ABSTRACT

This invention provides fusion proteins comprising consecutive amino acids which beginning at the amino terminus of the protein correspond to consecutive amino acids present in (i) a cytomegalovirus human MHC-restricted peptide, (ii) a first peptide linker, (iii) a human β-2 microglobulin, (iv) a second peptide linker, (v) a HLA-A2 chain of a human MHC class I molecule, (vi) a third peptide linker, (vii) a variable region from a heavy chain of a scFv fragment of an antibody, and (viii) a variable region from a light chain of such scFv fragment, wherein the consecutive amino acids which correspond to (vii) and (viii) are bound together directly by a peptide bond or by consecutive amino acids which correspond to a fourth peptide linker, wherein the antibody from which the scFv fragment is derived specifically binds to mesothelin. This invention provides nucleic acid constructs encoding same, processes for producing same, compositions, and uses thereof.

A. scHLA-A2/SS1(scFv)

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β2-M  ─── HLA-A2 ─── scFv/SS1
       (GGGGS)_3 (SEQ ID NO: 8)
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B. pep/scHLA-A2/SS1(scFv)

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CMV  ─── β2-M  ─── HLA-A2  ─── scFv/SS1
       (GGGGS)_4 (SEQ ID NO: 6) (GGGGS)_3 (SEQ ID NO: 8)
```
Fig 4

\[ R^2 = 0.9995 \]

\[ \text{Log. (TV4451)} \]
Fig 5
Fig 6

A

B

relative cryptoschy

A431K5 A431 + TV4451 A431K5 + CTLs abse

relative apoptosis

A431K5 CELLS A431 CELLS

TV4451 µg/100ul

5 1 0.5 0.1 0.05 0.01
Fig 11

A

relative killing (%)

A431K5 cells + TV4450 + CTLs
A431 cells + TV4450 + CTLs
A431K5 Cells + CTLs

B

relative killing (%)

TV4450 (ug)

0 5 1 0.5 0.1 0.06 0.01
Fig 12

![Graph showing relative killing (%) against Fusion Molecule (µg)]
A. scHLA-A2/SS1(scFv)

B. M1/scHLA-A2/SS1(scFv)

β2-M

HLA-A2

scFv/SS1

(GGGGS)_3

(GGGGS)_3
Fig 16
Fig 17

A

K1 mAb

B

C

M1/scHLA-A2/SS1(scFv)

D
Fig 18
FUSION PROTEINS, USES THEREOF AND PROCESSES FOR PRODUCING SAME

RELATED APPLICATION/S

[0001] This application is a Continuation of U.S. patent application Ser. No. 11/804,541 filed on May 17, 2007, which claims the benefit of U.S. Provisional Patent Application No. 60/801,798 filed on May 19, 2006. The contents of the above Applications are all incorporated herein by reference.

BACKGROUND OF THE INVENTION

[0002] Throughout this application, certain publications are referenced. Full citations for these publications may be found immediately preceding the claims. The disclosures of these publications are hereby incorporated by reference into this application in order to more fully describe the state of the art to which this invention relates.

[0003] According to current immune surveillance theory, the immune system continuously locates and destroys transformed cells. However, some cells escape from an apparently effective immune response and consequently become tumors (1-4). Tumor evasion from immune response is a well established phenomenon demonstrated in numerous studies and is caused by a wide variety of suggested mechanisms (1-4). Among these mechanisms are: the production of suppressive cytokines, the loss of immunodominant peptides, the resistance to killing mechanisms (apoptosis), and the loss of MHC class I (1-4). One of the evasion mechanisms shown to be strongly correlated with tumor progression is the loss or down regulation of MHC class I molecules. This evasion mechanism is abundant in many tumors and can result from a number of different mutations. Several studies revealed weak spots in the MHC class I loading and presentation route including loss of beta-2-microglobulin, TAP1/TAP2 mutations, LMP mutations, loss of heterozygocity in the MHC genes, and down regulation of specific MHC alleles.

[0004] Current cancer immunotherapy strategies typically employ the two arms of the immune system: the humoral and the cellular systems. In the first, systemic injection of high affinity monoclonal antibodies (mAbs) directed against cell surface tumor associated antigens has demonstrated statistically significant anti-tumor activity in clinical trials (5,6). Furthermore, anti-tumor mAbs that carry effectors such as cytokines or toxins are currently being evaluated in clinical trials (7). The second major approach for specific cancer immunotherapy employs the cellular arm of the immune system, mainly the CD8+ cytotoxic T-lymphocytes. Two major strategies are currently being investigated to increase the anti-tumor effectiveness of the cellular arm of the immune system: (i) active immunization of patients with peptides known to be recognized by T-lymphocytes, and (ii) adoptive transfer therapies that enable the selection, activation, and expansion of highly active T-cell subpopulations with improved anti-tumor potency. In the first approach, MHC-restricted peptides derived from recently identified tumor associated antigens (such as gp100, the MAGE group, NY-ESO-1) are used to vaccinate patients. These tumor specific antigen-derived peptides are highly specific due to their exclusive expression in specific tissues (8-11). The second strategy, adoptive cell transfer, has recently shown impressive results in metastatic melanoma patients in which highly selected, tumor-reactive T-cells against different over-expressed self-derived differentiation antigens were isolated, expanded ex-vivo and reintroduced to the patients. In this approach, a persistent clonal repopulation of T-cells, proliferation in vivo, functional activity, and trafficking to tumor sites were demonstrated (12-14).

