil C., Co M. S., and Berg E. L., Humanization and pharmacokinetics of a monoclonal antibody with specificity for both E- and P-selectin, *J. Immunol.* (1998), 160(2), 1029-35; and LinksOber RJ, Radu CG, Ghetie V, Ward ES. Differences in promiscuity for antibody-FcRn interactions across species: implications for therapeutic antibodies. *Int Immunol.* (2001) 13(12), 1551-9). As described above, the known methods for modulating the plasma kinetics of the IgG antibody involved modification of the binding activity to FcRn by modification of amino acid residues constituting the Fc region. Specific examples of the above amino acid residues include the amino acid residues H250, H253, H310, H311, H314, H428, H435 and H436, according to the Kabat numbering. In addition, the amino acid residues H254, H255, H257, H288, H296, H307, H309, H315, H415, H433, which indirectly involve the interaction between IgG antibodies and FcRn, were thought to be the target for modification. These amino acid residues correspond to, for example, the 130, 133, 190, 191, 194, 308, 315 and 316th amino acid residues, and the 134, 135, 137, 168, 176, 187, 189, 195, 295 and 313rd amino acid residues in SEQ ID NO:30, as well as the 133, 136, 193, 194, 197, 311, 318 and 319th amino acid residues and the 137, 138, 140, 171, 179, 190, 192, 198, 298 and 316th amino acid
residues in SEQ ID NO:31, respectively. However, as shown in the examples provided below, it has now found by the present invention that the plasma half-life of an antibody (e.g. anti-glypican 3 antibody) depends on the pI with a high correlation. Thus, it is now shown that the plasma half-life of the antibody (e.g. anti-glypican 3 antibody) can be modulated without modifying the amino acid residues constituting FcRn binding region, whose modification would invoke the immunogenicity, in particular the amino acid residues H250, H253, H310, H311, H314, H428, H435 and H436, as well as H254, H255, H257, H288, H296, H307, H309, H315, H415 and H433 according to the Kabat numbering. It was also a surprising result that the modification in the amino acid residues other than H250, H253, H310, H311, H314, H428, H435 and H436, as well as H254, H255, H257, H288, H296, H307, H309, H315, H415 and H433 exhibited a decrease in the pI value and change in the binding activity to FcRn.

While not wishing to be bound to a particular theory, the present inventors hold the following view at the present time. The rate of nonspecific uptake of the IgG antibody by endothelial cells is thought to depend on physicochemical Coulombic interaction between the IgG antibody and the negatively charged cell surface. It is therefore thought that decrease (increase) in the Coulombic interaction by lowering (raising) the pI of the IgG antibody may cause decrease (increase) in nonspecific uptake by the endothelial cell,
which in turn causes decrease (increase) in metabolism at the endothelial cell resulted in the modulation of blood kinetics. As used herein, "decreasing the Coulombic interaction" means an increase in the Coulombic forth as expressed in a repulsive force. Since Coulombic interaction between the antibody and the cell surface negative charge of the endothelial cell is a physicochemical interaction, it is believed that this interaction does not primarily depend on the amino acid sequence per se that constitutes the antibody. Therefore, the method of modulating the plasma kinetics discovered in the present invention can be broadly applied to any antibodies or anti-glypican 3 antibodies but not limited to only a specific antibody or an anti-glypican 3 antibody.

When an IgG antibody is used as the antibody (e.g. anti-glypican 3 antibody) of the present invention, any subtype may be used as long as it is an IgG-type antibody molecule. A bispecific IgG antibody may also be used. When the antibody (e.g. anti-glypican 3 antibody) of the present invention is a bispecific antibody, the antibody can also specifically bind both the corresponding antigen (glypican 3 molecule in the case of anti-glypican 3 antibody) and an epitope other than that antigen. For example, in order to recruit NK cells, cytotoxic T-cells, LAK cells, and so forth, a surface antigen that specifically binds to these cells may be suitably used as another antigen. It has been shown that cytotoxicity by LAK cells is exhibited against bile duct cancer using a bispecific
antibody produced from the MUSE11 antibody recognizing MUC1 (an adenocarcinoma-related antigen), and OKT3 antibody recognizing LAK cell surface antigen (Katayose Y., Kudo T., Suzuki M., Shinoda M., Saijyo S., Sakurai N., Saeki H., Fukuhara K., Imai K., and Matsuno S., MUC1-specific targeting immunotherapy with bispecific antibodies: inhibition of xenografted human bile duct carcinoma growth, Cancer Res. (1996) 56(18), 4205-12). The antibody (e.g. anti-glypican 3 antibody) having improved plasma kinetics of the present invention can be suitably used in place of the MUSE11 antibody recognizing MUC1. In addition, antibody that recognizes different epitopes of the antigen to which the antibody binds (glypican 3 molecule in the case of anti-glypican 3 antibody) can also be suitably used as the bispecific antibody (e.g. anti-glypican 3 antibody) of the present invention. In the case of low molecular weight antibodies for which renal excretion is the main metabolic pathway, such as scFv and Fab, the plasma kinetics of such antibodies cannot be modulated by the pI as described above. However, the present invention can be applied to any antibody molecule type if it is an Fc-coupled protein for which renal excretion is not the main metabolic pathway. Examples of such molecules include scFv-Fc, dAbs-Fc and Fc fusion proteins. Since the main metabolic pathway of these molecules is not via metabolism by renal excretion, the plasma kinetics of these molecules can be modulated by changing the pI according to the method of the
present invention. Antibody-like molecules are included in the antibody molecules envisaged by the invention. Antibody-like molecules are molecules that function by binding to a target molecule (Binz H. K., Amstutz P., and Pluckthun A., Engineering novel binding proteins from nonimmunoglobulin domains, *Nat. Biotechnol.* (2005) 23(10), 1257-68); and examples include DARPin, Affibody, and Avimer.

The term "modulated plasma kinetics" as used herein means that the plasma kinetics are modified in a desired direction when the plasma kinetics of the antibody after modification of the amino acids constituting the antibody (e.g. anti-glypican 3 antibody) are compared with the plasma kinetics prior to modification. Thus, when it is desired to increase the plasma half-life of the antibody (e.g. anti-glypican 3 antibody), "modulation of the plasma kinetics" refers to an increase in the plasma half-life of the antibody. When it is desired to decrease the plasma half-life of the antibody (e.g. anti-glypican 3 antibody), "modulation of the plasma kinetics" refers to decrease in the plasma half-life of the antibody.

Whether the plasma kinetics of the antibody (e.g. anti-glypican 3 antibody) of the present invention have been modified in the desired direction, that is, whether the plasma kinetics have been modulated as desired, can be appropriately evaluated by pharmacokinetic tests using, for example, mouse, rat, rabbit, dog, monkey, and so forth. In addition, an "extension of the plasma half-life" or a "decrease of the
plasma half-life" as used herein may also be comprehended via parameters other than the plasma half-life parameter, such as the mean plasma residence time and the plasma clearance (Analysis by Pharmacokinetic Practice, (Nanzando)). For example, the "modulation of the plasma kinetics" according to the present invention can be suitably evaluated with these parameters by carrying out noncompartmental analysis according to the instructions accompanying the WinNonlin (Pharsight) in vivo pharmacokinetic analysis software.

The phrase "amino acid residue that can be exposed on the surface" as used herein generally denotes an amino acid residue that resides on the surface of the polypeptide constituting the antibody (e.g. anti-glypican 3 antibody). The phrase "amino acid residue that resides on the surface of the polypeptide" refers to an amino acid residue whose side chain can come into contact with solvent molecules (typically water molecules). All of its side chain need not come into contact with solvent molecules. If even a portion of the side chain of an amino acid residue comes into contact with solvent molecules, such an amino acid residue is considered to be an amino acid residue that resides on the surface. Those skilled in the art can construct a homology model of the polypeptide or antibody using commercially available homology modeling software. Based on this homology model, amino acid residues on the surface of the polypeptide constituting the antibody (e.g. anti-glypican 3 antibody) can be appropriately selected as an
"amino acid residue that resides on the surface of the polypeptide".

The "amino acid residue that can be exposed on the surface" is not particularly limited in the present invention, but is preferably an amino acid residue residing outside the FcRn binding region of the antibody (e.g. anti-glypican 3 antibody). The FcRn binding region is preferably the Fc region, but also include, for example, a region consisting of one or more amino acid residues H250, H253, H310, H311, H314, H428, H435 and H436 according to the Kabat numbering. In addition, the amino acid residues H254, H255, H257, H288, H296, H307, H309, H315, H415, H433, which indirectly involve the interaction between IgG antibodies and FcRn, were thought to be the target for modification. These amino acid residues correspond to, for example, the 130, 133, 190, 191, 194, 308, 315 and 316th amino acid residues, and the 134, 135, 137, 168, 176, 187, 189, 195, 295 and 313rd amino acid residues in SEQ ID NO:30, as well as the 133, 136, 193, 194, 197, 311, 318 and 319th amino acid residues and the 137, 138, 140, 171, 179, 190, 192, 198, 298 and 316th amino acid residues in SEQ ID NO:31, respectively.

The amino acid residue to be subjected to charge modification in the antibody (e.g. anti-glypican 3 antibody) according to the present invention is preferably an amino acid residue constituting the heavy chain (H chain) variable region or the light chain (L chain) variable region of the antibody.
Preferred specific examples of these variable regions are the complementarity-determining region (CDR) and framework region (FR).

Those skilled in the art can appropriately select a surface amino acid residue in the antibody’s variable region based on a homology model built by homology modeling. Thus, a surface amino acid residue in the antibody’s variable region can be suitably selected from H1, H3, H5, H8, H10, H12, H13, H15, H16, H19, H23, H25, H26, H39, H42, H43, H44, H46, H68, H71, H72, H73, H75, H76, H81, H82b, H83, H85, H86, H105, H108, H110, and H112, which are amino acid residues according to the Kabat numbering. For example, in the FR of the humanized anti-glypican 3 antibody heavy chain shown in SEQ ID NO: 1, the surface amino residues may include, but not limited to, the amino acid residues at positions 1, 3, 5, 8, 10, 12, 13, 15, 16, 19, 23, 25, 26, 39, 42, 43, 44, 46, 69, 72, 73, 74, 76, 77, 82, 85, 87, 89, 90, 107, 110, 112, and 114. A surface amino acid residue in the heavy chain CDR can also be selected using the same homology model. Thus, H97, an amino acid residue according to the Kabat numbering, is exposed at the surface for almost all antibodies. For example, the serine at position 101 in the heavy chain CDR of the humanized anti-glypican 3 antibody shown in SEQ ID NO: 1 corresponds to that amino acid residue. Suitable examples of other amino acid residues in the heavy chain CDR of the humanized anti-glypican 3 antibody shown in SEQ ID NO: 1 are the amino acid residues at positions
52, 54, 62, 63, 65, and 66.

With respect to the light chain FR, surface amino acid residues in the antibody's variable region can be suitably selected from L1, L3, L7, L8, L9, L11, L12, L16, L17, L18, L20, L22, L38, L39, L41, L42, L43, L45, L46, L49, L57, L60, L63, L65, L66, L68, L69, L70, L74, L76, L77, L79, L80, L81, L85, L100, L103, L105, L106, and L107 of amino acid residues according to the Kabat numbering. For example, surface amino acids may include, but not limited to, 1, 3, 7, 8, 9, 11, 12, 16, 17, 18, 20, 22, 43, 44, 45, 46, 48, 49, 50, 54, 62, 65, 68, 70, 71, 73, 74, 75, 79, 81, 82, 84, 85, 86, 90, 105, 108, 110, 111, and 112 of the humanized anti-glypican 3 antibody shown in SEQ ID NO: 7. Surface amino acid residues in the light chain CDR can be selected using the same homology model as the homology model with which the surface amino acid residues in the heavy chain CDR were determined. Suitable examples of amino acid residues in the CDR of the humanized anti-glypican 3 antibody light chain shown in SEQ ID NO: 7 are the amino acid residues at positions 24, 27, 33, 55, and 59.

The term "modification" in an amino acid residue in the method of the present invention specifically denotes, inter alia, substitution of an original amino acid residue with another amino acid residue, deletion of an original amino acid residue, and addition of a new amino acid residue, and preferably indicates the substitution of an original amino acid residue with another amino acid residue. Thus,
"modification in the charge of an amino acid residue" in the present invention is preferably an amino acid substitution.

In order to carry out "modification in the charge of an amino acid residue" on the anti-glypican 3 antibody of the present invention, for example, the charge is preferably modified for at least one amino acid residue selected from the amino acid residues at positions 19, 43, 52, 54, 62, 63, 65, 66, and 107 in the heavy chain variable region constituting the humanized anti-glypican 3 antibody shown in SEQ ID NO: 1. In addition, the charge is preferably modified, for example, for at least one amino acid residue selected from the amino acid residues at positions 17, 24, 27, 33, 55, 59, 79, 82, 105 and 112 in the light chain variable region constituting the humanized anti-glypican 3 antibody shown in SEQ ID NO: 7.

Among the amino acid residues listed above, amino acid residues other than those modified in its charge need not be modified as long as the desired modulating effect on the plasma kinetics is being obtained; however, such amino acid residues may conventionally modified so as to have no charge or to have the same type of charge as the modified amino residue(s).

Charge-bearing amino acids are known to be present. In general, lysine (K), arginine (R), and histidine (H) are known as positively charged amino acids. Aspartic acid (D) and glutamic acid (E) are known as negatively charged amino acids. The amino acids other than those are known as the uncharged
amino acids.

Preferably the aforementioned "modified amino acid residue" is conventionally selected from, but not limited to, the amino acid residues present in either of the following groups (a) and (b).

(a) glutamic acid (E), aspartic acid (D)

(b) lysine (K), arginine (R), and histidine (H)

When the original (pre-modification) amino acid residue already bears a charge, modification so as to provide an uncharged amino acid residue is also a preferred embodiment of the present invention. Thus, modification in the present invention encompasses (1) substitution of a charged amino acid with an uncharged amino acid, (2) substitution of a charged amino acid with an amino acid bearing an opposite charge, and (3) substitution of an uncharged amino acid with a charged amino acid.

Modification of an amino acid residue constituting the antibody (e.g. anti-glypican 3 antibody) so as to change the isoelectric point (pI) of the antibody is preferred for the present invention. In addition, in those instances where a plurality of amino acid residues will be modified, the amino acid residues subjected to modification may include a small number of uncharged amino acid residues.

Suitable examples of the "modification in the charge of an amino acid residue" in the anti-glypican 3 antibody of the present invention are as follows. With regard to a
modification that increases the pI value, for example, at least one substitution selected from Q43N, D52N, and Q107R in the heavy chain variable region constituting the humanized anti-glypican 3 antibody shown in SEQ ID NO: 1 can be made, and a modification to the amino acid sequence shown in SEQ ID NO: 4 or 6 is particularly preferred. In addition, for example, at least one substitution selected from E17Q, Q27R, and Q105R in the light chain variable region constituting the humanized anti-glypican 3 antibody shown in SEQ ID NO: 7 can be made, and a modification to the amino acid sequence shown in SEQ ID NO: 10 or 12 is particularly preferred. With regard, on the other hand, to a modification that decreases the pI value, at least one substitution selected from K19T, Q43E, G62E, K63S, K65Q, and G66D in the heavy chain variable region constituting the humanized anti-glypican 3 antibody shown in SEQ ID NO: 1 can be made, and modification to the amino acid sequence shown in SEQ ID NO: 3, 5 or 27 is particularly preferred. In addition, for example, at least one substitution selected from R24Q, Q27E, K79T, R82S and K112E in the light chain variable region constituting the humanized anti-glypican 3 antibody shown in SEQ ID NO: 7 can be made, and modification to the amino acid sequence shown in SEQ ID NO: 9, 11, 28 or 29 is particularly preferred. In addition, the modification that decreases the pI value also include substitution of one or more amino acid residues in the heavy chain constant region designated by H268, H274, H355, H356, H358 and H419 according
to the Kabat numbering. Preferred examples of substitution is at least one modification in the heavy chain constant region shown in SEQ ID NO: 31 includes, for example, substitution of H that is the 151st amino acid residue with Q, substitution of K that is the 157th amino acid residue with Q, substitution of R that is the 238th amino acid residue with Q, substitution of D that is the 239th amino acid residue with E, substitution of L that is the 241st amino acid residue with M, and substitution of Q that is the 302nd amino acid residue with E. The above substitution resulted in a chimera of the constant region of IgG1 and the constant region of IgG4 of a human antibody. Thus such a substitution allows for preparation of a modified antibody with a desired pI value without affecting the immunogenicity of the antibody.

There are no particular limitations in the present invention on the number of amino acid residues submitted to modification; when, for example, the variable region of the antibody is being modified, preferably the fewest number of amino acid residues necessary to achieve the desired modulated plasma kinetics is modified, in order to avoid lowering the binding activity with antigen and in order to avoid raising the immunogenicity. It may also be suitable to implement a suitable combination with an amino acid residue modification that causes a decline in immunogenicity and/or an amino acid residue modification that causes an increase in the binding activity with antigen.
Known techniques can be used to measure the antigen binding activity of the antibody. For example, enzyme-linked immunosorbent assay (ELISA), enzyme immunoassay (EIA), radioimmunoassay (RIA), or a fluorescent immunoassay can be used. These methods are described in a common textbook, *Antibodies: A Laboratory Manual*, Ed Harlow and David Lane, Cold Spring Harbor laboratory, 1988.

The methods described on pages 359 to 420 of *Antibodies: A Laboratory Manual* (Ed Harlow and David Lane, Cold Spring Harbor laboratory, 1988) are examples of methods that can be used to measure an antibody's binding activity for cells. The binding activity may be evaluated based on a FACS (fluorescence activated cell sorting) or ELISA principle using the cells as an antigen. In the ELISA format, an antibody's binding activity for cells is quantitatively evaluated by comparing the signal levels generated by an enzymatic reaction. Thus, a test antibody is added to an ELISA plate on which the over-expressing cells have been immobilized and the antibody bound to the cells is detected by an enzyme-labeled antibody that recognizes the test antibody. In the case of FACS, the binding activity for cells can be compared by constructing a dilution series with the test antibody and comparing the antibody binding titers for the over-expressing cells.

Binding between an antigen expressed on the surface of cells suspended in a buffer and not anchored on a carrier (such as an ELISA plate) and an antibody for this antigen can
be measured by the FACS format. Flow cytometers used in such measurements may include FACSCanto™ II, FACSAria™, FACSArray™, FACSVantage™ SE, and FACSCalibur™ (all from BD Bioscience) and the EPICS ALTRA HyperSort, Cytomics FC 500, EPICS XL-MCL ADC, EPICS XL ADC, and Cell Lab Quanta/Cell Lab Quanta SC (all from Beckman Coulter).

In an example of a suitable method for measuring the binding activity of a test anti-glypican 3 antibody to an antigen, the test antibody is reacted with a cell expressing glypican 3; the cells are stained with an FITC-labeled secondary antibody that recognizes the test antibody; the fluorescent intensity is measured with FACSCalibur (BD); and analyzed using CELL QUEST software (BD). According to this method, the test antibody bound to the glypican 3 on the surface of glypican 3-expressing cells is stained by FITC-labeled secondary antibody that specifically recognizes the test antibody, and the fluorescent intensity if measured by FACSCalibur, then the geometric mean value (test geo-mean value) obtained by analysis of the resulting fluorescent intensity is compared with the control geo-mean value obtained from a control antibody using the CELL QUEST software. The computational formulas that yield the geo-mean value (geometric mean) are described in the CELL QUEST Software User’s Guide (BD Biosciences).