[0005] A new immunotherapeutic approach recently presented takes advantage of two well-established areas: (i) the known effectiveness of CD8+ cytotoxic T-lymphocytes in the elimination of cells presenting highly immunogenic MHC/peptide complexes, and (ii) the tumor-specific cell surface antigens targeting via recombinant fragments of antibodies, mainly single chain Fv fragments (scFvs). This approach utilizes a recombinant fusion protein composed of two functionally distinct entities: (i) a single-chain MHC class I molecule that carries a highly immunogenic tumor or viral-derived peptide, and (ii) a tumor-specific, high-affinity scFv fragment (15). Several groups have previously shown that a biotinylated MHC peptide multimerized on streptavidin or monomeric HLA-A2/influenza (Flu) matrix peptide complexes coupled via chemical conjugation to tumor-specific antibodies could induce in vitro T-lymphocyte-mediated lysis of coated tumor cells (16-20). However, these approaches utilize chemical conjugation and use whole antibodies or larger fragments, e.g. Fab fragments. However, production and homogeneity owing to the coupling strategy as well as tumor penetration capability are limited due to the large size of such molecules. Lev et al. describe a genetic fusion created between a single-chain recombinant HLA-A2 and tumor specific scFvs. These fusions were shown to be functional in vitro and in vivo, being able to specifically induce T-lymphocyte mediated lysis and in vivo lysis of target-coated tumor cells (15). The stability of the new chimeric molecule is highly dependent on the presence of the peptide in the MHC groove. Therefore, dissociation of the peptide from the scHLA-A2 domain of the chimeric molecule can impair its stability. Oved et al. addressed this problem by constructing new chimeric molecules in which the peptide is connected to the scHLA-A2/scFv construct via a short linker. This new fusion protein was tested for its in vitro biochemical and biological activity (21).

[0006] There is a widely recognized need for a new fusion protein that can maintain its dual activity: bind tumor target cells through the scFv moiety as well as mediate potent, effective and specific cytotoxicity through the recruitment of CD8+ T-cells whose specificity is governed by the covalently linked HLA-A2-restricted peptide.

[0007] The MHC class I-restricted CD8+ cytotoxic T-cell (CTL) effector arm of the adaptive immune response is best equipped to recognize tumor cells as foreign and initiate the cascade of events resulting in tumor destruction. However, tumors have developed sophisticated strategies to escape immune effector mechanisms, of which the best-studied is the downregulation of MHC class I molecules which prevent the antigens recognized by CTLs.

[0008] To overcome the limitation of previous approaches and develop new approaches for immunotherapy, a recombinant molecule was constructed in which a single-chain MHC is specifically targeted to tumor cells through its fusion to cancer specific-recombinant antibody fragments or a ligand that binds to receptors expressed by tumor cells.

SUMMARY OF THE INVENTION

[0009] This invention provides a fusion protein comprising consecutive amino acids which, beginning at the amino terminus of the protein, correspond to consecutive amino acids present in (i) a cytomegalovirus human MHC-restricted pep-
tide, (ii) a first peptide linker, (iii) a human β-2 microglobulin, (iv) a second peptide linker, (v) a HLA-A2 chain of a human MHC class I molecule, (vi) a third peptide linker, (vii) a variable region from a heavy chain of a scFv fragment of an antibody, and (viii) a variable region from a light chain of such scFv fragment, wherein the consecutive amino acids which correspond to (vii) and (viii) are bound together directly by a peptide bond or by consecutive amino acids which correspond to a fourth peptide linker and the scFv fragment is derived from an antibody which specifically binds to mesothelin.

This invention also provides compositions comprising the fusion protein and a carrier.

This invention further provides a nucleic acid construct encoding a fusion protein comprising consecutive amino acids which, beginning at the amino terminus of the protein, correspond to consecutive amino acids present in (i) a cytomegalovirus human MHC-restricted peptide, (ii) a first peptide linker, (iii) a human β-2 microglobulin, (iv) a second peptide linker, (v) a HLA-A2 chain of a human MHC class I molecule, (vi) a third peptide linker, (vii) a variable region from a heavy chain of a scFv fragment of an antibody, and (viii) a variable region from a light chain of such scFv fragment, wherein the consecutive amino acids which correspond to (vii) and (viii) are bound together directly by a peptide bond or by consecutive amino acids which correspond to a fourth peptide linker and the scFv fragment is derived from an antibody which specifically binds to mesothelin.