In order to avoid increasing the in vivo immunogenicity for the human receiving the antibody, the modified amino acid
sequence is preferably, but not limited to, a human sequence (sequence seen in a naturally occurring antibody of human origin). In addition, mutations can be suitably introduced at locations other than the modifications introduced to change the isoelectric point, so as to modify each of the plurality of FRs (FR1, FR2, FR3, FR4) into a human sequence. A method that converts each of the FRs into a human sequence in this manner is reported by Ondo K., Ohtomo T., Yoshida K., Yoshimura Y., Kawai S., Koishihara Y., Ozaki S., Kosaka M., and Tsuchiya M., The humanized anti-HM1.24 antibody effectively kills multiple myeloma cells by human effector cell-mediated cytotoxicity, Mol. Immunol. (1999) 36(6), 387-395. In addition, in order to change the isoelectric point of the antibody, each of the FR sequence may be converted to another human FR sequence in order to change the charge of a particular FR (for example, FR3 may be exchanged with another human FR in order to lower the isoelectric point of the antibody). Such a humanization method is reported in Dall'Acqua W. F., Damschroder M. M., Zhang J., Woods R. M., Widjaja L., Yu J., and Wu H., Antibody humanization by framework shuffling, Methods (2005) 36(1), 43-60.

In those instances where the desired modulated plasma kinetics are not achieved by a modest modification of the surface charge, an antibody (e.g. anti-glypican 3 antibody) exhibiting the desired modulated plasma kinetics can be suitably obtained by the repetitive execution of surface
charge modification and evaluation of plasma kinetics.

The plasma kinetics of chimeric EP5C7.g4, a chimeric anti-E, P-selectin antibody (IgG4), have been compared with those of HuEP5C7.g4, a humanized antibody (IgG4), and the two were shown to have the same plasma kinetics in the rhesus monkey (He X. Y., Xu Z., Melrose J., Mullowney A., Vasquez M., Queen C., Vexler V., Klingbeil C., Co M. S., and Berg E. L., Humanization and pharmacokinetics of a monoclonal antibody with specificity for both E- and P-selectin, J. Immunol. (1998), 160(2), 1029-35). In addition, the plasma kinetics of ch5d8, a chimeric anti-CD154 antibody, have been compared with those of the humanized antibody Hu5c8 in cynomolgus monkey and the two were shown to have the same plasma kinetics (Gobburu J. V., Tenhoor C., Rogge M. C., Frazier D. E. Jr., Thomas D., Benjamin C., Hess D. M., and Jusko W. J., Pharmacokinetics/dynamics of 3c3, a monoclonal antibody to CD154 (CD40 ligand) suppression of an immune response in monkeys, J. Pharmacol. Exp. Ther. (1998) 286(2), 925-30). The plasma kinetics of the chimeric antibody cCC49 were shown to be the same as those of the humanized antibody HuCC49 in mice (Kashmiri S. V., Shu I., Padlan E. A., Milenic D. E., Schlom J., and Hand P. H., Generation, characterization, and in vivo studies of humanized anticanicoma antibody CC49, Hybridoma (1995) 14(5), 461-73). The plasma kinetics and distribution of mouse antibody and humanized antibody were shown to be the same in evaluations in mice (Graves S. S., Goshorn S. C.,
Stone D. M., Axworthy D. B., Reno J. M., Bottino B., Searle S., Henry A., Pedersen J., Rees A. R., and Libby R. T., Molecular modeling and preclinical evaluation of the humanized NR-LU-13 antibody, *Clin. Cancer Res.* (1999) 5(4), 899-908; Couto J. R., Blank E. W., Peterson J. A., and Ceriani R. L., Anti-BA46 monoclonal antibody Mc3: humanization using a novel positional consensus and in vivo and in vitro characterization, *Cancer Res.* (1995) 55(8), 1717-22. It is thought that the plasma kinetics and distribution of the chimeric antibodies and humanized antibodies are the same due to the fact that the murine Fc and human Fc are both cross-reactive with the murine FcRn. As shown by these examples, the plasma kinetics are the same between chimeric antibodies and humanized antibodies that have the same CDR. This means that humanization by the known methods provided in Ghetie V., Popov S., Borvak J., Radu C., Matesoi D., Medesan C., Ober R. J., and Ward E. S., Increasing the serum persistence of an IgG fragment by random mutagenesis, *Nat. Biotechnol.* (1997) 15(7), 637-40 and so forth, may provide the same plasma kinetics as the chimeric antibody, and thus a humanized antibody having modulated plasma kinetics therefore cannot be produced by the known methods.

In contrast, when a chimeric antibody (e.g. chimeric anti-glypican 3 antibody) is humanized according to the method of the present invention, the pl of the antibody is changed by modification to an amino acid residue that can be exposed on the surface of the chimeric antibody to construct a humanized
antibody (e.g. humanized anti-glypican 3 antibody) that exhibits modulated plasma kinetics (i.e., an increase or decrease in its plasma half-life) in comparison to the original chimeric antibody. For the purpose of modulating the plasma kinetics, the modification of amino acid that can be exposed on the surface of the humanized antibody (e.g. humanized anti-glypican 3 antibody) may be carried out at the same time as the humanization of the antibody, or the pI of the humanized antibody may be modified by modification of surface-exposed amino acid starting from the humanized antibody (e.g. humanized anti-glypican 3 antibody).

It has been established that trastuzumab, bevacizumab, and pertuzumab, which are three humanized antibodies that have been humanized using the same human antibody FR sequence, have about the same plasma kinetics (Adams C. W., Allison D. E., Flagella K., Presta L., Clarke J., Dybdal N., McKeever K., and Sliwkowski M. X., Humanization of a recombinant monoclonal antibody to produce a therapeutic HER dimerization inhibitor, pertuzumab, Cancer Immunol Immunother. (2006) 55(6), 717-27). Thus, the plasma kinetics are about the same when antibodies are humanization using the same FR sequence. According to the method of the present invention, the plasma kinetics of the antibody (e.g. anti-glypican 3 antibody) can be modulated in the humanization step where the pI of the antibody (e.g. anti-glypican 3 antibody) is modified by adding a modification to an amino acid residue that can be exposed on the surface of
the antibody.

The method of the present invention can also be applied to human antibodies. A human antibody (e.g. human anti-glypican 3 antibody) having modulated plasma kinetics relative to the plasma kinetics of the initially prepared human antibody (i.e., an increase or decrease in the former’s plasma half-life) can be constructed by modification of the pI of a human antibody (e.g. human anti-glypican 3 antibody) by adding a modification to an amino acid residue that can be exposed on the surface of a human antibody constructed from a human antibody library, a human antibody-producing mouse, and so forth.

The plasma half-life of an antibody is increased by lowering the pI value of the antibody. In contrast, it is known that the plasma half-life is decreased and the antibody’s tissue translocation characteristics are improved by raising the antibody’s pI value (Vaisitti T., Deaglio S., and Malavasi F., Cationization of monoclonal antibodies: another step towards the "magic bullet"?, J. Biol. Regul Homoest. Agents (2005) 19(3-4), 105-12; Pardridge W. M., Buciak J., Yang J., and Wu D., Enhanced endocytosis in cultured human breast carcinoma cells and in vivo biodistribution in rats of a humanized monoclonal antibody after cationization of the protein, J Pharmacol Exp Ther. (1998) 286(1), 548-54). However, due to fact that such an antibody exhibits an increased immunogeneity and an enhanced
internalization into the cell, additional improvements are required for application as an antibody that exhibits anti-cancer effect via a mechanism such as a cytotoxic activity, because internalization into the cell is a hindrance to the manifestation of its cytotoxic activity, such as ADCC activity, CDC activity, and so forth. Thus, with regard to antibody that exhibits anti-cancer effect via a mechanism such as a cytotoxic activity, where internalization into the cell is a hindrance to the manifestation of its cytotoxic activity, such as ADCC activity, CDC activity, and so forth, it had not been determined whether an increase in the pI value of an antibody or a reduction in the pI value of an antibody causes enhancement of the tumor-inhibiting effect. In the present invention, modified humanized anti-glypican 3 antibody with a reduced pI value and modified humanized anti-glypican 3 antibody with an increased pI value were constructed, and then the question of which modification has the higher tumor-inhibiting activity was examined by subjecting both antibodies to comparative testing of the antitumor effect. As a result, it was surprisingly shown that the humanized anti-glypican 3 antibody with the reduced pI value exhibited the better effect against liver cancer.

The term "anti-glypican 3 antibody" as used herein encompasses anti-glypican 3 antibody obtained by subjecting anti-glypican 3 antibody that has already been subjected to amino acid residue charge modification as described above, to
additional modification of its amino acid sequence, for example, by subjecting the amino acid residues constituting this anti-glypican 3 antibody to substitution, deletion, addition, and/or insertion. In addition, the term "anti-glypican 3 antibody" as used herein also encompasses anti-glypican 3 antibody obtained starting from anti-glypican 3 antibody that has already been subjected to amino acid residue substitution, deletion, addition, and/or insertion, or to modification of its amino acid sequence by, for example, chimerization, humanization, and so forth, and additionally subjecting the amino acid residues constituting this anti-glypican 3 antibody to charge modification.

A preferred example of a modification whose goal is to improve the characteristics of the antibody (e.g. anti-glypican 3 antibody) of the present invention is a modification whose goal is to raise the stability of the antibody (referred to below as a stability modification). In aqueous solution, the antibody equilibrates between two states, its native state and an inactive denatured state. The stability of the native state, as shown by the second law of thermodynamics (ΔG = ΔH - TΔS), depends on the Gibbs free energy change ΔG of the system and the balance between the components of ΔG, i.e., the enthalpy change ΔH (attributable to changes in, for example, hydrophobic interactions and hydrogen bond in the polypeptide chain) and the entropy change
\( \Delta S \) (attributable to changes in solvation and the degrees of freedom in the three-dimensional structure). A positive value for \( \Delta G \) indicates that the native state of the protein is more stable than the protein's denatured state, and the stability of the native state of the protein rises as \( \Delta G \) assumes larger positive values. The forces contributing to this stability must be disrupted in order to denature the protein. For example, exposing the protein solution to high temperatures results in an increase in the degrees of freedom in the three-dimensional structure and a weakening of the factors that contribute to protein stabilization, causing a thermal denaturation of the protein. In this case the \( -T\Delta S \) term governs the denaturation. The \( \Delta H \) of the unfolding due to thermal denaturation of the protein can be directly measured by differential scanning calorimetry (DSC), as is specifically described in the examples provided herein. The DSC curve for the protein thermal denaturation process takes the form of an endothermic peak that frames a temperature, known as the denaturation midpoint (Tm), that is characteristic of the tost protein. The denaturation enthalpy change is obtained by integration of this peak. The Tm value is generally an indicator of thermal stability. The change in the heat capacity (\( \Delta C_p \)) can also be measured during thermal denaturation of the protein by DSC. The change in the heat capacity that occurs during denaturation is caused mainly by
hydration that occurs when amino acid residues that are not exposed on the molecule's surface when the protein is in its native state become exposed to solvent molecules during the course of protein denaturation.

As described above, amino acid residue "modification" in the method of the present invention specifically encompasses, inter alia, substitution of an original amino acid residue with another amino acid residue, deletion of an original amino acid residue, and the addition of a new amino acid residue. The substitution of an original amino acid residue with another amino acid residue is preferred. Thus, modification by amino acid substitution is preferably used in the present invention when a modification of antibody stability is sought. The Tm value of the antibody (e.g. anti-glypican 3 antibody) is increased as a result of the execution of the stability modification on the amino acid residues constituting the antibody. Thus, the Tm value is suitably used as an indicator of the stability modification of the antibody (e.g. anti-glypican 3 antibody) has occurred.

In order to carry out the aforementioned "stability modification" with the anti-glypican 3 antibody of the present invention, modification is preferably carried out, for example, on at least one amino acid residue selected from the amino acid residues at positions 37, 40, 48, and 51 in the humanized anti-glypican 3 antibody heavy chain variable region shown in SEQ ID NO: 1. In addition, modification is preferably carried
out on at least one amino acid residue selected from the amino acid residues at positions 2, 25, 42, 48, 50, 83, and 84 in the humanized anti-glypican 3 antibody light chain variable region shown in SEQ ID NO: 7. The amino acid residues other than the aforementioned amino acid residues on which stability modification has been carried out, need not be modified as long as the desired Tm value has been obtained; however, it can be appropriate to carry out a suitable modification thereon so as to provide a Tm value that is about the same as or higher than the Tm value of the humanized anti-glypican 3 antibody submitted to the modification.

The stability modification can be carried out by randomly modifying amino acid residues constituting the antibody (e.g. humanized anti-glypican 3 antibody) submitted to the modification. In addition, stability modification can also be carried out by replacing a portion of the amino acid sequence constituting the humanized antibody (e.g. antibody anti-glypican 3 antibody) submitted to the modification, with an amino acid sequence found in an antibody that already has a high Tm value and that corresponds - from the standpoint of the correlation of antibody three-dimensional structure - with said portion of the amino acid sequence constituting the humanized antibody (e.g. humanized anti-glypican 3 antibody) submitted to the modification. There are no limitations on the position of the amino acid residue or residues undergoing substitution; however, amino acid residue or residues in the
FR are preferably modified. Amino acid residue modification can even be carried out as appropriate in the CDR region as long as there is no associated reduction in the binding activity for the antigen. In addition, the number of amino acid residues subjected to modification is not particularly limited, and modification can even be implemented by replacing a particular segment of the FR with a desired segment. With regard to such segments, all of the segments within the FR (FR1, FR2, FR3, FR4) can be modified, or one or more of the segments modification may be combined.

The FR2 of the heavy chain or the light chain is a preferred example in those instances where a segment of the FR is modified. A preferred specific example in this regard is an amino acid residue modification in which humanized anti-glypican 3 antibody heavy chain FR2 in the VH1b subclass (shown in SEQ ID NO: 1) is modified to the VH4 subclass, i.e., V37I (valine at position 37 is replaced by isoleucine) as well as A40P, M48I, and L51T modifications. Other preferred specific examples are a modification of humanized anti-glypican 3 antibody light chain FR2 region in the VK2 subclass (shown in SEQ ID NO: 7) to the VK3 subclass, i.e., L42Q, S48A, and Q50R modifications. Also preferred is V2I modification, which corresponds to modification to a germline sequence for FR1.

The execution of substitution, deletion, addition, and/or insertion with respect to the amino acid residues constituting
the antibody (e.g. anti-glypican 3 antibody) and the modification of the amino acid sequence by, for example, chimerization and humanization, can be carried out as appropriate by any method known to those skilled in the art. The execution of substitution, deletion, addition, and/or insertion with respect to the amino acid residues constituting the antibody’s variable region and constant region can similarly be carried out as appropriate during construction of the antibody (e.g. anti-glypican 3 antibody) of the present invention as a recombinant antibody.

Any antibodies of animal origin, e.g., mouse antibody, human antibody, rat antibody, rabbit antibody, goat antibody, camel antibody may preferably be used as the antibody (e.g. anti-glypican 3 antibody) of the present invention. Also preferred for use is a modified antibody (e.g. anti-glypican 3 antibody) as obtained by substitution in the amino acid sequence of chimeric antibody and humanized antibody. Also preferred for use are antibody modifications in which any of various molecules is attached.

"Chimeric antibody" refers to an antibody constructed by combining sequences originating from different animals. Suitable examples in this regard are antibodies constructed from the heavy chain and light chain variable regions from mouse antibody and the heavy chain and light chain constant regions from human antibody. Methods for constructing chimeric antibodies are known. For example, recombinant DNA is
generated by the in-frame fusion of DNA encoding a mouse antibody variable region and DNA encoding a human antibody constant region, and can be incorporated into a conventional expression vector. A host cell transfected or transformed with this vector is cultured and the chimeric antibody can be prepared or isolated from a culture medium by appropriate means.

Humanized antibodies are also known as reshaped human antibodies. These are antibodies in which a complementarity-determining region (CDR) from an antibody isolated from a nonhuman mammal, such as mouse, is ligated to a framework region (FR) from a human antibody. The DNA sequence encoding the humanized antibody can be synthesized by an overlapping PCR reaction using a plurality of oligonucleotides as primers. The starting materials and procedures for overlapping PCR reactions are described, for example, in WO 98/13388 and elsewhere. DNA encoding the variable region of a humanized antibody of the present invention is obtained by overlapping PCR from a plurality of oligonucleotides constructed so as to have nucleotide sequences that overlap with each other, which is in turn ligated with DNA encoding a human antibody constant region so as to form an in-frame codon sequence. The DNA ligated as described is then operably inserted into an expression vector followed by transfection into a host.

Methods for identifying CDRs are known (Kabat et al., Sequences of Proteins of Immunological Interest (1987),

60
National Institute of Health, Bethesda, Md.; Chothia et al.,
procedures therefor are also known (refer to EP 125023 A and
WO 96/02576). Once a CDR from an antibody of a nonhuman animal,
for example, a murine antibody, has been determined, these
known methods can be used to construct DNA encoding a
recombinant antibody in which the CDR and FR from a human
antibody are ligated. The human antibody FRs ligated with the
CDR are selected in such a manner that the CDR forms a high-
quality antigen binding site. As necessary, an amino acid
residue or residues in the antibody variable regions FRs may
be modified as appropriate so as to enable the CDR of the
reshaped human antibody to form a suitable antigen binding
site (Sato et al., _Cancer Res._ (1993) 53, 851-6). The amino
acid residues in the FRs submitted to modification may include
residues that directly bind to the antigen by noncovalent
bonds (Amit et al., _Science_ (1986) 233, 747-53), residues that
influence or act on the CDR structure (Chothia et al., _J. Mol.
Biol._ (1987) 196, 901-17), and residues related to VH-VL
(heavy chain variable region-light chain variable region)
interactions (EP 239400 B).