This invention still further provides an isolated preparation of bacterially-expressed inclusion bodies comprising over 30 percent by weight of a fusion protein in accordance with the invention.

This invention also provides a process for producing a fusion protein comprising culturing a transformed cell comprising the fusion protein, so that the fusion protein is expressed, and recovering the fusion protein so expressed.

This invention further provides a method of selectively killing a tumor cell which comprises contacting the cell with the fusion protein of the invention in an amount effective to initiate a CTL-mediated immune response against the tumor cell so as to thereby kill the tumor cell.

Finally, this invention further provides a method of treating a tumor cell which expresses mesothelin on its surface, which comprises contacting the tumor cell with the fusion protein according to the invention in an amount effective to initiate a CTL-mediated immune response against the tumor cell so as to thereby treat the tumor cell.

As an exemplary molecule of the present invention, a single-chain MHC molecule composed of β2 microglobulin fused to the α1, α2 and α3 domains of HLA-A2 via a short peptide linker (15 amino acids) was fused to the scFv SS1 which targets mesothelin. To construct a fusion protein with covalently linked peptide a 9 amino acids peptide derived from the CMV pp 65 protein NLYPMVATV (SEQ ID NO:4) was fused to the N-terminus of the scFV-HLA-A2/SS1(scFv) fusion protein via a 20 amino acid linker GGGGSGGGSGGGSGGGGS (SEQ ID NO:6). The fusion protein was expressed in E. coli and functional molecules were produced by in vitro refolding in the presence of CMV/scFV-HLA-A2/SS1(scFv). Flow cytometry studies revealed the ability to decorate antigen-positive, HLA-A2-negative human tumor cells with HLA-A2-peptide complexes in a manner that was entirely dependent upon the specificity of the targeting antibody fragment.

This invention further provides a method of selectively killing a tumor cell which comprises contacting the cell with the fusion protein of the invention in an amount effective to initiate a CTL-mediated immune response against the tumor cell so as to thereby kill the tumor cell.

Brief description of the figures

Figs. 1A-B Schematic representation of scFV-HLA-A2/SS1 (scFv) and a pep(CMV)scFV-HLA-A2/SS1 (scFv) (Compound A).

Fig. 1A illustrates the C-Terminal of the scFV-HLA-A2 fused to the N-terminus of SS1 (scFv) via a 4 amino acid linker. Fig. 1B illustrates that the CMV pp 65 peptide, i.e. NLYPMVATV (SEQ ID NO:4) was fused to the N-terminus of the scFV-HLA-A2/SS1 (scFv) via a 20 amino acid linker GGGGSGGGSGGGSGGGS (SEQ ID NO:6).

Fig. 2 Nucleic acid sequence encoding Compound A (SEQ ID NO:1).

Figs. 3A-B Expression and purification of Compound A.

Fig. 3A shows the SDS/PAGE analysis of isolated inclusion bodies. Fig. 3B shows the SDS/PAGE analysis of Compound A after purification on ion-exchange chromatography.

Fig. 4 Binding of Compound A to recombinant mesothelin.

Mesothelin was immobilized on immuno-plates and dose-dependent binding of Compound A was monitored by conformative sensitive mAb W6 (33, 34).

Fig. 5A-D Binding of Compound A to mesothelin-expressing cells.

Fig. 5A-B demonstrates the flow cytometry analysis of the binding of Compound A to mesothelin-positive
HLA-A2-negative A431K5 cells and mesothelin-negative HLA-A2-negative A431 cells. FIG. 5A shows the binding of the K1 mAb (31,32) to A431K5 cells, and FIG. 5B shows the presence of the K1 mAb to A431 cells. FIG. 5C shows the binding of Compound A to A431K5 cells, and FIG. 5D shows the absence of binding of Compound A to A431 cells. The binding was monitored using anti-HLA-A2 specific antibody BB7.2 (35) and a FITC-labeled secondary antibody.

[0027] FIG. 6A-B Potentiation of CTL-mediated lysis of HLA-A2 negative tumor cells by Compound B. In FIG. 6A, the mesothelin-transfected A431K5 cells and the parental mesothelin-negative A431 cells were incubated with Compound A (10 μg) and CMV specific CTLs in a [35S]methionine release assay. FIG. 6B demonstrates dose-dependent activity of Compound A when mesothelin-transfected A431K5 cells and the parental mesothelin-negative A431 cells were incubated with different concentrations of Compound A and CMV-specific CTLs in a [35S]methionine release assay.

[0028] FIG. 7 Schematic representation of the pep/scHLA-A2/SS1(scFv) (Compound B). In Compound B, the peptide NLVPVMVAIV (SEQ ID NO:4) was fused to the N-terminus of scHLA-A2/SS1 (scFv) via a 15 amino acid linker GGGSSGGGSSGGGGS (SEQ ID NO:8).

[0029] FIG. 8 Nucleic acid sequence encoding Compound B (SEQ ID NO:22).

[0030] FIGS. 9A-B Expression and purification of Compound B.

[0031] FIG. 9A shows SDS/PAGE analysis of isolated inclusion bodies. FIG. 9B shows SDS/PAGE analysis of Compound B after purification on ion-exchange chromatography.

[0032] FIGS. 10A-F Binding of Compound B to mesothelin-expressing cells.

[0033] FIGS. 10A-F demonstrate the flow cytometry analysis of the binding of Compound B to mesothelin-positive HLA-A2-negative A431K5 cells and mesothelin-negative HLA-A2-negative A431 cells. FIG. 10A shows the binding of K1 mAb to A431K5 cells, and FIG. 10B shows the absence of binding of K1 mAb to A431 cells (B). FIG. 10C shows the binding of Compound B to A431K5 cells, and FIG. 10D shows the absence of binding of Compound B to A431 cells. FIG. 10E shows the comparison between the binding of Compound A and Compound B to A431K5 cells, and FIG. 10F shows the absence of binding of Compound A and Compound B to A431 cells. The binding was monitored using anti-HLA-A2 specific antibody BB7.2 and a FITC-labeled secondary antibody.

[0034] FIGS. 11A-B Potentiation of CTL-mediated lysis of HLA-A2-negative tumor cells by Compound B.

[0035] In FIG. 11A, mesothelin-transfected A431K5 cells and the parental mesothelin-negative A431 cells were incubated with Compound B (10 μg) and CMV-specific CTLs in a [35S]methionine release assay. FIG. 11B demonstrates dose-dependent activity of Compound B, when mesothelin-transfected A431K5 cells and the parental mesothelin-negative A431 cells were incubated with different concentrations of Compound A and CMV-specific CTLs in a [35S]methionine release assay.

[0036] FIG. 12

[0037] Potentiation of CTL-mediated lysis of HLA-A2 negative tumor cells by Compound B and Compound A. Mesothelin-transfected A431K5 cells and the parental mesothelin-negative A431 cells were incubated with different concentrations of Compound B or Compound A and with CMV-specific CTLs in a [35S]methionine release assay. The figure shows results of incubation of Compound A with A431K5 cells, incubation of Compound B with A431K5 cells, incubation of Compound A with A431 cells, and incubation of Compound B with A431 cells.

[0038] FIGS. 13A-B

[0039] Schematic representation of scHLA-A2/SS1(scFv) and M1cov/scHLA-A2/SS1(scFv).

[0040] FIG. 13A shows the C-terminus of the scHLA-A2 fused to the N-terminus of scFv via 4 amino acid linker. FIG. 13B shows the M158-66 peptide fused to the N-terminus of the scHLA-A2/SS1(scFv) via a 15 amino acid linker GGGSSGGGSSGGGGS (SEQ ID NO:8).

[0041] FIG. 14 Nucleic acid sequence encoding the M1cov/scHLA-A2/SS1(scFv) fusion protein (SEQ ID NO:23).

[0042] FIGS. 15A-B Expression and purification of the M1-cov/scHLA-A2/SS1(scFv) fusion protein.

[0043] FIG. 15A shows the SDS/PAGE analysis of isolated inclusion bodies. FIG. 15B shows the SDS/PAGE analysis of M1-cov/scHLA-A2/SS1 (scFv) fusion protein after purification on ion-exchange chromatography.

[0044] FIG. 16 Binding of the M1-cov/scHLA-A2/SS1 (scFv) fusion protein to recombinant Mesothelin. Mesothelin was immobilized onto immuno-plates and dose-dependent binding of M1-cov/scHLA-A2/SS1 (scFv) was monitored by confocal detection of mAb (W6).

[0045] FIGS. 17A-D Binding of M1-cov/scHLA-A2/SS1 (scFv) fusion protein to Mesothelin expressing cells. FIGS. 17A-D show flow cytometry analysis of the binding of M1-cov/scHLA-A2/SS1 (scFv) to mesothelin-positive HLA-A2-negative A431K5 cells and mesothelin-negative HLA-A2-negative A431 cells. FIG. 17A shows the binding of K1 mAb to A431K5 cells, and FIG. 17B shows the absence of binding of K1 mAb to A431 cells. FIG. 17C shows the binding of M1-cov/scHLA-A2/SS1 (scFv) fusion protein to A431K5 cells, and FIG. 17D shows the absence of binding of M1-cov/scHLA-A2/SS1 (scFv) fusion protein to A431 cells. The binding was monitored using anti-HLA-A2 specific antibody BB7.2 and a FITC-labeled secondary antibody.