The humanized antibody encoded by the DNA is produced by
the host cell that has been transformed or transfected by the
generally used expression vector into which the DNA has been
inserted and is isolated from the culture medium yielded by
the culture of the host cell.
When the antibody of the present invention is a chimeric antibody, humanized antibody or human antibody, a constant region of human antibody origin is preferably used as the constant region of the antibody. For example, when the anti-glypican 3 antibody of the present invention is a chimeric anti-glypican 3 antibody, humanized anti-glypican 3 antibody or human anti-glypican 3 antibody, a constant region of human antibody origin is preferably used as the constant region of the anti-glypican 3 antibody. For example, \( \gamma_1 \), \( \gamma_2 \), \( \gamma_3 \), and \( \gamma_4 \) are each suitable for use as the heavy chain constant region, while \( \kappa \) and \( \lambda \) are each suitable for use as the light chain constant region. In addition, the human antibody constant region can be modified as appropriate in order to improve the antibody (e.g. anti-glypican 3 antibody) or improve the stability of its production. Chimeric antibody (e.g. chimeric anti-glypican 3 antibody) of the present invention suitably comprises a variable region from an antibody of a nonhuman mammal and a constant region from a human antibody. On the other hand, the humanized antibody preferably comprises a CDR from an antibody of a nonhuman mammal and a constant region and FR from a human antibody. For example, a humanized anti-glypican 3 antibody preferably comprises a CDR from an anti-glypican 3 antibody of a nonhuman mammal and a constant region and FR from a human antibody. The human antibody suitably comprises a CDR from an antibody of
human origin and a constant region and FR from a human antibody. For example, the human anti-glypican 3 antibody suitably comprises a CDR from an anti-glypican 3 antibody of human origin and a constant region and FR from a human antibody. The human antibody constant region is composed of an amino acid sequence that is characteristic of the particular isotype, i.e., IgG (IgG1, IgG2, IgG3, IgG4), IgM, IgA, IgD, and IgE. The constant region from antibody belonging to any of these isotypes is suitably used as the constant region of the humanized antibody (e.g. anti-glypican 3 antibody) of the present invention. The use of the constant region from human IgG is preferred, but not limited. There are also no particular limitations on the human antibody FR used as the FR of the humanized antibody (e.g. humanized anti-glypican 3 antibody) or human antibody (e.g. human anti-glypican 3 antibody), and the FR from antibody belonging to any isotype is suitably used.

With the aim of lowering the immunogenicity, all or a portion of the amino acid residues constituting the FR can also be replaced by germline sequences using the method described in Ono K. et al., loc. cit., or a similar method. Based on the rational prediction that germline sequences will have a low immunogenicity, the amino acid sequence constituting the FR of the humanized antibody may be compared, by alignment, with germline amino acid sequences (Abhinandan K. R. and Martin C. R., J. Mol. Biol. (2007) 369, 852-862). The
amino acid residues of the humanized antibody FR that differ in this comparison can be replaced, within a range that does not impair the antigen binding characteristics, with amino acid residues from a germline sequence. The following are specific examples for the amino acid residues constituting the heavy chain variable region shown in SEQ ID NO: 1:

modification that replaces the L at position 70 with I,
modification that replaces the T at position 87 with R, and
modification that replaces the T at position 97 with A. In addition, a modification that replaces the S at position 25 with A is an example for the amino acid residues constituting the light chain variable region shown in SEQ ID NO: 7.

With regard to the variable region and constant region of the modified chimeric antibody, humanized antibody, and human antibody of the present invention, deletion, substitution, insertion, and/or addition can be carried out as appropriate at one or more of the amino acids constituting the variable region and/or constant region of the antibody that was submitted to modification, insofar as binding specificity for the antigen is exhibited. In particular, the variable region and constant region of the modified chimeric anti-glypican 3 antibody, humanized anti-glypican 3 antibody, and anti-glypican 3 human antibody of the present invention, deletion, substitution, insertion, and/or addition can be carried out as appropriate at one or more of the amino acids constituting the variable region and/or constant region of the anti-glypican 3
antibody that was submitted to modification, insofar as binding specificity for the antigen glypican 3 molecule is exhibited.

Because chimeric anti-glypican 3 antibodies utilizing human-derived sequences, humanized anti-glypican 3 antibodies, and human anti-glypican 3 antibodies have a reduced immunogenicity in the human body, they are believed to be useful for application as therapeutic antibodies that are administered to humans, for example, with a therapeutic objective.

Known sequences can be used in the method of the present invention for the sequences that encode the antibody heavy chain and light chain for introduction of mutation. In addition, novel antibody gene sequences can be obtained by methods known to those skilled in the art. For example, genes can be suitably obtained from antibody libraries. Moreover, genes can also be obtained by cloning using known procedures, e.g., RT-PCR using as template mRNA from a monoclonal antibody-producing hybridoma.

Numerous antibody libraries are already known within the sphere of antibody libraries. Moreover, as methods for constructing antibody libraries are also known, those skilled in the art will be able to obtain or construct relevant antibody libraries. Examples of suitable antibody libraries are the antibody phage libraries disclosed in the literature, for example, Clackson et al., Nature (1991) 352, 624-8; Marks
et al., J. Mol. Biol. (1991) 222, 581-97; Waterhouses et al., Nucleic Acids Res. (1993) 21, 2265-6; Griffiths et al., EMBO J. (1994) 13, 3245-60; Vaughan et al., Nature Biotechnology (1996) 14, 309-14; and Japanese Patent Application Laid-open No. 2008-504970. A method of constructing a library in eukaryotic cells (WO 95/15393) and known methods such as ribosome display and so forth are also suitably used. Technology for obtaining human antibodies by panning techniques using a human antibody library as the starting material is also known to those skilled in the art. Thus, single-chain antibody (scFv), comprising the variable regions of human antibody heavy and light chains fused in-frame, is expressed on a phage surface using phage display techniques. Genes encoding antigen-binding scFv are isolated from this phage by selecting for phage that binds to the antigen. Identification of the sequence of such genes enables the determination of the DNA sequence that encodes the heavy and light chain variable regions of a human anti-glypican 3 antibody that binds to the antigen glypican 3. Human anti-glypican 3 antibody is suitably prepared by inserting the antibody gene having this sequence in a suitable expression vector and allowing for expression in a suitable host cell as described below. Such methods are well-known in the art and include those disclosed in WO 92/01047, WO 92/20791, WO 93/06213, WO 93/11236, WO 93/19172, WO 95/01438, and WO 95/15388.
Known technologies can be used to obtain an antibody-encoding gene from a hybridoma that produces the antibody, particularly to obtain an anti-glypican 3 antibody-encoding gene from a hybridoma that produces anti-glypican 3 monoclonal antibody. Briefly, an animal is immunized with glypican 3 (the desired sensitizing antigen) using a standard immunization technique and the immunocytes obtained from the animal are subjected to cell fusion with known parent cells by a standard cell fusion technique, which is described in detail below. Using standard screening techniques, the monoclonal antibody-producing cells (hybridomas) are screened, and cDNA for the variable region (V region) of anti-glypican 3 antibody can be synthesized with reverse transcriptase using the mRNA isolated from the selected hybridoma as a template. An anti-glypican 3 antibody gene is suitably prepared by the in-frame fusion of this cDNA with DNA encoding the desired antibody constant region (C region).

Specific examples are provided in the following, but the present invention is not limited to these examples. The sensitizing antigen used to generate the antibody of the present invention may be the complete immunogenic antigen or may be an incomplete antigen, including, for example, a nonimmunogenic hapten. For example, the full length glypican 3 protein or a partial polypeptide or peptide thereof can be suitably used. The soluble GPC3 core polypeptide shown in SEQ ID NO: 13 is a preferred example. Otherwise, the use of

67
substances comprising polysaccharide, nucleic acid, lipid, and so forth as an antigen is also known. Antigen to which the anti-glypican 3 antibody of the present invention binds is not particularly limited to the embodiments described above. Antigen production is suitably carried out by methods known to those skilled in the art. For example, a method using baculovirus (for example, WO 98/46777 and so forth) can be suitably used. When the antigen has a low immunogenicity, the animal can suitably be immunized with such an antigen attached to a very large immunogenic molecule, such as albumin. When the sensitizing antigen is a molecule that spans the cell membrane, such as glypican 3, a polypeptide fragment from the molecule’s extracellular domain is suitably used as necessary as the sensitizing antigen. Or, a cell expressing such a molecule on the cell surface is suitably used as the sensitizing antigen. In addition, when the sensitizing antigen is an insoluble molecule, solubilization may be effected by attaching the molecule with another water-soluble molecule and the solubilized molecule may then be suitably used as the sensitizing antigen.

Antibody-producing cells (e.g. anti-glypican 3 antibody-producing cells) are suitably obtained by immunizing an animal using a suitable sensitizing antigen as described above. Or, antibody-producing cells (e.g. anti-glypican 3 antibody-producing cells) can be obtained by in vitro immunization of lymphocytes that are capable of producing antibody. Various
vertebrates and mammals can be used as the animal to be immunized. In particular, rodents, lagomorphs, and primates can be generally used as the animal to be immunized. The rodents may include mouse, rat, and hamster; the lagomorphs may include rabbit; and the primates may include monkeys such as the cynomolgus monkey, rhesus monkey, hamadryas baboon, and chimpanzee. In addition, transgenic animals that maintain a repertoire of human antibody genes in their genome are also known, and human antibody can be suitably obtained using such animals (WO 96/34096; Mendez et al., Nat. Genet. (1997) 15, 146-56). Rather than using such transgenic animals, a desired human antibody that exhibits binding activity to a desired antigen may be suitably obtained by, for example, sensitizing human lymphocytes in vitro with the desired antigen or with cells that express the desired antigen followed by cell fusion with human myeloma cells, for example, U266 (Japanese Patent Publication No. Hei 1-59878). In addition, a desired human antibody (e.g. human anti-glypican 3 antibody) can be suitably obtained by the immunization with a desired of antigen of a transgenic animal that maintains the entire repertoire of human antibody genes in its genome (WO 93/12227, WO 92/03918, WO 94/02602, WO 96/34096, and WO 96/33735).

Immunization of the animal is carried out, for example, by suitable dilution and suspension of the sensitizing antigen in phosphate-buffered saline (PBS), physiological saline solution, and so forth; as necessary emulsification by the
admixture of adjuvant; and then intraabdominal or subcutaneous injection of the sensitizing antigen into the animal. This is followed preferably by several administrations of the sensitizing antigen mixed with Freund’s incomplete adjuvant on a 4 to 21 day interval. The production in the immunized animal of antibody against the sensitizing antigen can be measured using known analytical techniques, for example, enzyme-linked immunosorbent assay (ELISA) and flow cytometry (FACS).

A hybridoma can be prepared by fusing an anti-glypican 3 antibody-producing cell obtained from an animal or lymphocyte immunized with the sensitizing antigen of interest, with myeloma cells using a fusing agent conventionally used for cell fusion, e.g., polyethylene glycol (Goding, Monoclonal Antibodies: Principles and Practice, Academic Press (1986) 59-103). Hybridoma production can be suitably carried out according to, for example, the method of Milstein et al. (G. Kohler and C. Milstein, Methods Enzymol. (1981) 73, 3-46). Culture and proliferation of the hybridoma cells prepared by this method yields monoclonal antibody that is produced by the hybridoma and specifically binds to glypican 3. The binding specificity exhibited by this monoclonal antibody for glypican 3 can be suitably measured by known analytical techniques, such as immunoprecipitation, radioimmunoassay (RIA), enzyme-linked immunosorbent assay (ELISA), flow cytometry (FACS), and so forth. As necessary, a hybridoma that produces anti-glypican 3 antibody with the desired specificity, affinity, or
activity may then be suitably subcloned by, for example, limit dilution, and the monoclonal antibody produced by this hybridoma can be isolated.

The gene encoding the selected antibody can then be cloned from the aforementioned hybridoma or antibody-producing cell (e.g., sensitized lymphocyte) using a probe capable of specifically binding to the gene (for example, oligonucleotide complementary to the sequence encoding the antibody's constant region). Cloning can also be carried out by RT-PCR using mRNA isolated from the hybridoma or antibody-producing cell (e.g., sensitized lymphocyte) as a template. Immunoglobulins are divided into 5 different classes, i.e., IgA, IgD, IgE, IgG, and IgM, based on differences in their structure and function. Furthermore, the individual classes are divided into several subclasses (isotypes) (for example, IgG1, IgG2, IgG3, and IgG4; IgA1 and IgA2; and so forth). The antibody of the present invention may originate from antibody belonging to any of these classes and subclasses and is not particularly limited to any class or subclass; however, antibody belonging to the IgG class is particularly preferred.

Genes encoding the amino acid sequences constituting the heavy chain and light chain of the antibody (e.g. anti-glypican 3 antibody) can be modified as appropriate by the techniques of genetic engineering. For example, a recombinant antibody that has been subjected to artificial modification with the aim of, for example, reducing the xeno-
immunogenicity against humans (e.g., chimeric antibody such as chimeric anti-glypican 3 antibody, humanized antibody such as humanized anti-glypican 3 antibody, and so forth), can be suitably produced by modification of the nucleic acid residues that encode the amino acid sequence constituting the antibody, for example, a mouse antibody, rat antibody, rabbit antibody, hamster antibody, sheep antibody, camel antibody, and so forth. A chimeric antibody is an antibody constituted of the heavy chain and light chain variable regions of antibody originating from a nonhuman mammal such as mouse, and the heavy chain and light chain constant regions of human antibody. A chimeric antibody can be obtained as follows: DNA encoding the variable region of a mouse-originated antibody is ligated with DNA encoding the constant region of a human antibody and inserted into an expression vector, and the resulting recombinant vector is introduced into a host and allowed for expression. Humanized antibody is also known as reshaped human antibody and is an antibody in which the complementarity-determining region (CDR) of an antibody (e.g. anti-glypican 3 antibody) isolated from a nonhuman mammal such as mouse is ligated with a human antibody framework region so as to form an in-frame codon sequence. The DNA sequence encoding a humanized antibody can be synthesized by an overlapping PCR reaction using a plurality of oligonucleotides as templates. The starting materials and procedures for the overlapping PCR reaction are described in, for example, WO 98/13388 and elsewhere.
The DNA encoding the variable region of the recombinant antibody (e.g. recombinant anti-glypican 3 antibody) of the present invention is obtained by overlapping PCR from a plurality of oligonucleotides constructed so as to have nucleotide sequences that overlap with each other, which is in turn ligated with DNA encoding human antibody constant region so as to form an in-frame codon sequence. The DNA ligated in this manner is then inserted in an operable manner into an expression vector, and introduced into a host. The antibody (e.g. anti-glypican 3 antibody) encoded by the DNA is expressed by culturing the host. The expressed antibody (e.g. anti-glypican 3 antibody) is isolate by purifying from the host culture medium (EP 239400, WO 96/02576, and so forth). The FRs of the humanized antibody (e.g. anti-glypican 3 antibody) ligated with CDR are selected in such a manner that the complementarity-determining region forms a high-quality antigen-binding site for the antigen. As necessary, the amino acid sequence may be modified by suitable substitution of the amino acid residues constituting the antibody variable region FRs, so as to enable the complementarity-determining region of reshaped antibody to form a suitable antigen-binding site for the antigen (K. Sato et al., Cancor Res. (1993) 53, 851-856)

In addition to modifications related to humanization as discussed above, further modifications can be introduced, for example, to improve the biochemical characteristics of the antibody, e.g., the binding activity with the antigen.
recognized by the antibody (e.g. anti-glypican 3 antibody). Modifications in the context of the present invention can be suitably carried out by known methods such as site-specific mutagenesis (refer, for example, to Kunkel, Proc. Natl. Acad. Sci. USA (1985) 82, 488), PCR mutation, cassette mutation, and so forth. The amino acid sequence of the modified antibody with improved biochemical characteristics generally has at least 70%, more preferably at least 80%, and even more preferably at least 90% (for example, at least 95%, 97%, 98%, 99%, and so forth) identity and/or similarity with the amino acid sequence constituting the antibody submitted to the modification (that is, the antibody on which the modified antibody is based). As used herein, the sequence identity and/or similarity refers to the proportion of amino acid residues that are identical (identical residues) or similar (amino acid residues classified into the same group based on the general characteristics of the amino acid side chain) after aligning the sequences and inserting gaps as necessary in such a manner that the sequence identity assumes a maximum. The naturally occurring amino acid residues can generally be classified into the following groups based on the properties of the side chain: (1) hydrophobic: alanine, isoleucine, valine, methionine, and leucine; (2) neutral hydrophilic: asparagine, glutamine, cysteine, threonine, and serine; (3) acidic: aspartic acid and glutamic acid; (4) basic: arginine, histidine, and lysine; (5) residues that affect chain
orientation: glycine and proline; and (6) aromatic: tyrosine, tryptophan, and phenylalanine.

A specific suitable embodiment of a modification for enhancing antibody function includes an improvement in the cytotoxicity exhibited by the antibody, such as humanized anti-glypican 3 antibody. Preferred examples of cytotoxicity are antibody-dependent cell-mediated cytotoxicity (ADCC) and complement-dependent cytotoxicity (CDC). In the present invention, CDC activity denotes cytotoxicity due to the complement system, while ADCC refers to an activity of damaging the target cell when a specific antibody attaches to a cell surface antigen on the target cell and an Fcγ receptor-bearing cell (e.g., an immunocyte) binds to the Fc portion of the antibody through the Fcγ receptor. Whether a test antibody has ADCC activity or has CDC activity can be measured by known methods (for example, Current Protocols in Immunology, Chapter 7, Immunologic studies in humans. Editor, John E. Coligan et al., John Wiley & Sons, Inc., (1993)).

In specific terms, effector cells, a complement solution, and target cells are first prepared.

(1) Preparation of effector cells

The spleen is isolated from, for example, CBA/N mouse, and the spleen cells are separated on RPMI1640 medium (Invitrogen). After washing with the same medium containing 10% fetal bovine serum (FBS, HyClone) the cell concentration
is adjusted to 5 x 10^6 cells/mL to prepare the effector cells.

(2) Preparation of complement solution

The complement solution can be prepared by a 10X dilution with 10% FBS-containing medium (Invitrogen) of Baby Rabbit Complement (CEDARLANE).

(3) Preparation of target cells

Cells that express an antigen protein to which the test antibody binds are cultured with 0.2 mCi ^{51}Cr sodium chromate (GE Healthcare Bioscience) for 1 hour at 37°C in DMEM medium containing 10% FBS in order to radiolabel the target cells. The following, inter alia, can be used as the cells that express an antigen protein to which the test antibody binds: cells transformed with a gene that encodes an antigen protein to which the test antibody binds, ovarian cancer cells, prostate cancer cells, breast cancer cells, uterine cancer cells, liver cancer cells, lung cancer cells, pancreatic cancer cells, kidney cancer cells, bladder cancer cells, and colon cancer cells. After radiolabeling, the cells are washed three times with RPMI1640 medium containing 10% FBS and the cell concentration is adjusted to 2 x 10^5 cells/mL to prepare the target cells.

The ADCC activity and the CDC activity can be measured by the following methods. In order to measure the ADCC activity, 50 µL of the target cells and 50 µL of the test antibody are added to a 96-well U-bottom plate (Becton Dickinson) and reacted for 15 minutes on ice. To the reaction mixture is
added 100 μL effector cells and incubated for 4 hours in a carbon dioxide incubator. A final test antibody concentration is preferably within the range from 0 to 10 μg/mL. After incubation, 100 μL of the supernatant is recovered and the radioactivity of the supernatant is measured using a gamma counter (COBRA II AUTO-GAMMA, MODEL D5005, Packard Instrument Company). The cytotoxicity can be calculated using the radioactivity according to the following equation:

\[
\frac{(A - C)}{(B - C)} \times 100
\]

wherein A is the radioactivity (cpm) for the sample using the particular test antibody; B is the radioactivity (cpm) of the sample to which 1% NP-40 (Nacalai Tesque) has been added; and C is the radioactivity (cpm) for the sample containing only the target cells.