[0046] FIG. 18 Potentiation of CTL-mediated lysis of HLA-A2-negative tumor cells by M1-cov/scHLA-A2/SS1 (scFv) fusion protein. Mesothelin-transfected A431K5 cells and the parental mesothelin-negative A431 cells were incubated with different concentrations of M1-cov/scHLA-A2/SS1(scFv) and with M1 specific HLA-A2-restricted CTLs in a [35S]methionine release assay.

DETAILED DESCRIPTION OF THE INVENTION

[0047] This invention provides a fusion protein comprising consecutive amino acids which, beginning at the amino terminus of the protein, correspond to consecutive amino acids present in (i) a cytomegalovirus human MHC-restricted peptide, (ii) a first peptide linker, (iii) a human β-2 microglobulin, (iv) a second peptide linker, (v) a HLA-A2 chain of a human MHC class I molecule, (vi) a third peptide linker, (vii) a variable region from a heavy chain of a scFv fragment of an antibody, and (viii) a variable region from a light chain of such scFv fragment, wherein the consecutive amino acids which correspond to (vii) and (viii) are bound together directly by a peptide bond or by consecutive amino acids which correspond to a fourth peptide linker and the scFv fragment is derived from an antibody which specifically binds
to mesothelin. In one embodiment, the first peptide linker has the amino acid sequence GGGGGSGGGGGSGGGGSGGGGG (SEQ ID NO:6). In another embodiment, the second peptide linker has the amino acid sequence GGGGGSGGGGGSGGGGSGGGG (SEQ ID NO:8). In another embodiment, the third peptide linker has the amino acid sequence ASGG (SEQ ID NO:10). In another embodiment, the fourth peptide linker has the amino acid sequence GVGGSGGGGGSGGGG (SEQ ID NO:19). In another embodiment, the cytomegalovirus human MHC-restricted peptide has the amino acid sequence NLVP-MVATV (SEQ ID NO:4).

[0048] As used herein, “first peptide linker”, “second peptide linker” and “fourth peptide linker” refer to peptides composed of a monomeric peptide whose amino acid sequence is GGGGG (SEQ ID NO:20) or a multimer thereof, wherein X may be any amino acid. These peptide linkers may be a multimer of 2-10 of such monomeric peptide. In any such multimer, each monomeric peptide may be the same as or different from other monomeric peptides in the multimer depending on the identity of amino acid X. In one embodiment, X in the monomeric peptide is the amino acid valine (V). In another embodiment, X in the monomeric peptide is the amino acid glycine (G). In presently preferred embodiments, the peptide linker comprises a multimer of three or four monomeric peptides, particularly a multimer of three monomeric peptides in which the most N-terminal X is the amino acid V, and the second and third X are the amino acid G.

[0049] In one embodiment, the sequence of the consecutive amino acids corresponding to (vii), followed by the fourth peptide linker, followed by (viii) is set forth in SEQ ID NO:12.

[0050] In another embodiment, the consecutive amino acids of the fusion protein, Compound A, have the amino acid sequence set forth in SEQ ID NO:2.

[0051] This invention also provides a composition comprising a fusion protein in accordance with the invention and a carrier. In one embodiment, the fusion protein is present in the composition in a therapeutically effective amount and the carrier is a pharmaceutically acceptable carrier.

[0052] This invention also provides a nucleic acid construct encoding a fusion protein comprising consecutive amino acids which, beginning at the amino terminus of the protein, correspond to consecutive amino acids present in (i) a cytomegalovirus human MHC-restricted peptide, (ii) a first peptide linker, (iii) a human β-2 microglobulin, (iv) a second peptide linker, (v) a HLA-A2 chain of a human MHC class I molecule, (vi) a third peptide linker, (vii) a variable region from a heavy chain of a scFv fragment of an antibody, and (viii) a variable region from a light chain of such scFv fragment, wherein the consecutive amino acids which correspond to (vii) and (viii) are bound together directly by a peptide bond or by consecutive amino acids which correspond to a fourth peptide linker and the scFv fragment is derived from an antibody which specifically binds to mesothelin. In one embodiment, the nucleic acid construct has the nucleic acid sequence set forth in SEQ ID NO:1.

[0053] This invention also provides a vector comprising the nucleic acid construct of the invention. Examples of such vectors are plasmids, viruses, phages, and the like.

[0054] This invention further provides an expression vector comprising the nucleic acid construct of the invention and a promoter operatively linked thereto.