To measure the CDC activity, 50 μL of the target cells and 50 μL of the test antibody are added to a 96-well U-bottom plate (Becton Dickinson) and reacted for 15 minutes on ice. To the reaction mixture is added 100 μL complement solution and incubated for 4 hours in a carbon dioxide incubator. A final test antibody concentration is preferably within the range from 0 to 3 μg/mL. After cultivation, 100 μL of the supernatant is recovered and the radioactivity of the supernatant is measured with a gamma counter. The cytotoxicity can be calculated in the same manner as for measurement of the ADCC activity.
When the cytotoxicity due to an antibody conjugate is to be measured, 50 μL of the target cells and 50 μL of the test antibody conjugate are each added to a 96-well flat-bottom plate (Becton Dickinson) and reacted for 15 minutes on ice. The plate is then incubated for from 1-4 hours in a carbon dioxide incubator. A final test antibody concentration is preferably within the range from 0 to 3 μg/mL. After incubation, 100 μL of the supernatant is recovered and the radioactivity of the supernatant is measured with a gamma counter. The cytotoxicity can be calculated in the same manner as for measurement of the ADCC activity.

The heavy chain and light chain variable regions of the antibody (e.g. anti-glypican 3 antibody) are, as described above, generally composed of 3 CDRs and 4 FRs. The amino acid residues submitted to "modification" in a preferred embodiment of the present invention can be suitably selected, for example, from the amino acid residues constituting the CDRs or FRs. Modification of the amino acid residues constituting the CDRs may in some cases cause a decline in the antigen binding capacity of the antibody (e.g. anti-glypican 3 antibody) involved in the modification. Accordingly, the antibody (e.g. anti-glypican 3 antibody) amino acid residues submitted to "modification" in the present invention are preferably selected from the amino acids residues constituting the FRs, but not limited to. When it is confirmed that modification of
amino acid residues located in the CDRs do not cause a decline in the antigen binding capacity of the antibody (e.g. anti-glypican 3 antibody) involved in the modification, then such amino acid residues may be selected for modification.

Those skilled in the art can readily find in a public database, such as the Kabat database, amino acid sequences constituting the antibody variable region FR that actually occur in an organism such as mouse or human.

A preferred embodiment of the present invention provides a humanized antibody (e.g. anti-glypican 3 antibody) having plasma kinetics modulated by the method of the present invention. For example, the humanized antibody (e.g. anti-glypican 3 antibody) is a humanized antibody comprising a complementarity-determining region (CDR) derived from a nonhuman animal, a framework region (FR) of human origin, and a human constant region, wherein at least one amino acid residue that can be exposed on the antibody surface in the CDR or FR has a different charge from that of the amino acid residue in the corresponding position of the CDR or FR of the original antibody and wherein the humanized antibody has plasma kinetics modulated in comparison to the chimeric antibody having the same constant region.

Another preferred embodiment of the present invention provides a human antibody (e.g. human anti-glypican 3 antibody) having plasma kinetics modulated by the method of the present invention. The human antibody, for example, is a
human antibody comprising a complementarity-determining region (CDR) of human origin, a framework region (FR) of human origin, and a human constant region, wherein at least one amino acid residue that can be exposed on the antibody surface in the CDR or FR has a different charge from that of the amino acid residue in the corresponding position of the CDR or FR of the original antibody and wherein the human antibody has plasma kinetics modulated in comparison to a chimeric antibody having the same constant region.

The aforementioned human constant region preferably denotes a region comprising a wild-type human Fc region, but a modified Fc can also be suitably used. Such a "modified Fc" may include a modified Fc as prepared by the modification of an amino acid residue constituting the Fc as well as a modified Fc as prepared by a modification of a modification already executed in the Fc moiety. A preferred example of such a modification of a modification is the modification of the nature of the sugar chain modification attached to the Fc portion. One preferred specific example is the "antibody having reduced content of fucose attached to the Fc region" that is specifically disclosed herein as a reference example.

The term "antibody having reduced content of fucose attached to the Fc region" denotes an antibody for which the amount of bound fucose has been significantly reduced in comparison to the control antibody, and preferably fucose is undetectable. Fucose is generally bound to the N-glycoside
linked sugar chains that are bound at the two sugar chain-binding sites present in the Fc region of the two heavy chain molecules that form a single antibody molecule. The term "antibody having reduced content of fucose attached to the Fc region" denotes an antibody that, in comparison with the ordinary antibody as the control, has a fucose content no greater than 50%, preferably no greater than 25%, more preferably no greater than 10%, and particularly preferably no greater than 0% of the total sugar chain content of the control antibody. The fucose content can be measured using the analytical procedure provided in the reference examples below. The method for producing such a fucose-depleted antibody may include the method described in the reference examples of the present invention, as well as a method for producing antibodies with fucose transferase-deficient animal cells (Biotechnol. Bioeng. (2004) 87(5), 614-22) and a method for producing antibodies with animal cells in which the complex branched sugar chain modification is modified (Biotechnol. Bioeng. (2006) 93(5), 851-61). In addition, suitable examples of production methods using non-animal cells as the host cells may include a method for producing antibodies with plant cells (Nature Biotechnology (2006) 24, 1591-7) and with yeast cells (Nature Biotechnology (2006) 24, 210-5).

A preferred embodiment of the preparation method of the present invention is a method of preparing an antibody (e.g. anti-glypican 3 antibody) having modulated plasma kinetics,
comprising (a) modifying a nucleic acid that encodes a polypeptide comprising at least one amino acid residue that can be exposed on the surface of the antibody (e.g. anti-glypican 3 antibody), such that the charge of the amino acid residue(s) is changed; (b) culturing a host cell in such a manner that the nucleic acid modified in the step (a) is expressed; and (c) recovering the antibody (e.g. anti-glypican 3 antibody) from the host cell culture.

In the method of the present invention, the term "modifying a nucleic acid" refers to a modification of the nucleic acid sequence so as to provide a codon that corresponds to the amino acid residue that is introduced by the "modification" of the present invention. More specifically, this term refers to a modification of a nucleic acid that contains a codon that will undergo modification, wherein said modification changes the codon corresponding to the pre-modification amino acid residue to the codon of the amino acid residue that is introduced by the modification. Generally this term means the implementation of a genetic process or mutagenic treatment so as to replace at least one base of the codon-comprising nucleic acid to provide a codon that encodes the target amino acid residue. That is, a codon encoding the amino acid residue submitted to modification is replaced by a codon encoding the amino acid residue that is introduced by the modification. The nucleic acid modification can be suitably carried out by those skilled in the art using known
technology, for example, site-specific mutagenesis, PCR mutagenesis, and so forth.

The nucleic acid prepared in the present invention is generally placed (inserted) into a suitable vector and introduced into a host cell. There are no particular limitations on the vector as long as it has the capability to stably maintain the inserted nucleic acid. For example, with reference to the use of E. coli as the host, pBluescript vector (Stratagene) is preferred as a cloning vector, although various commercially available vectors can be used. An expression vector is particularly useful in those instances where the vector is used to produce polypeptide of the present invention. There are no particular limitations on the expression vector as long as it can express polypeptide in vitro, in E. coli, in cell culture, or within an organism. Preferred examples include pBEST vector (Promega) for in vitro expression, pET vector (Invitrogen) for E. coli, pME18S-FL3 vector (GenBank Accession No. AB009864) for cell culture, and pME18S vector (Mol. Cell Biol. (1988) 8, 466-472) for an organism. DNA according to the present invention can be inserted into the vector by standard methods, for example, by the ligase method utilizing restriction enzyme sites (Current Protocols in Molecular Biology, edited by Ausubel et al. (1987) Publish. John Wiley & Sons, sections 11.4-11.11).

There are no particular limitations on the host cell referenced above, and various host cells can be used depending
on the purpose. Examples of cells for the expression of polypeptide include bacterial cells (e.g., Streptococcus, Staphylococcus, E. coli, Streptomyces, Bacillus subtilus), fungi cells (e.g., yeast, Aspergillus), insect cells (e.g., Drosophila S2, Spodoptera Sf9), animal cells (e.g., CHO, COS, HeLa, C127, 3T3, BHK, HEK293, Bowes melanoma cells), and plant cells. The vector can be introduced into the host cell by known methods, for example, calcium phosphate precipitation, electropulse poration (Current Protocols in Molecular Biology, edited by Ausubel et al. (1987) John Wiley & Sons, sections 9.1-9.9), lipofection, and microinjection.

An appropriate secretory signal can be suitably incorporated into the antibody (e.g. anti-glypican 3 antibody) in order to bring about secretion of the antibody expressed in the host cell into a vesicle lumen, the periplasmic space, or the extracellular environment. Such a signal can suitably be the native signal sequence characteristic of the antibody (e.g. anti-glypican 3 antibody) or can be a heterologous signal sequence.

The antibody (e.g. anti-glypican 3 antibody) produced as described above can be recovered by collecting the culture medium in those instances where the antibody of the present invention is secreted into the culture medium. In those instances where the antibody (e.g. anti-glypican 3 antibody) of the present invention is produced within the cell, the cell is first lysed and then the antibody is recovered.
Known methods can be suitably used to purify the antibody (e.g. anti-glypican 3 antibody) of the invention recovered from the recombinant cell culture, for example, ammonium sulfate and ethanol precipitation, acid extraction, anion- or cation-exchange chromatography, phosphocellulose chromatography, hydrophobic interaction chromatography, affinity chromatography, hydroxyapatite chromatography, and lectin chromatography.

The present invention also relates to a composition (a drug) comprising a pharmaceutically acceptable carrier and the antibody (e.g. anti-glypican 3 antibody) (for example, IgG antibody) having plasma kinetics modulated by the method of the invention.

The pharmaceutical composition in the context of the present invention generally refers to an agent for the treatment or prevention of a disease or for the detection or diagnosis of a disease.

The pharmaceutical composition of the present invention can be formulated by a method known to those skilled in the art. For example, the formulation can be used by a non-oral route in the form of an injectable, i.e., a suspension or aseptic solution with water or other pharmaceutically acceptable liquid. For example, the formulation can be prepared by combining the antibody with a pharmacologically acceptable carrier or medium, in particular with sterile water or physiological saline, plant oil, emulsifying agent,
suspending agent, surfactant, stabilizer, flavorant, diluent, carrier, preservative, binder, and so forth, and mixing in the unit dosage state required by generally established pharmaceutical practice. The amount of effective component in the formulations is selected in such a manner that a suitable dose in the indicated range is obtained.

A sterile composition for injection can be suitably formulated in accordance with standard pharmaceutical practice using a carrier such as distilled water for injection.

The aqueous solution for injection can be, for example, an isotonic solution containing physiological saline, dextrose, and other adjuvants (for example, D-sorbitol, D-mannose, D-mannitol, sodium chloride). Suitable dissolution auxiliaries (e.g., an alcohol such as ethanol or a polyalcohol such as propylene glycol or polyethylene glycol) and nonionic surfactant (e.g., Polysorbate 80<sup>®</sup>, HCO-50) can also be used as appropriate.

Oily liquids include sesame oil and soy oil. Benzyl benzoate and/or benzyl alcohol can also be suitably used as dissolution auxiliaries. A buffer (for example, phosphate buffer solution, sodium acetate buffer solution), soothing agent (for example, procaine hydrochloride), stabilizer (for example, benzyl alcohol and phenol), and oxidation inhibitor can also be incorporated as appropriate. The injection solution prepared as described above is generally filled into suitable ampoules.
The pharmaceutical composition of the present invention can be administered preferably by non-oral administration. For example, it can be formulated as an injectable composition, a transnasal composition, a composition for inhalation administration, or a transdermal composition. It can be suitably administered systemically or locally by, for example, intravenous injection, intramuscular injection, intraperitoneal injection, or subcutaneous injection.

The method of administration can be selected as appropriate depending on the patient's age and symptoms. The dosage of the pharmaceutical composition comprising the antibody or polynucleotide encoding the antibody can be selected, for example, in the range of 0.0001 mg to 1000 mg per 1 kg body weight per unit dose. Or, the dosage can be formulated or set at a dosage of 0.001 to 100,000 mg per patient; however, the present invention is not necessarily limited to these numerical values. The dosage and method of administration will vary depending on the patient's body weight, age, and symptoms, and an appropriate dosage and method of administration can be selected by those skilled in the art based on a consideration of these factors.

The present invention also provides nucleic acid encoding the antibody (e.g. anti-glypican 3 antibody) (for example, humanized anti-glypican 3 antibody) having plasma kinetics modulated by the method of the present invention. A vector carrying such a nucleic acid is also encompassed by the
present invention.

The present invention also provides a host cell that comprises the aforementioned nucleic acid. The type of the host cell is not particularly limited, and it may be, for example, a bacterial cell such as E. coli or any of various animal cells. The host cell can be used as appropriate in a production system for the production or expression of the antibody (e.g. anti-glypican 3 antibody) of the present invention. Thus, the present invention also provides a production system that may be used to produce the antibody (e.g. anti-glypican 3 antibody) using the aforementioned host cell. An in vitro or in vivo production system can suitably be used as the production system. Eukaryotic cells and prokaryotic cells are suitably employed as the host cells used in the in vitro production system.

The eukaryotic cells used as the host cell may include animal cells, plant cells, and fungi cells. The animal cells may include mammalian-type cells such as CHO (J. Exp. Med. (1995) 108, 945), COS, HEK293, 3T3, myeloma, BHK (baby hamster kidney), HeLa, Vero, and so forth; amphibian cells such Xenopus laevis oocytes (Valle et al., Nature (1981) 291, 338-340); and insect cells such as Sf9, Sf21, and Tn5. For example, CHO-DG44, CHO-DX11B, COS7 cells, HEK293 cells, and BHK cells are suitably used for expression of the anti-glypican 3 antibody of the present invention. The use of CHO cells as the host cell is particularly preferred when high levels of
expression in animal cells is intended. Transfection of the recombinant vector into the host cell is suitably carried out using the calcium phosphate technique, DEAE dextran technique, techniques that employ the cationic liposome DOTAP (Boehringer Mannheim), electroporation technique, lipofection technique, and so forth.

With regard to plant cells, Lemna minor and cells derived from Nicotiana tabacum are known as protein production systems, and the anti-glypican 3 antibody of the present invention can be produced by callus culture techniques using these cells. With regard to fungi cells, protein expression systems using yeast cells, e.g., Saccharomyces (e.g., Saccharomyces cerevisiae, Schizosaccharomyces pombe), and protein expression systems using filamentous fungi, e.g., Aspergillus (e.g., Aspergillus niger), are known and can be used as host cells to produce the anti-glypican 3 antibody of the present invention.

With regard to the use of prokaryotic cells, any production systems using bacterial cells are suitably used. Production systems that use E. coli as described above or B. subtilus are known for the bacterial cells, and any of these bacterial cells can be suitably used to produce the antibody (e.g. anti-glypican 3 antibody) of the present invention.

In order to produce the antibody (e.g. anti-glypican 3 antibody) using a host cell of the present invention, the host cell transformed with the expression vector comprising polynucleotide encoding the antibody of the present invention
is cultured to express a polynucleotide coding for the antibody (e.g. anti-glypican 3 antibody). The host cells can be cultured according to known methods. When an animal cell is used as the host, for example, DMEM, MEM, RPMI1640, or IMDM is suitably used as the culture medium. A serum auxiliary, e.g., FBS, fetal calf serum (FCS), and so forth, may be added. The cells can also be cultured in a serum-free medium. The cells can be cultured at a pH of about 6-8, depending on the host cell. Culture is generally run for about 15 to 200 hours at about 30 to 40°C, with medium replacement, aeration, and stirring as necessary.

On the other hand, production systems based on an animal or plant are available as in vivo systems for producing the antibody (e.g. anti-glypican 3 antibody) of the present invention. Polynucleotide encoding the antibody (e.g. anti-glypican 3 antibody) of the present invention is introduced into an animal or plant and the antibody is produced within the animal or plant and is recovered. The term "host" as used herein encompasses such animals and plants.

When an animal is used as the host, production systems based on mammals and insects are available. Goat, pig, sheep, mouse, cow, and so forth, are suitably used as the mammal (Vicki Glaser, SPECTRUM Biotechnology Applications (1993)). A transgenic animal may also be employed.

For example, a polynucleotide encoding the antibody (e.g. anti-glypican 3 antibody) of the present invention can be
produced in the form of a fusion gene with a gene encoding a polypeptide that is specifically produced in milk, such as goat β-casein. A polynucleotide fragment comprising the fusion gene is then injected into a goat embryo and is transplanted into a female goat. The antibody (e.g. anti-glypican 3 antibody) of interest is obtained from milk produced by the transgenic goat born out of the goat that received the embryo or from milk produced by the offspring of the transgenic goat. Suitable hormones may be administered as appropriate to the transgenic goat in order to increase the amount of antibody (e.g. anti-glypican 3 antibody)-containing milk produced from the transgenic goat (Ebert et al., Bio/Technology (1994) 12, 699-702).

The silkworm is an example of an insect that can be used to produce the antibody (e.g. anti-glypican 3 antibody) of the present invention. When the silkworm is employed, the silkworm may be infected with a baculovirus having a polynucleotide encoding the desired antibody (e.g. anti-glypican 3 antibody) inserted into its viral genome. The antibody (e.g. anti-glypican 3 antibody) of interest is obtained from the body fluids of the infected silkworm (Susumu et al., Nature (1985) 315, 592-4).

The tobacco plant is an example of a plant that may be used to produce the antibody (e.g. anti-glypican 3 antibody) of the present invention. When the tobacco plant is used, polynucleotide encoding the antibody (e.g. anti-glypican 3
antibody) of interest is inserted into a plant expression vector, such as pMON 530, and the resulting recombinant vector is transfected into a bacterium such as Agrobacterium tumefaciens. This bacterium is then used to infect a tobacco plant (for example, Nicotiana tabacum) (Ma et al., Eur. J. Immunol. 1994 24, 131-8), and the desired antibody (e.g. anti-glypican 3 antibody) is obtained from the leaves of the infected tobacco plant. Lemna minor can be similarly infected with such a bacterium and the desired antibody (e.g. anti-glypican 3 antibody) can be obtained from cells of the cloned infected Lemna minor (Cox K. M. et al., Nat. Biotechnol. 2006 24(12), 1591-7).

The antibody (e.g. anti-glypican 3 antibody) of the present invention obtained in the manner described above can be isolated from inside or outside of the host cells (e.g., culture medium or milk) and can be purified into a substantially pure homogeneous antibody. The separation and purification techniques generally used for polypeptide purification can be suitably used for antibody separation and purification in the present invention, but is not limited. For example, the antibody can be suitably separated and purified by a suitable selection and combination of column chromatography, filtration, ultrafiltration, salting out, solvent precipitation, solvent extraction, distillation, immunoprecipitation, SDS-polyacrylamide gel electrophoresis, isoelectric electrophoresis, dialysis, recrystallization, and
so forth.

The chromatographic techniques include affinity chromatography, ion-exchange chromatography, hydrophobic chromatography, gel filtration chromatography, reverse-phase chromatography, adsorption chromatography, and so forth (Strategies for Protein Purification and Characterization: A Laboratory Course Manual. Ed. Daniel R. Marshak et al. (1996) Cold Spring Harbor Laboratory Press). Chromatography can be carried out using liquid-phase chromatography, for example, HPLC, FPLC, and so forth. The column used in affinity chromatography includes, for example, a protein A column or protein G column. Hyper D, POROS, and Sepharose F. F. (Pharmacia) are examples of protein A based columns.

Another preferred embodiment of the present invention is a method of preparing the antibody (e.g. anti-glypican 3 antibody) with modulated plasma kinetics of the present invention, comprising the step of culturing a host cell of the present invention as described above and recovering the antibody (e.g. anti-glypican 3 antibody) from the cell culture.

All of the literature cited in the specification are incorporated herein by reference.

The present invention is specifically described in the following, but the present invention is not limited to the examples provided below.
Example 1

(1) Construction of point-mutation genes of the humanized H0L0 antibody

Various point-mutation genes were constructed starting from a gene encoding anti-glypican 3 antibody comprising the CDR of the humanized GC33 antibody disclosed in WO 2006/046751. OligoDNAs designed based on the sequences of the sense and antisense chains containing the modification sites were synthesized. A plurality of point-mutation genes were constructed using the commercial QuickChange Site-Directed Mutagenesis Kit (Stratagene). Construction of the point-mutation genes was carried out by PCR under the following conditions. After heating for 30 seconds at 95°C, a reaction mixture of 10 ng template plasmid, 10 pmol forward chain and reverse chain synthetic oligo-DNAs and 10X buffer, dNTP mix, and Pfu Turbo DNA Polymerase provided with the kit was subjected to 18 cycles of 95°C 30 sec, 55°C 1 min and 68°C 4 min. The DpnI provided with the kit was added to the reaction mixture, and restriction digestion with the restriction enzyme was carried for 1 hour at 37°C. DH5α competent cells (Toyobo) were transformed with the resulting reaction solution to obtain transformants. The introduction of point mutation was confirmed by determining the nucleotide sequence of the plasmid DNA isolated from the transformants. Each point-mutation gene was cloned into expression vectors capable of expressing the insert gene in animal cells. Modified genes
were prepared by modifications as described below.

Transient expression of the humanized HOLO antibody and its point mutation-modified antibodies was carried out using polyethyleneimine (Polysciences Inc.). HEK293 cells were separated by trypsin EDTA (Invitrogen), and seeded to a 10 cm² culture dish at 6 x 10⁶ cells/10 mL. The next day, SFMII culture medium and polyethyleneimine were mixed with a heavy chain expression plasmid DNA and a light chain expression plasmid DNA according to the manufacturer's instructions, and the resulting mixture was left stand for 10 minutes at room temperature. The entire mixture was added dropwise to the culture dish containing the HEK293 cells seeded as described above. The culture supernatant was recovered after approximately 72 hours and the expressed humanized HOLO antibody and its point mutation-modified antibodies were purified using rProteinA Sepharose™ Fast Flow (GE Healthcare) according to the manufacturer's instructions.

(1-1) Modification of the Tm value of the humanized HOLO antibody

The thermal denaturation midpoint temperature (Tm) was determined by the top of the denaturation peak in the thermogram (Cp versus T) obtained after heating the test sample solution at a constant programmed heating rate. The Tm value of the humanized HOLO antibody was measured using a sample solution for DSC measurement prepared as described in the following. The antibody solution (corresponding to 50 to
100 μg) filled in a dialysis membrane was first dialyzed for 24 hours against a dialysis external solution of 20 mol/L sodium acetate buffer solution (pH 6.0) containing 150 mmol/L sodium chloride. Subsequently, the sample solution was adjusted at its antibody concentration of 50 to 100 μg/mL with dialysis external solution and used as the sample solution for DSC measurement.

A suitable DSC instrument, for example, DSC-II (Calorimetry Sciences Corporation), is used for this experiment. The sample solution prepared as described above and the reference solution (dialysis external solution) were thoroughly degassed, and each of these test specimens was placed in a calorimeter cell and was thermally equilibrated at 40°C. A DSC scan was then run from 40°C to 100°C at a scan rate of approximately 1 K/minute. The results of this measurement are given as the top of the denaturation peak as a function of temperature. The thermal denaturation midpoint temperature of the humanized HOLO antibody was calculated by peak assignment of the Fab domain according to Rodolfo et al., *Immunology Letters* (1999), 47-52. As a specific example, the chart obtained from differential scanning calorimetry (DSC) on the Hspo2.2Lspo2.2 (Hu2.2Lu2.2) antibody is shown in Figure 1.

The humanized HOLO antibody, comprising the heavy chain shown in SEQ ID NO: 1 and the light chain shown in SEQ ID NO: 7, has a Tm value of 76.6°C as calculated by the method
described above. The Tm values of Synagis and Herceptin, provided as examples of existing antibodies, are measured at 85.4°C and 81.8°C, respectively. It was thus shown that the Tm value of the humanized H0L0 antibody is lower than that of existing antibodies.

Modified antibodies were therefore prepared from humanized H0L0 antibody with the aim of raising the Tm value. Modifications of V37I, A40P, M48I, and L51I were introduced into FR2 of the H0L0 antibody heavy chain shown in SEQ ID NO: 1 to prepare the antibody H1b (SEQ ID NO: 2), where its subclass was changed from VH1b to VH4. The Tm value was improved to 79.1°C. Also the H0L0 antibody light chain shown in SEQ ID NO: 7 was modified by introducing L42Q, S43A, and Q50R modifications into the FR2 which changed the subclass from VK2 to VK3, and introducing V2I modification to replace the V2 of FR1 with I (germline sequence), thereby L4 (SEQ ID NO: 8) was prepared. The Tm value of each antibody was measured as described above. The Tm value of H15L0 and H0L4 was 79.2°C and 77.2°C, respectively, which shows improvement from the Tm value 76.6°C of H0L0. The Tm value of the H15L4 antibody comprising the combination of these two modifications was improved to 80.5°C.

(1-2) Modification of the pI value of the humanized H0L0 antibody

The plasma half-life of an antibody is extended by lowering the pI value exhibited by the antibody. In contrast,
the tissue translocation characteristics of an antibody are improved by increasing the antibody’s pI. It is unknown whether either increase or decrease in the pI value of an antibody effective in cancer treatment would enhance the tumor-suppressing effect. Therefore, modified humanized HOLO antibody with a lowered pI and modified humanized HOLO antibody with an increased pI were prepared and the antitumor activity of them was compared to investigate if either modification had a higher tumor-suppressing activity.

The pI value of each antibody was calculated based on the analysis by isoelectric electrophoresis. Electrophoresis was carried out as described in the following. Using PhastSystem Cassette (Amersham Bioscience), Phast-Gel Dry IEF (Amersham Bioscience) gel was swollen for about 60 minutes with a swelling solution with the composition given below.

(a) Composition of the swelling solution for high pI:
1.5 mL 10% glycerol
100 μL Pharmalyte 8-10.5 for IEF (Amersham Bioscience)

(b) Composition of the swelling solution for low pI:
1.5 mL purified water
20 μL Pharmalyte 8-10.5 for IEF (Amersham Bioscience)
80 μL Pharmalyte 5-8 for IEF (Amersham Bioscience)

Approximately 0.5 μg antibody was loaded on the swollen gel and isoelectric electrophoresis was run using the PhastSystem (Amersham Bioscience) controlled by the program
described below. The sample was added to the gel at Step 2 of this program. A Calibration Kit for pI (Amersham Bioscience) was used for the pI markers.

Step 1: 2000 V, 2.5 mA, 3.5 W, 15°C, 75 Vh
Step 2: 200 V, 2.5 mA, 3.5 W, 15°C, 15 Vh
Step 3: 2000 V, 2.5 mA, 3.5 W, 15°C 410 Vh

After electrophoresis, the gel was fixed with 20% TCA and silver staining was then carried out using Silver Staining Kit, Protein (Amersham Bioscience) according to the instructions provided with the kit. After staining, the isoelectric point of each antibody (test sample) was calculated based on the already known isoelectric points of the pI markers. The electrophoretogram from the high pI isoelectric electrophoresis is shown in Figure 2 and the electrophoretogram from the low pI isoelectric electrophoresis is shown in Figure 3.

(a) Modifications that raised the pI

Hspu2.2 (Hu2.2) (SEQ ID NO: 6) was prepared, in which the Q43K, D52N, and Q107R modifications were additionally implemented in H15. Lspu2.2 (Lu2.2) (SEQ ID NO: 12) was also prepared, in which the E17Q, Q27R, Q105R and S25A modifications were implemented in L4, where the S25A modification replaces S25 in CDR2 with A (abundant in a germline). The Tm value of the Hspu2.2Lspu2.2 (Hu2.2Lu2.2) antibody composed of Hspu2.2 (Hu2.2) and Lspu2.2 (Lu2.2) was measured at 76.8°C and its pI value was measured at 9.6. Since
the pI of the HOLO antibody is 8.9, the pI of the Hspu2.2Lspu2.2 (Hu2.2Lu2.2) antibody was increased by 0.7.

(b) Modifications that lowered the pI

Hspd1.8 (Hd1.8) (SEQ ID NO: 5) was prepared, in which the K19T, Q43E, K63S, K65Q, and G66D modifications were additionally implemented in H15. Lspd1.6 (Ld1.6) (SEQ ID NO: 11) was prepared by making the following modifications: the Q27E modification in L4; modification of KISRVE at 79-84 of the FR3 in L4 to TISSLQ; and the S25A modification to achieve the same modification as for Lspu2.2 (Lu2.2). The pI value of the Hspd1.8Lspd1.6 (Hd1.8Ld1.6) antibody composed of Hspd1.8 (Hd1.8) and Lspd1.6 (Ld1.6), was measured at 7.4. Since the pI of the HOLO antibody is 8.9, the pI of the Hspd1.8Lspd1.6 (Hd1.8Ld1.6) antibody was reduced by 1.5.

(2) Evaluation by competitive ELISA of the binding activity of the point-mutation modified antibodies from the HOLO antibody

The HOLO antibody and its point mutation-modified antibodies purified in (1) was evaluated by competitive ELISA. 100 μL of the soluble GPC3 core polypeptide (SEQ ID NO: 13) at 1 μg/mL was added to each well of a 96-well plate. The soluble GPC3 core polypeptide was immobilized on the plate by allowing the plate to stand overnight at 4°C. The soluble GPC3 core polypeptide immobilized on the plate was washed 3 times with a washing buffer using Skan WASHER400 (Molecular Devices); and blocked with 200 μL blocking buffer at 4°C for at least 30 min.
The blocked plate on which soluble GPC3 core polypeptide was immobilized was then washed 3 times with washing buffer using the Skan WASHER400. Subsequently, each well of the plate received 200 μL of a mixture containing 100 μL of biotinylated H0L0 antibody (final concentration = 0.3 μg/mL) and 100 μL of the H0L0 antibody or its point mutation-modified antibody (at various concentrations). The H0L0 antibody was biotinylated using Biotin Labeling Kit (Roche) according to the instructions provided with the kit. The plate was left stand for 1 hour at room temperature, then washed 5 times with washing buffer using the Skan WASHER400 (Molecular Devices). 100 μL goat anti-streptavidin alkaline phosphatase (ZYMED), diluted 20,000X with substrate buffer, was added to each well, and the resulting plate was left stand for 1 hour at room temperature, and then washed 5 times with washing buffer using the Skan WASHER400. Phosphatase Substrate (Sigma) was prepared at 1 mg/mL in the substrate buffer, added at 100 μL per well for 1 hour. The absorbance at 405 nm of the reaction solution in each well was measured using Benchmark Plus (BIO-RAD), with the control absorbance at 655 nm.

As shown by Figure 4, the antigen binding activity of the H15L4 antibody was shown to be almost the same as that of the H0L0 antibody that had been submitted to the modification. In addition, as shown in Figure 5, the antigen binding activity of the Hspu2.2Lspu2.2 (Hu2.2Lu2.2) antibody was shown to be
about the same as that of the H0L0 antibody submitted to the modification. Moreover, as shown in Figure 6, the antigen binding activity of the Hspdl.8Lspdl.6 (Hdl.8Ld1.6) antibody was shown to be about the same as that of the H0L0 antibody submitted to the modification.

Reference Example 2

Disruption of the fucose transporter gene in CHO cells

(1) Construction of the targeting vector

(1-1) Construction of the KO1 vector

The BamHI site and the TGC3G sequence were added at the 5' side of the start codon of the hygromycin resistance gene (Hygr) by PCR using pcDNA3.1/Hygro (Invitrogen) with primers Hyg5-BH and Hyg3-NT to obtain the same sequence as on the 5' side of the start codon of the fucose transporter gene. Also the NotI site was added on the 3' side including the region up to the SV40 polyA addition signal. Then the Hygr gene was excised.

forward primer

Hyg5-BH 5'-GGATCTGCGCATGAAAAAGCCCTGAACCTCACCC-3' (SEQ ID NO: 14)

reverse primer

Hyg3-NT 5'-GCCGGCCGCTATTCTTTGCCCTCGGACG-3' (SEQ ID NO: 15)

Fucose transporter targeting vector ver. 1 (referred to as the KO1 vector) was constructed by inserting the 5' side of the fucose transporter (from the SmaI at No. 2,780 of SEQ ID NO: 16 to the BamHI at No. 4,232), the 3' side (from No. 4,284
to the SacI at No. 10,934), and the Hygr fragment into the pMC1DT-A vector (Yagi T., Proc. Natl. Acad. Sci. USA (1990) 87, 9918-22). A characteristic feature of this vector is that, when homologous recombination has occurred, Hygr is expressed under the promoter of the fucose transporter, since a promoter is not attached to the Hygr gene. However, when only one copy of the vector is introduced into the cell by homologous recombination, Hygr is not always expressed to a degree sufficient for exhibiting hygromycin B resistance. The KO1 vector was cleaved at the NotI site and was transfected into the cell. It is thought that 41 base pairs in exon 1, including the start codon, will be deleted from the fucose transporter by introduction of the KO1 vector, resulting in a loss of function.

(1-2) Construction of pBSK-pgk-1-Hygr

pBSK-pgk-1 was constructed by excising the pgk-1 gene promoter from the pKJ2 vector (Popo H., Biochemical Genetics (1990) 28, 299-308) with EcoRI-PstI and cloning it into the EcoRI-PstI sites of pBluescript (Stratagene). With regard to Hygr, an EcoT22I site and a Kozak sequence were added on the 5' side of Hygr and a BamHI site was added on the 3' side including the region up to the SV40 polyA addition signal by PCR using pcDNA3.1/Hygro with the Hyg5-AV and Hyg3-BH primers. Then the Hygr gene was excised. forward primer
Hyq5-AV 5'-ATGCATGCCACCATGAAAAAGCCTGAACTCACC-3' (SEQ ID NO: 17)
reverse primer

Hyg3-BH 5'-GGATCCAGGCTTTTACACTTTTGCTTC-3' (SEQ ID NO: 18)

pBSK-pgk-1-Hygr was constructed by inserting the Hygr fragment (EcoT22I-BamHI) into the PstI-BamHI sites of pBSK-pgk-1.

(1-3) Construction of the KO2 vector

Fucose transporter targeting vector ver. 2 (referred to hereafter as the KO2 vector) was constructed by inserting the 5' side of the fucose transporter (from the SmaI site at No. 2,780 of SEQ ID NO: 16 to the BamHI at No. 4,232), the 3' side (from No. 4,284 to the SacI site at No. 10,934), and the pgk-1-Hygr fragment into the pMC1DT-A vector. Unlike the case for the KO1 vector, the KO2 vector has the pgk-1 gene promoter attached to Hygr, thus will acquire hygromycin B resistance even when only one copy of the vector is introduced into the cell by homologous recombination. The KO2 vector was cleaved at NotI and was transfected into the cell. by introducing the KO2 vector, 46 base pairs in exon 1 including the start codon will be deleted from the fucose transporter, resulting in a loss of function.

(1-4) Construction of pBSK-pgk-1-Puror

The pPUR vector (3D Biosciences) was cleaved by PstI and BamHI and the excised fragment (Puror) was inserted into the PstI-BamHI sites of pBSK-pgk-1 to construct pBSK-pgk-1-Puror.
(1-5) Construction of the KO3 vector

Fucose transporter targeting vector ver. 3 (referred to hereafter as the KO3 vector) was constructed by inserting the 5’ side of the fucose transporter (from the SmaI site at No. 2,780 of SEQ ID NO: 16 to the BamHI at No. 4,232), the 3’ side (from No. 4,284 to the SacI site at No. 10,934), and the pgk-1-Puror fragment into the pMC1DT-A vector. A sequence for binding to the screening primer shown below was preliminarily attached to the 3’ end of pgk-1-Puror. The KO3 vector was cleaved by Not1 and was transfected into the cell. By introducing the KO3 vector, 46 base pairs in exon 1 including the start codon will be deleted from the fucose transporter, resulting in a loss of function.

reverse primer

RSGR-A 5’-GCTGTCTGGAGTACTGTGCATCTGC-3’ (SEQ ID NO: 19)

The fucose transporter gene was knocked out using the three targeting vectors described above.

(2) Introduction of vector into CHO cells

HT Supplement (100x) (Invitrogen) and penicillin-streptomycin (Invitrogen) were added to CHO-S-SFMII HT- (Invitrogen), each at 1/100th the volume of the CHO-S-SFMII HT- to prepare a culture medium (hereafter referred to as SFMII(+)). CHO DXB11 cells were subcultured in SFMII(+). 8 x 10^6 CHO cells were suspended in 0.8 mL Dulbecco’s phosphate buffer (Invitrogen) (referred to hereafter as PBS). 30 μg of
targeting vector was added to the cell suspension, and transferred to a Gene Pulser cuvette (4 mm) (BioRad). After holding for 10 minutes on ice, the vector was transfected into the cells by electroporation with GENE PULSER II (BioRad) at 1.5 kV and 25 μFD. After vector transfection, the cells were suspended in 200 mL SFMII(+) medium and were seeded into twenty 96-well flat-bottom plates (Iwaki) at 100μl/well. The plates were cultured for 24 hours at 37°C in a CO₂ incubator, then the reagent was added.

(3) Knockout: step 1

KO1 vector or KO2 vector was transfected into the CHO cells, and selected with hygromycin B (Invitrogen) 24 hours after vector transfection. The hygromycin B was dissolved in SFMII(+) at 0.3 mg/mL, and 100 μL was added per well.

(4) Screening for homologous recombinants by PCR

(4-1) Preparation of the PCR sample

Homologous recombinants were screened by PCR. The CHO cells were cultured on 96-well flat-bottom plates, the culture supernatant was removed and a cell lysis buffer was added at 50 μL/well, and incubated at 55°C for 2 hours. Then proteinase K was deactivated by heating at 95°C for 15 minutes to prepare a PCR template. The cell lysis buffer was composed of the following per well: 5 μL 10x LA Buffer II (supplied with Takara LA Taq), 2.5 μL 10% NP-40 (Roche), 4 μL proteinase K (20 mg/mL, Takara), and 38.5 μL distilled water (Nacalai
(4-2) PCR conditions

The PCR reaction mixture was comprised of 1 µL of the PCR sample as described above, 5 µL 10X LA Buffer II, 5 µL MgCl₂ (25 mM), 5 µL dNTP (2.5 mM), 2 µL primer (10 µM each), 0.5 µL LA Taq (5 IU/µL), and 29.5 µL distilled water (total of 50 µL). The TP-F4 and Thygro-R1 primers were used as the PCR primers to screen the KO1 vector-transfected cells, and the TP-F4 and Thygro-F1 primers were used as the PCR primers to screen the KO2 vector-transfected cells.