[0055] This invention also provides a transformed cell comprising a vector according to the invention. The transformed cell may be an eukaryotic cell, e.g. one selected from the group consisting of a mammalian cell, an insect cell, a plant cell, a yeast cell and a protozoan cell. Alternatively, the transformed cell may be a bacteriobial cell.

[0056] This invention provides an isolated preparation of bacterially-expressed inclusion bodies comprising over 30 percent by weight of a fusion protein according to the invention.

[0057] This invention also provides a process for producing a fusion protein comprising culturing the transformed cell of the invention so that the fusion protein is expressed, and recovering the fusion protein so expressed. In one embodiment, the recovery of the fusion protein comprises subjecting the expressed fusion protein to size exclusion chromatography. In another embodiment, the fusion protein is expressed in inclusion bodies. In one embodiment, the process further comprises treating the inclusion bodies so as to separate and refold the fusion protein and thereby produce the fusion protein in active form. In another embodiment, treating of the inclusion bodies to separate the fusion protein therefrom comprises contacting the inclusion bodies with a denaturing agent.

[0058] As used herein, an “active form” of the fusion protein means a three dimensional conformation of the fusion protein which permits the fusion protein to specifically bind to mesothelin when mesothelin is present on the surface of a tumor cell.

[0059] This invention also provides a method of selectively killing a tumor cell, which comprises contacting the cell with the fusion protein of the invention in an amount effective to initiate a CTL-mediated immune response against the tumor cell so as to thereby kill the tumor cell. In one embodiment, the tumor cell is in a patient and the contacting is effected by administering the fusion protein to the patient.

[0060] This invention further provides a method of treating a tumor cell which expresses mesothelin on its surface, which comprises contacting the tumor cell with the fusion protein according to the invention in an amount effective to initiate a CTL-mediated immune response against the tumor cell so as to thereby treat the tumor cell. In one embodiment, the tumor cell is present in a solid tumor. In another embodiment, the solid tumor is a tumor associated with ovarian, lung, pancreatic or head/neck cancer, or mesothelioma.

[0061] The present invention provides (i) novel fusion proteins; (ii) processes of preparing same; (iii) nucleic acid constructs encoding same; and (iv) methods of using same for selective killing of cells, cancer cells in particular.

[0062] The principles and operation of the present invention may be better understood with reference to the figures and description set forth herein.

[0063] It is to be understood that the invention is not limited in its application to the details set forth in the description or as exemplified. The invention encompasses other embodiments and is capable of being practiced or carried out in various ways. Also, it is to be understood that the phraseology and terminology employed herein is for the purpose of description and should not be regarded as limiting.

[0064] Tumor progression is often associated with the secretion of immune-suppressive factors and/or the down-regulation of MHC class I antigen-presentation functions (2). Even when a specific CTL response is demonstrated in patients, this response is low because the anti-tumor CTL
population is rare, very infrequent, and in some cases the CLTs are not functional or anergic (26). Moreover, it is well-established that the number of MHC-peptide complexes on the surface of tumor cells that present a particular tumor-associated peptide is low (27). Significant progress toward developing vaccines that can stimulate an immune response against tumors has involved the identification of the protein antigens associated with a given tumor type and epitope mapping of tumor antigens for MHC class I and class II restricted binding motifs were identified and are currently being used in various vaccination programs (14, 11, 8). MHC class I molecules presenting the appropriate peptides are necessary to provide the specific signals for recognition and killing by CTLs. However, the principal mechanism of tumor escape is the loss, downregulation or alteration of HLA profiles that may render the target cell unresponsive to CTL lysis, even if the cell expresses the appropriate tumor antigen.

[0065] The present invention provides a new approach to circumvent this problem. While reducing the present invention to practice, tumor-specific targeting of class I MHC-peptide complexes on tumor cells was shown to be an effective and efficient strategy to render HLA-A2-negative cells susceptible to lysis by relevant HLA-A2-restricted CTLs. This new strategy of redirecting CTLs against tumor cells takes advantage of the use of recombinant anti-mesothelin antibody fragment and CMV ligand that can localize on malignant cells that express a tumor with a relatively high degree of specificity.

[0066] The anti-mesothelin antibody targeting fragment and CMV ligand are fused to a single-chain HLA-A2 molecule that can be folded efficiently and functionally.

[0067] The results presented herein provide a clear demonstration of the usefulness of the approach of the present invention to recruit active CTLs for tumor cell killing via cancer-specific antibody or ligand guided targeting of scMHC-peptide complexes. These results pave the way for the development of a new immunotherapeutic approach based on naturally occurring cellular immune responses which are redirected against the tumor cells.

[0068] It will be appreciated that the fusion protein of the present invention or portions thereof can be prepared by several ways, including solid phase protein synthesis. However, in the preferred embodiment of the invention, at least major portions of the molecules, e.g., the scHLA-A2 domain (with or without the CMV peptide) and the scFV domain are generated by translation of a respective nucleic acid construct or constructs encoding the molecule.

[0069] Accordingly, one to three open reading frames are required to synthesize the molecules of FIG. 1B via translation. These open reading frames can reside on a single, two or three nucleic acid molecules. Thus, for example, a single nucleic acid construct can carry one, two or all three open reading frames. One to three cis-acting regulatory sequences can be used to control the expression of the one to three open reading frames. For example, a single cis-acting regulatory sequence can control the expression of one, two or three open reading frames, in a cistron-like manner. In the alternative, three independent cis-acting regulatory sequences can be used to control the expression of the three open reading frames. Other combinations are also envisaged.

[0070] The open reading frames and the cis-acting regulatory sequences can be carried by one to three nucleic acid molecules. For example, each open reading frame and its cis-acting regulatory sequence are carried by a different nucleic acid molecule, or all of the open reading frames and their associated cis-acting regulatory sequences are carried by a single nucleic acid molecule. Other combinations are also envisaged.

[0071] Expression of the fusion protein can be effected by transformation/transfection and/or co-transformation/co-transfection of a single cell or a plurality of cells with any of the nucleic acid molecules, serving as transformation/transfection vectors (e.g., as plasmids, phages, phagemids or viruses).

[0072] It will be appreciated that the fusion protein whose amino acid sequence is set forth in SEQ ID NO:2 and includes the N-terminal amino acid methionine, likely represents the fusion protein as expressed in a bacterial cell. Depending on the specific bacterial cell employed to express the fusion protein, the N-terminal methionine may be cleaved and removed. Accordingly, it is contemplated that fusion proteins in accordance with this invention encompass both those with, and those without, a N-terminal methionine. In general, when a fusion protein in accordance with the invention is expressed in a eukaryotic cell, it would lack the N-terminal methionine. Therefore, it is to be appreciated that the amino acid sequence of expressed fusion proteins according to the invention may include or not include such N-terminal methionine depending on the type of cells in which the proteins are expressed.

[0073] Whenever and wherever used, the linker peptide is selected of an amino acid sequence which is inherently flexible, such that the polypeptides connected thereby independently and natively fold following expression thereof, thus facilitating the formation of a functional or active single chain (sc) human β₂M/HLA complex, antibody targeting or human β₂M/HLA-CMV restricted antigen complex.

[0074] Any of the nucleic acid constructs described herein comprise at least one cis-acting regulatory sequence operably linked to the coding polynucleotides therein. Preferably, the cis-acting regulatory sequence is functional in bacteria. Alternatively, the cis-acting regulatory sequence is functional in yeast. Still alternatively, the cis-acting regulatory sequence is functional in animal cells. Yet alternatively, the cis acting regulatory sequence is functional in plant cells.

[0075] The cis-acting regulatory sequence can include a promoter sequence and additional transcriptional or a translational enhancer sequences all of which serve for facilitating the expression of the polynucleotides when introduced into a host cell. Specific examples of promoters are described hereinbelow in context of various eukaryotic and prokaryotic expression systems and in the examples section which follows.

[0076] It will be appreciated that a single cis-acting regulatory sequence can be utilized in a nucleic acid construct to direct transcription of a single transcript which includes one or more open reading frames. In the latter case, an internal ribosome entry site (IRES) can be utilized so as to allow translation of the internally positioned nucleic acid sequence.

[0077] Whenever co-expression of independent polypeptides in a single cell is of choice, the construct or constructs employed must be configured such that the levels of expression of the independent polypeptides are optimized, so as to obtain highest proportions of the final product.

[0078] Preferably a promoter (being an example of a cis-acting regulatory sequence) utilized by the nucleic acid construct(s) of the present invention is a strong constitutive promoter such that high levels of expression are attained for the polynucleotides following host cell transformation.
It will be appreciated that high levels of expression can also be effected by transforming the host cell with a high copy number of the nucleic acid construct(s), or by utilizing cis acting sequences which stabilize the resultant transcript and as such decrease the degradation or “turn-over” of such a transcript.

As used herein, the phrase “transformed cell” describes a cell into which an exogenous nucleic acid sequence is introduced to thereby stably or transiently genetically alter the host cell. It may occur under natural or artificial conditions using various methods well known in the art some of which are described in detail hereinbelow in context with specific examples of host cells.

The transformed host cell can be a eukaryotic cell, such as, for example, a mammalian cell, an insect cell, a plant cell, a yeast cell and a protozoa cell, or alternatively, the cell can be a bacterial cell.

When utilized for eukaryotic host cell expression, the nucleic acid construct(s) according to the present invention can be a shuttle vector, which can propagate both in E. coli (wherein the construct comprises an appropriate selectable marker and origin of replication) and be compatible for expression in eukaryotic host cells. The nucleic acid construct(s) according to the present invention can be, for example, a plasmid, a bacmid, a phagemid, a cosmid, a phage, a virus or an artificial chromosome.