PCR screen for the KO1 vector-transfected cells was carried out by preheating for 1 minute at 95°C; 40 cycles of 95°C/30 seconds, 60°C/30 seconds, and 60°C/2 minutes; and 7 minutes at 72°C. PCR screening for the KO2 vector-transfected cells was carried out by preheating for 1 minute at 95°C; 40 cycles of 95°C/30 seconds and 70°C/3 minutes; and 7 minutes at 70°C.

The primers are given below. In cell samples where homologous recombination had occurred, approximately 1.6 kb DNA is amplified for the KO1 vector and approximately 2.0 kb DNA is amplified for the KO2 vector. TP-F4 primer was designed in the genomic region of the fucosе transporter on the 5' side outside the vector, and the Thygro-F1 and Thygro-R1 primers were designed in the Hygr within the vector.

forward primer (KO1, KO2)
TP-F4 5'-GGAATGCGCTTCTTCAAGGACTCGC-3' (SEQ ID NO: 20)
reverse primer (K01)

Thygro-R1 5’-TGCATCAGGTGGAGACGGCTGGAAC-3’ (SEQ ID NO: 21)
reverse primer (K02)

Thygro-F1 5’-GCACTCGTCCGAGGCAAGAATAGC-3’ (SEQ ID NO: 22)

(5) Results of the PCR screening

918 K01 vector-transfected cells were analyzed, and 1 cell was thought to be a homologous recombinant (homologous recombination efficiency = approximately 0.1%). 537 K02 vector-transfected cells were analyzed, and 17 cells were thought to be homologous recombinants (homologous recombination efficiency = approximately 3.2%).

(6) Southern blotting analysis

Further confirmation was carried out by Southern blotting. 10 μg genomic DNA was prepared from the cultured cells by a standard method and was submitted to Southern blotting. Using the two primers given below, a 387 bp probe was prepared by PCR spanning the region from No. 2,113 to No. 2,500 in the nucleotide sequence shown in SEQ ID NO: 16, and used for Southern blotting. The genomic DNA was cleaved with BglIII.

forward primer

Bgl-F 5’-TGTGCTGGGAATTGAACCCCAGGAC-3’ (SEQ ID NO: 23)
reverse primer

Bgl-R 5’-CTACTTGCTGCTTTTCTTCC-3’ (SEQ ID NO: 24)

Cleavage by BglIII provides an approximately 3.0 kb band
from the fucose transporter chromosome, an approximately 4.6 kb band from the chromosome where homologous recombination occurred with the KO1 vector, and an approximately 5.0 kb band from the chromosome where homologous recombination occurred with the KO2 vector. One cell obtained by homologous recombination with the KO1 vector and 7 cells obtained by homologous recombination with the KO2 vector were used in the experiments. The only cell obtained with the KO1 vector was designated 5C1; however, it was shown by subsequent analysis to comprise multiple cell populations. Thus cloning by limit dilution was carried out before using it in the subsequent experiments. One of the cells obtained with the KO2 vector was designated 6E2.

(7) Knockout: step 2

In order to establish cell lines having a completely defective fucose transporter gene, one of three vectors was introduced into the cells in which homologous recombination had been achieved by the KO1 vector or KO2 vector. The vector/cell combinations were as follows: method 1: KO2 vector and 5C1 cells (KO1); method 2: KO2 vector and 6E2 cells (KO2); method 3: KO3 vector and 6E2 cells (KO2). The vector was transfected into the respective cell, and selection with hygromycin B and puromycin (Nacalai Tesque) was begun 24 hours after vector transfection. The final hygromycin B concentration was 1 mg/mL in method 1 and 7 mg/mL in method 2. In method 3, the hygromycin B was added at a final
concentration of 0.15 mg/mL and the puromycin was added at a final concentration of 8 μg/mL.

(8) PCR screening of the homologous recombinants

The samples were prepared as described above. In the screening for method 1, both cells in which homologous recombination had occurred with the KO1 vector and cells in which homologous recombination had occurred with the KO2 vector were detected by means of PCR. The PCR primers shown below were designed for method 2. TPS-F1 corresponds to the region from No. 3,924 to No. 3,950 in the nucleotide sequence shown in SEQ ID NO: 16, while SHygro-R1 corresponds to the region from No. 4,248 to 4,274. These PCR primers will amplify 350 bp of the fucose transporter gene region that is otherwise deleted by the KO2 vector. Therefore, in the PCR screening in method 2, cells producing no 350 bp amplification product are considered to be the cells completely lacking the fucos transporter gene. The PCR conditions were as follows: preheating for 1 minute at 95°C; 35 amplification cycles of 95°C/30 seconds and 70°C/1 minute; and reheating at 70°C for 7 minutes.

forward primer
TPS-F1 5'-CTCGACTCGTCCCTATTAGGCAACAGC-3' (SEQ ID NO: 25)

reverse primer
SHygro-R1 5'-TCAGAGGCAGTGGAGCCTCCAGTCAGC-3' (SEQ ID NO: 26)

In the case of method 3, TP-F4 was used as the forward
primer and RSGR-A was used as the reverse primer. The PCR conditions were as follows: preheating for 1 minute at 95°C; 35 amplification cycles of 95°C/30 seconds, 60°C/30 seconds, and 72°C/2 minutes; reheating for 7 minutes at 72°C. Approximately 1.6 kb DNA will be amplified from the cell samples in which homologous recombination by the KO3 vector has occurred. By this PCR, the cells in which homologous recombination occurred by the KO3 vector were detected, and it was also confirmed that the cells retained the homologous recombination by the KO2 vector while detecting

(9) Results of PCR screening

18 cells out of 616 cells analyzed by method 1 were considered to be homologous recombinants (homologous recombination efficiency = 2.9%). 2 cells out of 524 cells analyzed by method 2 were considered to be homologous recombinants (homologous recombination efficiency = approximately 0.4%). 7 cells out of 382 cells analyzed by method 3 were considered to be homologous recombinants (homologous recombination efficiency = approximately 1.8%).

(10) Southern blotting analysis

Southern blotting analysis was carried out by the method described above, and 1 cell was identified to have the fucose transporter gene being completely lost. The analytical results from PCR and Southern blotting were consistent in the knockout step 1, but not in the knockout step 2.

(11) Fucose expression analysis
26 cells found to be homologous recombinants were analyzed for fucose expression by PCR. 1 x 10^6 cells were stained in 100 μL PBS containing 5 μg/mL Lens culinaris agglutinin, FITC conjugate (Vector Laboratory), 2.5% FBS, and 0.02% sodium azide (referred to hereafter as FACS solution) for 1 hour with ice cooling. The cells were then washed 3 times with FACS solution and were analyzed by FACSCalibur (Becton Dickinson). The results showed that fucose expression was reduced only in FTP-KO cells, which were found to exhibit a complete loss of the fucose transporter gene in the Southern blot analysis.

Reference Example 3
Establishment of antibody-producing cells derived from the FTP-KO line and purification of antibody produced by these cells

The fucose transporter-deficient line obtained in Example 1 (FTP-KO cells, clone name: 3F2) were subcultured in SFMII(+) containing hygromycin B at a final concentration of 1 mg/mL. 8 x 10^6 3F2 cells were suspended in 0.8 mL Dulbecco’s phosphate buffer. 25 μg of a humanized anti-glypican 3 antibody expression vector was added to the cell suspension and transferred to a Gene Pulser cuvetto. After holding for 10 minutes on ice, the vector was transfected into the cells by electroporation with GENE PULSER II at 1.5 kV and 25 μFD. After vector transfection, the cells were suspended in 40 mL
SFMII(+) medium and were seeded on a 96-well flat-bottom plate (Iwaki) at 100 μL/well. After the plate was incubated for 24 hours at 37°C in a CO₂ incubator, geneticin (Invitrogen) was added to a final concentration of 0.5 mg/mL. The antibody production of the antibody-resistant cells was measured to obtain humanized anti-glypican 3 antibody-producing cell lines.

The culture supernatant from the antibody-expressing lines was collected and applied to Hitrap rProtein A column (Pharmacia) using P-1 pump (Pharmacia). The column was washed with a binding buffer (20 mM sodium phosphate (pH 7.0)), and the bound antibody was subsequently eluted with elution buffer (0.1 M glycine-HCl (pH 2.7)). The eluate was immediately neutralized with neutralization buffer (1 M Tris-HCl (pH 9.0)). The eluted antibody fractions were selected using DC protein assay (BioRad) and pooled, and was concentrated to about 2 mL with Centriprep-YM10 (Millipore). The concentrated solution was then submitted to gel filtration on Superdex200 26/60 (Pharmacia) equilibrated with 20 mM acetic acid buffer (pH 6.0) containing 150 mM NaCl. The peak of the monomer fraction in the eluate was collected and concentrated with Centriprep-YM10. The concentrated solution was filtered using a MILLEX-GW 0.22 μm filter unit (Millipore) and was stored at 4°C. The purified antibody concentration was determined by calculation from the molar absorption coefficient based on the absorbance at 280 nm.
Reference Example 4

Analysis of sugar chains attached to the humanized anti-glypican 3 antibody produced by FTP-KO cells

(1) Preparation of 2-aminobenzamide-labeled sugar chains (2-AB-labeled sugar chains)

The antibody of the present invention produced by FTP-KO cells and antibody produced by CHO cells (control) were treated with N-glycosidase F (Roche Diagnostics) to release the antibody-bound sugar chains (Weitzhandler M. et al., Journal of Pharmaceutical Sciences (1994) 83(12), 1670-5). After deproteination with ethanol (Schenk B. et al., The Journal of Clinical Investigation (2001), 108(11), 1687-95), the released sugar chains were concentrated to dryness and fluorescent-labeled with 2-aminopyridine (Bigge J. C. et al., Analytical Biochemistry (1995) 230(2), 229-238). The resulting 2-AB-labeled sugar chains were separated from the reagent by solid-phase extraction using a cellulose cartridge, concentrated by centrifugation to obtain purified 2-AB-labeled sugar chains. The purified 2-AB-labeled sugar chains was treated with β-galactosidase (Seikagaku Kogyo Co., Ltd.) to prepare agalactosyl 2-AB-labeled sugar chains which were analysed as described below.

(2) Analysis of agalactosyl 2-AB-labeled sugar chains by normal-phase HPLC

The agalactosyl 2-AB-labeled sugar chains prepared by the above-described method, the sugar chains released from the
antibody of the present invention produced by FTP-KO cells and antibody produced by CHO cells (control), were analyzed by normal-phase HPLC by TSKgel Amide-80 amide column (TOSOH Corp.) and the chromatograms were compared. The following assessment was made with respect to the antibody produced by the CHO cells: G(0) was present as the main component of the sugar chains, while G(0)-Fuc lacking fucose accounted for about 4% of the total sugar chains, as assessed by calculation from the peak area ratio. In the case of the antibody produced by FTP-KO cells, G(0)-Fuc was the main component. For all the antibody producing cell lines, at least 90% of the total sugar chain in the produced antibody was fucose-free sugar chains, as assessed by on calculation from the peak area ratio.

Table 1.

Relative ratios of each sugar chain estimated from normal-phase HPLC analysis of agalactosyl 2-AB-labeled sugar chains

<table>
<thead>
<tr>
<th>sugar chain</th>
<th>CHO</th>
<th>FTP-KO-a</th>
<th>FTP-KO-b</th>
<th>FTP-KO-c</th>
</tr>
</thead>
<tbody>
<tr>
<td>G(0)-Fuc</td>
<td>4.0%</td>
<td>92.4%</td>
<td>92.5%</td>
<td>93.2%</td>
</tr>
<tr>
<td>G(0)</td>
<td>96.0%</td>
<td>7.6%</td>
<td>7.5%</td>
<td>6.8%</td>
</tr>
</tbody>
</table>

Example 5

Establishment of cell lines stably expressing humanized HOLO antibody and point mutation-modified antibodies

The genes encoding Hspu2.2Lspu2.2 (Hu2.2Lu2.2) antibody, Hspd1.8Lspd1.6 (Hd1.8Ld1.6) antibody (HOLO antibody-sourced modified antibodies produced by the method described in
Example 1) or the HOLO antibody submitted to the modifications were cloned into an expression vector. In the cloning, the genes encoding the heavy chain and light chain were inserted into different expression vectors in order to express each gene encoding the heavy chain and light chain of the antibody. Two expression vectors were selected so as to provide a desired combination of the genes encoding the heavy chain and light chain as described above, digested with PvuI and transfected by electroporation into the FTP-KO line produced in Reference Example 2.

Transformants that stably produce HOLO antibody or modified antibody were prepared by electroporation using the Gene Pulser II (BioRad). 0.75 mL CHO cells (1 x 10^7 cells/mL) suspended in PBS was mixed with 10 μg each of expression plasmid DNA providing the desired combination of heavy chain and light chain, and the mixture was left stand on ice for 10 minutes. The mixture was transferred to a Gene Pulser II cuvette and a 1.5 kV electrical pulse was applied at a capacitance of 25 μFD. The pulsed mixture was incubated for 10 minutes at room temperature and suspended in CHO-S-SFMII/1% HT/1% PS medium. 100 μL of the diluted suspension (5 fold, 10 fold, and 50 fold, made in the same medium) was added to each well of 96-well culture plates. The plates were incubated for 24 hours in CO₂ incubator maintained at 5% CO₂. Then, geneticin (Gibco) was added to each well to a final concentration of 500
μg/mL and zeocin (Invitrogen) was added to each well to a final concentration of 600 μg/mL. The plates were incubated for 2 weeks. The colonies of transformed cells that exhibited geneticin and zeocin resistance were further selected by subculture on the same medium containing 500 μg/mL geneticin (Gibco) and 600 μg/mL zeocin (Invitrogen). The culture supernatant of the transformed cells selected in this manner was evaluated for the antibody concentration using BiacoreQ (BIACORE) to establish transformant cell lines expressing the desired antibody at a high level. The antibody concentration in the culture supernatant was measured according to the instructions provided with the BiacoreQ (BIACORE). The cell line thus established was cultured in CHO-S-SFMII medium (Invitrogen) containing 500 μg/ml of Geneticin (Invitrogen) and 600 μg/ml of Zeocin (Invitrogen). After cultivating for an appropriate period of time, the culture supernatant was collected, and purified using rProteinA-Sepharose column (GE Healthcare). The purified antibody was concentrated with Amicon Ultra-4 (MILLIPORE) and subjected to buffer exchange on PD-10 Desalting column (Amersham Biosciences) with 20mM acetic acid buffer (pH6.0) containing 200mM NaCl. The purified antibody was quantified by the absorption at 280nm in ND-1000 Spectrophotometer (NanoDrop) or DU-600 Spectrophotometer (BECKMAN). The antibody concentration was calculated with the RACE method.
Example 6

Therapeutic efficacy in an in vivo model of the humanized HOLO antibody and the point mutation-modified antibodies

(1) Maintenance of the cell line used for transplantation into the in vivo model

HepG2 cells (ATCC) were used. The HepG2 cells were maintained by subculture in Eagle’s minimum essential medium (Sigma) containing sodium pyruvate (Invitrogen) at 1 mmol/L MEM and non-essential amino acids (Invitrogen) at 1 mmol/L MEM (this medium is referred to as the subculture medium).

(2) Preparation of an HepG2 cell-transplanted mouse model

Using a solution containing subculture medium and MATRIGEL Matrix (BD Bioscience) at 1 : 1, a suspension of HepG2 cells was prepared at 5 x 10^7 cells/mL. 100 µL of the cell suspension (5 x 10^7 cells/mouse) was transplanted subcutaneously in the abdominal region of SCID mice (male, 5 week old, CLEA Japan, Inc.). On the day prior to cell transplantation, the mice received 100 µL anti-asialo GM1 antibody (content of 1 vial was dissolved in 5 mL of the aforementioned solution, Wako Pure Chemical Industries, Ltd.) by intraabdominal administration. The tumor volume was calculated according to the following formula.

\[ \text{tumor volume} = \text{long diameter} \times \text{short diameter} \times \text{short diameter}/2 \]

When the average tumor volume reached 130 to 330 mm^3, the
mouse was used as a model.

(3) Preparation of administration sample containing the test antibody

The administration samples containing H0L0 antibody, Hu2.2L2.2 antibody, or H2l.8Ld1.6 antibody at 0.5 mg/mL (5 mg/kg group) or 0.1 mg/mL (1 mg/kg group) were prepared with physiological saline on the day of administration.

(4) Administration of the antibody-containing administration sample

The administration sample prepared according to (3) was administered at a dose of 10 mL/kg through the tail vein to the mouse model prepared in (2) at once per week for three weeks beginning 27 days after HepG2 cell transplantation. As a negative control, physiological saline was administered at a dose of 10 mL/kg through the tail vein at once per week for three weeks. All the groups contained 5 animals, and each test antibody-containing administration sample was administered to a respective group. At about the same time as the administration, venous blood was collected from 3 animals in each group was analyzed for the murine blood concentration of the antibody. Specifically, blood was collected from a dorsal foot vein at two time points: 0.5 hour after the initial administration and immediately before the second administration. 20 μL of collected blood was treated with heparin and the plasma was isolated by centrifugation.

(5) Evaluation of the antitumor activity of the test
antibodies

The antitumor activity of each test antibody was evaluated in the mouse model transplanted with human liver cancer. The tumor volume was measured at one week after the last day of the sample administration. The results are shown in Figure 7. Hspd1.8Lspd1.6 (Hd1.8Ld1.6) antibody showed a stronger therapeutic efficacy, while the Hspu2.2Lspu2.2 (Hu2.2Lu2.2) antibody showed a weak therapeutic efficacy.

(6) Blood concentrations of the test antibodies

The concentration of the test antibody in murine plasma was measured by an ELISA-based method described in Example 1. Standard samples were prepared at the plasma concentration of 12.8, 6.4, 3.2, 1.6, 0.8, 0.4, and 0.2 μg/mL. The standard samples and the murine plasma test samples (suitably diluted to a desired concentration) were added to immunoplates (Nunc-ImmuNoPlate, MaxiSorp (Nalge Nunc International)) on which soluble glypican 3 core (Chugai Seiyaku Kabushiki Kaisha) is immobilized, and the plates were incubated for 1 hour at room temperature. Goat Anti-Human IgG-BIOT (Southern Biotechnology Associates) and then streptavidin-alkaline phosphatase conjugate (Roche Diagnostics) were added and a chromogenic reaction was effected using the BluePhos Microwell Phosphatase Substrates System (Kirkgaard & Perry Laboratories) as a substrate. Using a microplate reader, the color of the reaction solution in each well was determined by measuring the absorbance of the reaction solution at 650 nm. The murine
plasma antibody concentration was then calculated using SOFTmax PRO analytical software (Molecular Devices) with reference to the standard curve prepared with the absorbance values obtained from the standard samples.

The murine plasma concentrations after 30 minutes and 7 days from administration are shown in Figure 8. A higher antibody concentration was observed in murine plasma after 7 days from administration of the test antibody having lower pI values for both administration doses tested.

Example 7

ADCC activity of the test antibodies measured using human peripheral blood monocytes as the effector cell

The ADCC activity of the test antibodies was tested as described below using a human peripheral blood monocyte (referred to as PBMC) as the effector cell).

(1) Preparation of a human PBMC solution

Using a syringe previously loaded with 200 μL of a 1000 unit/mL heparin solution (Novo Heparin Injection 5000 Units, Novo Nordisk), 50 mL peripheral blood was collected from a healthy volunteer (adult male) from Chugai Seiyaku Kabushiki Kaisha. The blood was diluted twofold with PBS(-), divided into 4 equal parts, and introduced into a Leucosep lymphocyte separation tube (Greiner Bio-one) that was previously loaded with 15 mL Ficoll-Paque PLUS and subjected to centrifugation. The separation tube loaded with the peripheral blood was centrifuged for 10 minutes at room temperature at 2150 rpm.
and the monocyte fraction layer was collected. The cells contained in the layer were washed once with Dulbecco's Modified Eagle's Medium (Sigma) containing 10% FBS (referred to below as 10% FBS/D-MEM) and suspended in 10% FBS/D-MEM at a cell density of 5 x 10⁶/mL. The cell suspension was submitted to the following experiment as a human PBMC solution.

(2) Preparation of target cells

HepG2 cells were detached from a dish and seeded on a 96-well U-bottom plate at 1 x 10⁴ cells/well. The plate was incubated overnight at 37°C in a 5% CO₂ incubator. On the next day, 5.55 MBq Cr-51 was added to each well of the plate and the plate was incubated for 3 hours at 37°C in a 5% CO₂ incubator. The HepG2 cells contained in the wells of the plate were used as target cells in the ADCC activity assay as described below.

(3) Chromium release assay (ADCC activity)

The ADCC activity was evaluated from the specific chromium release rate determined by the chromium release method. The target cells prepared as in (2) were washed with the medium, and 100 μL of each antibody (HOL0 antibody, Hu2.2Lu2.2 antibody, or Hd1.8Id1.6 antibody) was added at a concentration of 0, 0.004, 0.04, 0.4, 4, or 40 μg/mL. The plate was reacted for 15 minutes at room temperature, and the antibody solution was removed. 100 μL of subculture medium was added to each well and the plate was incubated for 1 hour at
37°C in a 5% CO₂ incubator. 100 μL of the human PBMC solution prepared as in (1) was added to each well (5 x 10⁵ cells/well), and the plate was incubated for 4 hours at 37°C in a 5% CO₂ incubator and centrifuged. The radioactivity in 100 μL of the culture supernatant in each well of the plate was measured by a gamma counter. The specific chromium release rate was determined according to the following formula.

\[
\text{Specific chromium release rate (\%) = } \frac{(A - C) \times 100}{(B - C)}
\]

wherein A represents the mean value of the radioactivity (cpm) of the 100 μL culture supernatant in each well; B represents the mean value of the radioactivity (cpm) of the 100 μL culture supernatant in wells where 100 μL of 2% aqueous NP-40 solution (Nonidet P-40, Nacalai Tesque) and 50 μL of 10% FBS/D-MEM medium were added to the target cells; and C represents the mean value of the radioactivity (cpm) of the 100 μL culture supernatant in wells where 150 μL of 10% FBS/D-MEM medium was added to the target cells. The tests were carried out in triplicate, and the mean value and standard deviation of the specific chromium release rate (\%), which is reflective of the ADCC activity of the antibody, were calculated form the test results.

(4) Evaluation of the ADCC activity of the test antibodies

The results of the evaluation of the ADCC activity exhibited by human PBMCs via the test antibody revealed that
all of the test antibodies exhibited ADCC activity. The results are shown in Figure 9. Significance test revealed that no significant difference was observed between test antibodies at any concentrations tested in the specific chromium release rate of the test antibodies. The statistical analyses were run using the SAS Preclinical Package (SAS Institute Inc.). These results showed that there was no difference between the ADCC activities of the pI-modified test antibodies.

Example 8
Preparation and characterization of pI modified antibody by point mutation

(1) Selection of modification sites for decreasing pI

To improve the tumor suppression activity of the Hdl.8Ld1.6 antibody, modification sites were selected for the ability of decreasing in the pI value of the variable region. Amino acid residues involving the decrease in the pI value of the variable region were found, which are summarized in Table 2 (heavy chain) and Table 3 (light chain). Specific examples of these modifications for decreasing the pI value are pH7pL14 antibody and pH7pL16 antibody. These pI modification antibodies were prepared as follows.

The modification sites were created by Assemble PCR. Oligo DNAs designed based on the sense and antisense sequences containing the modification site were synthesized. A pair of an antisense oligo DNA containing the modification site and a sense oligo DNA corresponding to the vector bearing the gene
to be modified, or a pair of a sense oligo DNA containing the modification site and an antisense oligo DNA corresponding to the vector bearing the gene to be modified was used in PCR with PrimeSTAR (TAKARA) to obtain 5'-side and 3'-side fragments containing the modification site. The two fragments were linked using Assemble PCR to prepare each mutant.

The mutant thus obtained was inserted into an expression vector which allows for expression of the inserted gene in animal cells. The nucleotide sequence of the expression vector was determined by a method known in the art. Introduction of the point mutation was confirmed by the nucleotide sequence of the plasmid DNA. The gene containing the point mutation was cloned into an expression vector which allows for expression of the inserted gene in animal cells. The expression and purification of the antibody was according to the method described in Example 1 or a similar method.

Starting from Hdl.8, the 61st glutamine (Q) (according to the Kabat numbering) present in CDR1 of Hdl.8 was substituted with glutamic acid (E) to prepare pH7 (SEQ ID NO:27). Starting from Ldl.6, the 24th arginine (R) (according to the Kabat numbering) present in CDR1 of Ldl.6 was substituted with glutamine (Q), the 37th glutamine (Q) was substituted with leucine (L), the 43rd alanine (A) was substituted with serine (S), the 45th arginine (R) was substituted with glutamine (Q), the 74th threonine (T) was substituted with lysine (K), the 77th serine (S) was substituted with arginine (R), the 78th
leucine (L) was substituted with valine (V), and the 79th glutamine (Q) was substituted with glutamic acid (E), each present in FR2 and FR3, to prepare pl14 (SEQ ID NO:28).

Starting from pl14, the 104th leucine (L) (according to the Kabat numbering) was substituted with valine (V), the 107th lysine (K) was substituted with glutamic acid (E), each present in FR4 of pl14, to prepare pl16 (SEQ ID NO:29).

(2) Measurement of pI value of point mutation pI modified antibodies

The pI values of the Hd1.8Ld1.6 antibody, pH7pL14 antibody and pH7pL16 antibody were measured by electrophoresis with PhastGel IEF 4-6.5 (GE Healthcare) using the method described in Example 1 or similar method. The pI value of Hd1.8Ld1.6 antibody, pH7pL14 antibody and pH7pL16 antibody was 7.47, 7.07 and 6.52, respectively, indicating that the pI values of pH7pL14 antibody and pH7pL16 antibody were lower than that of the Hd1.8Ld1.6 antibody by 0.4 and 0.95, respectively.

(3) Measurement of Tm value of point mutation pI modified antibodies

The Tm values of Fab's obtained from Hd1.8Ld1.6 antibody, pH7pL14 antibody and pH7pL16 antibody were measured with VP-DSC (Micro Cal) using the method similar to Example 1. In this experiment, PBS was used as a dialysis solution, and the antibody concentration in the test solution for DSC measurement was adjusted to 25-100 µg/ml. DSC scanning was set
from 20°C to 115 °C at the scanning rate of about 4K/min, with
the reference solution (dialysis solution) and DSC measurement
test solution. The thermal denaturation midpoint temperature
of the Fabs of the Hdl.8Ld1.6 antibody, pH7pL14 antibody and
pH7pL16 antibody was 77.5, 78.0 and 74.7 °C, respectively.
(4) Evaluation of binding activity to antigen of point
mutation pI modified antibodies by competitive ELISA

The binding activity to the antigen glypican 3 of each
point mutation pI modified antibody was measured using the
method described in Example 1 (Figure 10). The binding
activity to glypican 3 of the pH7pH14 antibody and pH7pL16
antibody was shown to be comparative to that of the H0L0
antibody.

Example 9
Preparation of point mutation pI modified antibody using FTP-
KO cell line

The expression vector carrying the gene coding for each
point mutation pI modified antibody prepared in Example 8 was
introduced into the cells of FTP-KO cell line prepared in
Reference Example 2 using Polyethylenimine (Polysciences Inc.),
and the antibody was expressed. The modified antibody was
purified from the cell culture supernatant using rProtein A
Sepharose™ Fast Flow (Amersham Biosciences). The purified
antibody solution was prepared according to the method
described in Example 5 and the antibody concentration was
measured.
Example 10

Therapeutic efficacy in an in vivo model of humanized GC33 antibody and point mutation p1 modified antibodies

(1) Maintenance of the cell line used for transplantation into the in vivo model

Hep G2 cells (ATCC) were used. The Hep G2 cells were maintained by subculture in Minimum Essential Medium Eagle medium (SIGMA) supplemented with 10% FBS, 1 mmol/l MEM Sodium Pyruvate (Invitrogen), and 1 mmol/l MEM non-essential amino acids (Invitrogen) (referred to as the subculture medium).

(2) Preparation of HepG2 cell-transplanted mouse model

Using a solution containing subculture medium and MATRIGEL Matrix (BD Bioscience) at 1 : 1, a suspension of HepG2 cells was prepared at $5 \times 10^7$ cells/mL. 100 µL of the cell suspension ($5 \times 10^6$ cells/mouse) was transplanted subcutaneously in the abdominal region of SCID mice (male, 5 week old, CLEA Japan, Inc.). On the day prior to cell transplantation, the mice received 100 µL anti-asialo GM1 antibody (content of 1 vial was dissolved in 5 mL of the solution, Wako Pure Chemical Industries, Ltd.) by intraabdominal administration. The tumor volume was calculated according to the following formula.

tumor volume = long diameter x short diameter x short diameter/2

When the average tumor volume reached 400 mm$^3$, the mouse was used as a model.
(3) Preparation of administration sample containing the test antibody

The administration samples containing H0L0 antibody, Hd1.8Ld1.6 antibody, pH7pL14 antibody or pH7pL16 antibody at 0.1 mg/mL (1 mg/kg group) were prepared with physiological saline on the day of administration.

(4) Administration of the antibody-containing administration sample

The administration sample prepared according to (3) was administered at a dose of 10 mL/kg through the tail vein to the mouse model prepared in (2) at once per week for five weeks beginning 34 days after HepG2 cell transplantation. As a negative control, physiological saline was administered at a dose of 10 mL/kg through the tail vein at once per week for five weeks. All the groups contained 5 animals, and a each test antibody-containing administration sample was administered to a respective group. At about the same time as the administration, venous blood was collected from 3 animals in each group and was analyzed for the murine blood concentration of the antibody. Specifically, blood was collected from a dorsal foot vein at two time points: 0.5 hour after the initial administration and immediately before the second administration. 20 µL collected blood was treated with heparin and the plasma was isolated by centrifugation.

(5) Evaluation of the antitumor activity of the test antibodies
The antitumor activity of each test antibody was evaluated in the mouse model transplanted with human liver cancer. The tumor volume was measured at one week after the last day of the sample administration. As shown in Figure 11, a stronger therapeutic efficacy was found in pH7pL14 antibody and pH7pL16 antibody compared to H0L0 antibody and Hd1.8Ld1.6 antibody.

Example 11
The PK test of humanized GC33 antibody and point mutation antibodies using in vivo model

(1) Preparation of test antibody-containing administration sample

The administration samples containing H0L0 antibody, Hd1.8Ld1.6 antibody, pH7pL14 antibody, pH7pL16 antibody or pH7M85pL16 antibody at 0.5 mg/mL (5 mg/kg group) were prepared with physiological saline on the day of administration.

(2) Administration of the antibody-containing administration sample

The administration sample prepared according to (1) was administered at a dose of 10 mL/kg through the tail vein to the C.B-17/Icr-scid mouse. All the groups contained 3 animals, and each test antibody-containing administration sample was administered to a respective group. Venous blood was collected from the animals and was analyzed for the murine blood concentration of the antibody. Specifically, blood was collected from a dorsal foot vein at seven time points: 0.5
hours, 2 hours, 8 hours, 24 hours, 72 hours, 168 hours after
the initial administration. 20 μL of collected blood was
treated with heparin and the plasma was isolated by
centrifugation.

(3) Blood concentrations of the test antibodies

The concentration of the test antibody in murine plasma
was measured by an ELISA-based method described in Example 6.
Standard samples were prepared at a plasma concentration of
12.8, 6.4, 3.2, 1.6, 0.8, 0.4, and 0.2 μg/mL. The standard
samples and the murine plasma test samples (suitably diluted
to a desired concentration) were added to immunoplates (Nunc-
ImmuNoPlate, MaxiSorp (Nalge Nunc International)) on which
soluble glypican 3 core (Chugai Seiyaku Kabushiki Kaisha) is
immobilized, and the plates were incubated for 1 hour at room
temperature. Goat Anti-Human IgG-BIOT (Southern Biotechnology
Associates) and then streptavidin-alkaline phosphatase
conjugate (Roche Diagnostics) were added, and a chromogenic
reaction was effected using the BluePhos Microwell Phosphatase
Substrates System (Kirkgaard & Perry Laboratories) as a
substrate. Using a microplate reader, the color of the
reaction solution in each well was determined by measuring the
absorbance of the reaction solution at 650 nm. The murine
plasma antibody concentration was then calculated using
SOFTmax PRO analytical software (Molecular Devices) with
reference to the standard curve prepared with the absorbance
values obtained from the standard samples.
The murine plasma concentrations after administration are shown in Figure 12. A higher antibody concentration was observed in the plasma of mice that received the test antibody having lower pI value.

Example 12

ADCC activity of the test antibodies using human peripheral blood monocytes as the effector cell

The ADCC activity of the test antibodies was tested as described below using a human peripheral blood monocyte (referred to as PBMC) as the effector cell.

1. Preparation of a human PBMC solution

Using a syringe previously loaded with 200 µL of a 1000 unit/mL heparin solution (Novo Heparin Injection 5000 Units, Novo Nordisk), 50 mL peripheral blood was collected from a healthy volunteer (adult male) from Chugai Seiyaku Kabushiki Kaisha. The blood was diluted twofold with PBS(-), divided into 4 equal parts and introduced into a Leucosep lymphocyte separation tube (Greiner Bio-one) that was previously loaded with 15 mL Ficoll-Paque PLUS and subjected to centrifugation. The separation tube loaded with the peripheral blood was centrifuged for 10 minutes at room temperature at 2150 rpm, and the monocyte fraction layer was collected. The cells contained in the layer were washed once with Dulbecco’s Modified Eagle’s Medium (Sigma) containing 10% FBS (referred to below as 10% FBS/D-MEM) and suspended in 10% FBS/D-MEM at a
cell density of $5 \times 10^5$/mL. The cell suspension was submitted to the following experiment as a human PBMC solution.

(2) Preparation of target cells

HepG2 cells were detached from a dish and seeded on a 96-well U-bottom plate at $1 \times 10^4$ cells/well. The plate was incubated overnight at 37°C in a 5% CO2 incubator. On the next day, 5.55 MBq Cr-51 was added to each well of the plate and the plate was incubated for 3 hours at 37°C in a 5% CO2 incubator. The HepG2 cells contained in the wells of the plate were used as target cells in the ADCC activity assay as described below.

(3) Chromium release assay (ADCC activity)

The ADCC activity was evaluated from the specific chromium release rate determined by the chromium release method. The target cells prepared as in (2) were washed with medium, and 100 μL of each antibody (H0L0 antibody, Hdl.8Ldl.6 antibody, pH7pL14 antibody or pH7pL16 antibody) was added at a concentration of 0, 0.004, 0.04, 0.4, 4, or 40 μg/mL. The plate was reacted for 15 minutes at room temperature, and the antibody solution was removed. 100 μL of subculture medium was added to each well and the plate was incubated for 1 hour at 37°C in a 5% CO2 incubator. 100 μL of the human PBMC solution prepared as in (1) was added to each well ($5 \times 10^5$ cells/well) and the plate was incubated for 4 hours at 37°C in a 5% CO2 incubator, and centrifuged. The radioactivity in 100 μL of the culture supernatant in each well of the plate was measured by
a gamma counter. The specific chromium release rate was determined according to the following formula.

Specific chromium release rate (%) = \((A - C) \times 100/(B - C)\)

wherein A represents the mean value of the radioactivity (cpm) of the 100 μL culture supernatant in each well; B represents the mean value of the radioactivity (cpm) of the 100 μL culture supernatant in wells where 100 μL of 2% aqueous NP-40 solution (Nonidet P-40, Nacalai Tesque) and 50 μL of 10% FBS/D-MEM medium were added to the target cells; and C represents the mean value of the radioactivity (cpm) of the 100 μL culture supernatant in wells where 150 μL of 10% FBS/D-MEM medium was added to the target cells. The tests were carried out in triplicate, and the mean value and standard deviation of the specific chromium release rate (%), which is reflective of the ADCC activity of the antibody, were calculated from the test results.

(4) Evaluation of the ADCC activity of the test antibodies

The results of the evaluation of the ADCC activity exhibited by human PBMCs via the test antibody revealed that all of the test antibodies exhibited ADCC activity. The results are shown in Figure 13. Significance test revealed that no significant difference was observed at any concentrations tested between test antibodies and the control HOLO antibody in the specific chromium release rate of the antibodies. The statistical analyses were run using the SAS Preclinical Package (SAS Institute Inc.). These results showed
that there was no difference between the ADCC activities of
the pI-modified test antibodies.

Example 13
Preparation and evaluation of modified antibodies capable of
decreasing in the pI value of the constant region

(1) Selection of modification sites for decreasing the pI
value of the constant region

IgG1ΔGK (SEQ ID NO:32) is the IgG1 constant region having
the amino acid sequence as shown in SEQ ID NO:31, which lacks
the 446th Gly and the 447th Lys (according to the EU
numbering) of the IgG1 constant region. Deletion of these two
amino acid residues allows for the decrease in heterogeneity
cauesed by the heavy chain terminal constant region of an
antibody.

The antibody was modified to have decreased pI value in
the constant region by substituting some of the amino acid
residues in IgG1ΔGK with the corresponding amino acid residues
in the sequence of the human IgG4 constant region according to
the EU numbering.

Specifically, the 268th histidine (H) of IgG1ΔGK
(according to the EU numbering) was substituted with glutamine
(Q) of the IgG4 sequence, the 274th lysine (K) was substituted
with glutamine (Q), the 355th arginine (R) was substituted
with glutamine (Q), the 356th aspartic acid (D) was
substituted with glutamic acid (E), the 358th leucine (L) was
substituted with methionine (M), and the 419th glutamine (Q)
was substituted with glutamic acid (E). These substitutions only contain sequence of 9-12 amino acids derived from a human constant region which can serve as a T-cell epitope and thus are expected to have lower risk of immunogenicity. These 6 modifications were introduced into IgGlΔGK to obtain M85 (SEQ ID NO:33).

The constant region M85 was combined with the variable region pH7 and HO to prepare pH7M85 (SEQ ID NO:34) and H0M85 (SEQ ID NO:35), respectively. H0M85L0 antibody was prepared from H0M85 as the heavy chain and L0 as the light chain, and pH7M85pL16 antibody was prepared from pH7M85 as the heavy chain and pL16 as the light light chain. Also, the H0L0 antibody and pH7pL16 antibody having the constant region of IgGl were prepared as in Example 1 and Example 8. The antibodies H0M85L0, pH7M85pL16, H0L0, and pH7pL16 were expressed in FTP-KO cell lino or HEK293 cells, and purified as described in Example 1 or 9.

(2) Measurement of pI value of constant region pI modified antibodies

The pI value of the H0L0 antibody, H0M85L0 antibody, pH7pL16 antibody, and pH7M85pL16 antibody was measured by electrophoresis with PhastGel IEF 4-6.5 (GE Healthcare) under the same electrophoresis conditions using the method similar to that described in Example 1. The pI values of the H0L0 antibody, H0M85L0 antibody, pH7pL16 antibody and pH7M85pL16 antibody was 8.85, 8.16, 6.52 and 5.78, respectively.
indicating that the modification in the constant region contributes to further decrease in the pI value without affecting to the immunogenicity of the antibody.

(3) Evaluation of binding activity of constant region pI modified antibodies by competitive ELISA

The binding activity to the antigen of each constant region pI modified antibody was measured using the method described in Example 1 (Figure 14). The binding activity to glypican 3 of the H0L0 antibody, H0M85L0 antibody, pH7pL16 antibody, pH7M85pL16 antibody were shown to be almost the same.

Example 14

ADCC activity of the constant region pI modified antibodies using human peripheral blood monocytes as the effector cell

The ADCC activity of the pH7pL16 antibody and pH7M85pL16 antibody was tested using the method as described in Example 12. The results are shown in Figure 15. Significance test revealed that no significant difference was observed between test antibodies at any concentrations tested in the specific chromium release rate of the test antibodies. The statistical analyses were run using the SAS Preclinical Package (SAS Institute Inc.). These results showed that there was no difference between the ADCC activities of the pH7pL16 antibody and pH7M85pL16 antibody.
DEMANDE OU BREVET VOLUMINEUX

LA PRÉSENTE PARTIE DE CETTE DEMANDE OU CE BREVET COMPREND PLUS D'UN TOME.

CECI EST LE TOME 1 DE 2
CONTENANT LES PAGES 1 À 137

NOTE : Pour les tomes additionnels, veuillez contacter le Bureau canadien des brevets

JUMBO APPLICATIONS/PATENTS

THIS SECTION OF THE APPLICATION/PATENT CONTAINS MORE THAN ONE VOLUME

THIS IS VOLUME 1 OF 2
CONTAINING PAGES 1 TO 137

NOTE: For additional volumes, please contact the Canadian Patent Office

NOM DU FICHIER / FILE NAME :

NOTE POUR LE TOME / VOLUME NOTE:
Claims:

1. A method for preparing an anti-glypican 3 antibody with modulated plasma kinetics, said method comprising the steps of:

(a) culturing a host cell bearing a nucleic acid that encodes the anti-glypican 3 antibody under conditions allowing for expression of the nucleic acid, wherein the anti-glypican 3 antibody has an amino acid sequence altered to causes a modification in the charge of at least one amino acid residue that can be exposed on the surface of the antibody; and

(b) recovering the anti-glypican 3 antibody from the host cell culture.

2. The method according to claim 1, wherein the modulation of the plasma kinetics is increase or decrease in a parameter selected from the plasma half life, the mean plasma residence time, and the plasma clearance.

3. The method according to claim 1, wherein the modification in the charge of the amino acid residue is achieved by amino acid substitution.

4. The method according to claim 1, wherein the amino acid residue that can be exposed on the surface of the anti-glypican 3 antibody is located in a region in the anti-glypican 3 antibody other than the FcRn binding region.

5. The method according to claim 4, wherein the FcRn binding region comprises the Fc region.
6. The method according to claim 4, wherein the FcRn binding region comprises the amino acid residues of the EU numbers 250, 253, 310, 311, 314, 428, 435, 436 according to the Kabat numbering.

7. The method according to claim 1, wherein the anti-glypican 3 antibody is an IgG antibody.

8. The method according to any one of claims 1-7, wherein the amino acid residue whose charge is modified is an amino acid residue present in the heavy chain variable region or the light chain variable region.

9. The method according to claim 8, wherein the anti-glypican 3 antibody comprises a complementarity-determining region (CDR) derived from a non-human animal, a framework region (FR) derived from human, and a constant region derived from human, and wherein the modification in the charge of the amino acid residue is achieved by substitution of at least one amino acid residue that can be exposed on the antibody surface in the CDR or FR of the antibody with an amino acid residue that has a charge different from that of the amino acid residue.

10. The method according to claim 9, wherein the modification in the charge of the amino acid residue is achieved by:

(1) at least one substitution in the heavy chain variable region shown in SEQ ID NO: 1 selected from:

(a) substitution of Q that is the 43rd amino acid residue with K,
(b) substitution of D that is the 52nd amino acid residue with N, and
(c) substitution of Q that is the 107th amino acid residue with R;
and/or
(2) at least one substitution in the light chain variable region shown in SEQ ID NO: 7 selected from:
(d) substitution of E that is the 17th amino acid residue with Q,
(e) substitution of Q that is the 27th amino acid residue with R, and
(f) substitution of Q that is the 105th amino acid residue with R.

11. The method according to claim 9, wherein the modification in the charge of the amino acid residue is achieved by:
(1) at least one substitution in the heavy chain variable region shown in SEQ ID NO: 1 selected from:
(a) substitution of K that is the 19th amino acid residue with T,
(b) substitution of Q that is the 43rd amino acid residue with E,
(c) substitution of Q that is the 62nd amino acid residue with E,
(d) substitution of K that is the 63rd amino acid residue with S,
(e) substitution of K that is the 65th amino acid residue with Q, and
(f) substitution of G that is the 66th amino acid residue with D;
and/or
(2) at least one substitution in the light chain variable region shown in SEQ ID NO: 7 selected from:
(g) substitution of R that is the 24th amino acid residue with Q,
(h) substitution of Q that is the 27th amino acid residue with E,
(i) substitution of K that is the 79th amino acid residue with T,
(j) substitution of R that is the 82nd amino acid residue with S, and
(k) substitution of K that is the 112nd amino acid residue with E.
12. The method according to claim 11, further comprising at least one modification in the heavy chain constant region shown in SEQ ID NO: 31 selected from:
(a) substitution of H that is the 151st amino acid residue with Q,
(b) substitution of K that is the 157th amino acid residue with Q,
(c) substitution of R that is the 238th amino acid residue with Q,
(d) substitution of D that is the 239th amino acid residue with E,

(e) substitution of L that is the 241st amino acid residue with M, and

(f) substitution of Q that is the 302nd amino acid residue with E.

13. The method according to any one of claims 9-12, wherein the anti-glypican 3 antibody has a reduced content of fucose attached to the Fc region of the antibody.

14. An anti-glypican 3 antibody prepared by the method according to any one of claims 1-13.

15. A method for preparing an antibody with modulated plasma kinetics, said method comprising the steps of:

   (a) culturing a host cell bearing a nucleic acid that encodes the antibody under conditions allowing for expression of the nucleic acid, wherein the antibody has an amino acid sequence altered to causes a modification in the charge of at least one amino acid residue in the constant region in the antibody other than the FcRn binding region; and

   (b) recovering the antibody from the host cell culture.

16. The method according to claim 15, wherein the modulation of the plasma kinetics is increase or decrease in a parameter selected from the plasma half life, the mean plasma residence time, and the plasma clearance.
17. The method according to claim 15, wherein the modification in the charge of the amino acid residue is achieved by amino acid substitution.

18. The method according to claim 17, wherein the antibody is an IgG antibody.

19. The method according to claim 18, wherein the antibody is an IgG1 antibody.

20. The method according to claim 17, wherein the modification in the charge of the amino acid residue is achieved by substitution of at least one amino acid residue of an IgG1 antibody with a corresponding amino acid residue of an IgG4 antibody.

21. The method according to any one of claims 15-20, wherein the FcRn binding region comprises the amino acid residues of the EU numbers 250, 253, 310, 311, 314, 428, 435, and 436 according to the Kabat numbering.

22. The method according to claim 20, wherein the modification in the charge of the amino acid residue is achieved by at least one substitution in the heavy chain constant region shown in SEQ ID NO: 31 selected from:

(a) substitution of H that is the 151st amino acid residue with Q,

(b) substitution of K that is the 157th amino acid residue with Q,

(c) substitution of R that is the 238th amino acid residue with Q,
(d) substitution of D that is the 239th amino acid residue with E,
(e) substitution of L that is the 241st amino acid residue with M, and
(f) substitution of Q that is the 302nd amino acid residue with E.

23. The method according to any one of claims 15-22, wherein the antibody is an anti-glypican 3 antibody.

24. A method of stabilizing an anti-glypican 3 antibody that comprises a complementarity-determining region (CDR) derived from a non-human animal, a framework region (FR) derived from human, and a constant region derived from human, said method comprising the steps of:

   (a) culturing a host cell bearing a nucleic acid that encodes the anti-glypican 3 antibody under conditions allowing for expression of the nucleic acid, wherein the anti-glypican 3 antibody has an amino acid sequence altered to increase in the Tm value of the antibody by a modification of at least one amino acid residue; and

   (b) recovering the antibody from the host cell culture.

25. The method according to claim 24, wherein the amino acid residue is present in the FR1 region and/or the FR2 region of the heavy chain or the light chain.

26. The method according to claim 25, wherein an amino acid residue in the FR2 region of the heavy chain is substituted with an amino acid residue of the FR2 region of the VH4
subclass.

27. The method according to claim 25, wherein an amino acid residue in the FR2 region of the light chain is substituted with an amino acid residue of the FR2 region of the VK3 subclass.

28. The method according to any one of claims 24-27, wherein the substitution of the amino acid residue is achieved by:
   (1) at least one substitution in the heavy chain variable region shown in SEQ ID NO: 1 selected from:
   (a) substitution of V that is the 37th amino acid residue with I,
   (b) substitution of A that is the 40th amino acid residue with P,
   (c) substitution of M that is the 48th amino acid residue with I, and
   (d) substitution of L that is the 51st amino acid residue with I;
   and/or
   (2) at least one substitution in the light chain variable region shown in SEQ ID NO: 7 selected from:
   (e) substitution of L that is the 42nd amino acid residue with Q,
   (f) substitution of S that is the 48th amino acid residue with A, and
   (g) substitution of Q that is the 50th amino acid residue with R.
29. A method for preparing an antibody with modulated cytotoxicity, comprising the steps of:

   (a) culturing a host cell bearing a nucleic acid that encodes the antibody under conditions allowing for expression of the nucleic acid, wherein the antibody has an amino acid sequence altered to causes a modification in the charge of at least one amino acid residue that can be exposed on the surface of a cytotoxic antibody; and

   (b) recovering the antibody from the host cell culture.

30. The method according to claim 29, wherein the modification in the charge of the amino acid residue is achieved by amino acid substitution.

31. The method according to claim 29, wherein the amino acid residue that can be exposed on the surface of the antibody is located in a region in the antibody other than the FcRn binding region.

32. The method according to claim 31, wherein the FcRn binding region comprises the Fc region.

33. The method according to claim 31, wherein the FcRn binding region comprises the amino acid residues of the EU numbers 250, 253, 310, 311, 314, 428, 435, 436 according to the Kabat numbering.

34. The method according to claim 29, wherein the antibody is an IgG antibody.

35. The method according to any one of claims 29-34, wherein the amino acid residue whose charge is modified is an amino
acid residue present in the constant region of the antibody.
36. The method according to any one of claims 29-34, wherein the amino acid residue whose charge is modified is an amino acid residue present in the heavy chain variable region or the light chain variable region of the antibody.
37. The method according to claim 36, wherein the antibody is an antibody that comprises a complementarity-determining region (CDR) derived from a non-human animal, a framework region (FR) derived from human, and a constant region derived from human, and wherein the modification in the charge of the amino acid residue is achieved by substitution of at least one amino acid residue that can be exposed on the antibody surface in the CDR or FR of the antibody with an amino acid residue that has a charge different from that of the amino acid residue.
38. The method according to claim 37, wherein the modification in the charge of the amino acid residue is achieved by:
(1) at least one substitution in the heavy chain variable region shown in SEQ ID NO: 1 selected from:
(a) substitution of K that is the 19th amino acid residue with T,
(b) substitution of Q that is the 43rd amino acid residue with E,
(c) substitution of Q that is the 62nd amino acid residue with E,
(d) substitution of K that is the 63rd amino acid residue with S,
(e) substitution of K that is the 65th amino acid residue with Q, and
(f) substitution of G that is the 66th amino acid residue with D;
and/or,
(2) at least one substitution in the light chain variable region shown in SEQ ID NO: 7 selected from:
(g) substitution of R that is the 24th amino acid residue with Q,
(h) substitution of Q that is the 27th amino acid residue with E,
(i) substitution of K that is the 79th amino acid residue with T,
(j) substitution of R that is the 82nd amino acid residue with S, and
(k) substitution of K that is the 112nd amino acid residue with E.
39. The method according to claim 38, further comprising at least one substitution in the heavy chain constant region shown in SEQ ID NO: 31 selected from:
(a) substitution of H that is the 151st amino acid residue with Q,
(b) substitution of K that is the 157th amino acid residue with Q,
(c) substitution of R that is the 238th amino acid residue with Q,
(d) substitution of D that is the 239th amino acid residue with E,
(e) substitution of L that is the 241st amino acid residue with M, and
(f) substitution of Q that is the 302nd amino acid residue with E.

40. The method according to claim 36, wherein the antibody comprises a complementarity-determining region (CDR) derived from a non-human animal, a framework region (FR) derived from human, and a constant region derived from human, and wherein the modification in the charge of the amino acid residue is achieved by substitution of at least one amino acid residue that can be exposed on the antibody surface in the constant region of the antibody with an amino acid residue that has a charge different from that of the amino acid residue.

41. The method according to claim 40, wherein the substitution is at least one substitution in the heavy chain constant region shown in SEQ ID NO: 31 selected from:
(a) substitution of H that is the 151st amino acid residue with Q,
(b) substitution of K that is the 157th amino acid residue with Q,
(c) substitution of R that is the 238th amino acid residue with Q,
(d) substitution of D that is the 239th amino acid residue with E,
(e) substitution of L that is the 241st amino acid residue with M, and
(f) substitution of Q that is the 302nd amino acid residue with E.

42. The method according to any one of claims 37-41, wherein the antibody has a reduced content of fucose attached to the Fc region of the antibody.

43. An antibody prepared by the method according to any one of claims 29-42.

44. The antibody according to claim 43, wherein the antibody is an anti-glypican 3 antibody.

45. An antibody comprising:
   (1) a heavy chain variable region shown in SEQ ID NO: 1 in which the amino acid sequence comprises at least one substitution selected from:
      (a) substitution of K that is the 19th amino acid residue with T,
      (b) substitution of Q that is the 43rd amino acid residue with E,
      (c) substitution of Q that is the 62nd amino acid residue with E,
      (d) substitution of K that is the 63rd amino acid residue with S,
(e) substitution of K that is the 65th amino acid residue with Q, and
(f) substitution of G that is the 66th amino acid residue with D;
and/or
(2) a light chain variable region shown in SEQ ID NO: 7 in which the amino acid sequence comprises at least one substitution selected from:
(g) substitution of R that is the 24th amino acid residue with Q,
(h) substitution of Q that is the 27th amino acid residue with E,
(i) substitution of K that is the 79th amino acid residue with T,
(j) substitution of R that is the 82nd amino acid residue with S, and
(k) substitution of K that is the 112nd amino acid residue with E.

46. The antibody according to claim 45, comprising the heavy chain shown in SEQ ID NO: 3 and the light chain shown in SEQ ID NO: 9.

47. The antibody according to claim 45, comprising the heavy chain shown in SEQ ID NO: 5 and the light chain shown in SEQ ID NO: 11.

48. The antibody according to claim 45 which comprises a heavy chain variable region shown in SEQ ID NO: 27 and a light
chain variable region shown in SEQ ID NO: 28.

49. The antibody according to claim 45 which comprises a heavy chain variable region shown in SEQ ID NO: 27 and a light chain variable region shown in SEQ ID NO: 29.

50. The antibody according to any one of claims 45-49 comprising a constant region of a human antibody.

51. The antibody according to claim 50, wherein the constant region comprises a sequence shown in SEQ ID NO: 32 or SEQ ID NO: 33.

52. An antibody comprising:

(1) a heavy chain variable region shown in SEQ ID NO: 1 in which the amino acid sequence comprises at least one substitution selected from:

(a) substitution of Q that is the 43rd amino acid residue with K,

(b) substitution of D that is the 52nd amino acid residue with N, and

(c) substitution of Q that is the 107th amino acid residue with R;

and

(2) a light chain variable region shown in SEQ ID NO: 7 in which the amino acid sequence comprises at least one substitution selected from:

(d) substitution of E that is the 17th amino acid residue with Q,

(e) substitution of Q that is the 27th amino acid residue with
R, and

(f) substitution of Q that is the 105th amino acid residue with R.

53. The antibody according to claim 52, comprising the heavy chain variable region shown in SEQ ID NO: 4 and the light chain variable region shown in SEQ ID NO: 10.

54. The antibody according to claim 52, comprising the heavy chain variable region shown in SEQ ID NO: 6 and the light chain variable region shown in SEQ ID NO: 12.

55. The antibody according to any one of claims 52-54 comprising a constant region of a human antibody.

56. An antibody comprising at least one substitution in the amino acid sequence of the heavy chain constant region shown in SEQ ID NO: 31 selected from:

(a) substitution of H that is the 151st amino acid residue with Q,

(b) substitution of K that is the 157th amino acid residue with Q,

(c) substitution of R that is the 238th amino acid residue with Q,

(d) substitution of D that is the 239th amino acid residue with E,

(e) substitution of L that is the 241st amino acid residue with M, and

(f) substitution of Q that is the 302nd amino acid residue with E.
57. An antibody comprising a heavy chain constant region shown in SEQ ID NO: 33.

58. The antibody according to any one of claims 45-57, wherein the antibody has a reduced content of fucose attached to the Fc region of the antibody.

59. A composition comprising the antibody according to any one of claims 45-58 and a pharmaceutically acceptable carrier.

60. An anticancer agent comprising as an effective component the antibody according to any one of claims 45-58.

61. The anticancer agent according to claim 60, wherein the cancer is liver cancer.

62. A nucleic acid that encodes a polypeptide of the antibody according to any one of claims 45-58.

63. A host cell comprising the nucleic acid according to claim 62.

64. The host cell according to claim 63, wherein the host cell is a fucose transporter-deficient animal cell, a fucosyltransferase-deleted animal cell, or an animal cell in which a complex branched sugar chain modification is modified.

65. A method for preparing an antibody comprising culturing the host cell according to claim 63 or 64 and recovering a polypeptide from the cell culture.
【FIG. 6】

![Graph showing OD405 655 vs. conc. (ug/mL)]

- H0 L0
- Hsp1 & Lspd1.6
FIG. 7

A

TUMOR VOLUME (mm³)

DAYS AFTER TUMOR INOCULATION (DAYS)

B

TUMOR VOLUME (mm³)

DAYS AFTER TUMOR INOCULATION (DAYS)
【FIG. 15】

- pH7, pH16
- p7, MS5, pH16

Specific Cr51 release (%)

antibody concentration (μg/mL)