Suitable mammalian expression systems include, but are not limited to, pCDNA3, pCDNA3.1(+) and pZeoSV2 (+/-), pSecTag2, pDisplay, pEF/myc/cyto, pCMV/myc/cyto, pCR3.1, which are available from Invitrogen™ Corporation (Carlsbad, Calif. USA), pCl which is available from Promega™ Corporation (Madison Wis. USA), pBK-RSV and pBR-CMV which are available from Stratagene® (La Jolla, Calif. USA), pTRES which is available from Clontech® Laboratories, Inc. (Mountain View, Calif. USA), and their derivatives.

Insect cell cultures can also be utilized to express the nucleic acid sequences of the present invention. Suitable insect expression systems include, but are not limited to the baculovirus expression system and its derivatives which are commercially available from numerous suppliers such as maxBac™ (Invitrogen™ Corporation, Carlsbad, Calif. USA), BacPak™ (Clontech® Laboratories, Inc. Mountain View, Calif. USA), or Bac-to-Bac™ (Invitrogen™/Gibco®, Carlsbad, Calif. USA).

Expression of the nucleic acid sequences of the present invention can also be effected in plants cells. As used herein, the phrase “plant cell” can refer to plant protoplasts, cells of a plant tissue culture, cells of plant derived tissues or cells of whole plants.

There are various methods of introducing nucleic acid constructs into plant cells. Such methods rely on either stable integration of the nucleic acid construct or a portion thereof into the genome of the plant cell, or on transient expression of the nucleic acid construct in which case these sequences are not stably integrated into the genome of the plant cell.

There are two principle methods of effecting stable genomic integration of exogenous nucleic acid sequences such as those included within the nucleic acid construct of the present invention into plant cell genomes:


The Agrobacterium system includes the use of plasmid vectors that contain defined DNA segments that integrate into the plant genomic DNA. Methods of inoculation of the plant tissue vary depending upon the plant species and the Agrobacterium delivery system. A widely used approach is the leaf disc procedure, see for example, Horsch et al. in Plant Molecular Biology Manual A5, Kluwer Academic Publishers, Dordrecht (1988) p. 1-9. A supplementary approach employs the Agrobacterium delivery system in combination with vacuum infiltration. The Agrobacterium system is especially viable in the creation of stably transformed dicotyledonous plants.

There are various methods of direct DNA transfer into plant cells. In electroporation, protoplasts are briefly exposed to a strong electric field. In microinjection, the DNA is mechanically injected directly into the cells using a very small micropipettes. In microparticle bombardment, the DNA is adsorbed on microprojectiles such as magnesium sulfate crystals, tungsten particles or gold particles, and the microprojectiles are physically accelerated into cells or plant tissues. Direct DNA transfer can also be utilized to transiently transform plant cells.

In any case suitable plant promoters which can be utilized for plant cell expression of the first and second nucleic acid sequences, include, but are not limited to CaMV 35S promoter, ubiquitin promoter, and other strong promoters which can express the nucleic acid sequences in a constitutive or tissue specific manner.

Plant viruses can also be used as transformation vectors. Viruses that have been shown to be useful for the transformation of plant cell hosts include CaV, TMV and BV. Transformation of plants using plant viruses is described in U.S. Pat. No. 4,855,237 (BVG), EP-A 67,553 (TMV), Japanese Published Application No. 63-149693 (TMV), EPA 194, 899 (BV), EPA 278,667 (BV); and Gluzman, Y. et al., Com- munications in Molecular Biology; Viral Vectors, Cold
Pseudovirus particles for use in expressing foreign DNA in many hosts, including plants, is described in WO 87/06261.


When the virus is a DNA virus, the constructions can be made to the virus itself. Alternatively, the virus can first be cloned into a bacterial plasmid for ease of constructing the desired viral vector with the nucleic acid sequences described above. The virus can then be excised from the plasmid. If the virus is a DNA virus, a bacterial origin of replication can be attached to the viral DNA, which is then replicated by the bacteria. Transcription and translation of this DNA will produce the coat protein which will encapsidate the viral DNA. If the virus is an RNA virus, the virus is generally cloned as a cDNA and inserted into a plasmid. The plasmid is then used to make all of the constructions. The RNA virus is then produced by transcribing the viral sequence of the plasmid and translation of the viral genes to produce the coat protein(s) which encapsidate the viral RNA.

Construction of plant RNA viruses for the introduction and expression in plants of non-viral exogenous nucleic acid sequences such as those included in the construct of the present invention is demonstrated by the above references as well as in U.S. Pat. No. 5,316,931.

Yeast cells can also be utilized as host cells by the present invention. Numerous examples of yeast expression vectors suitable for expression of the nucleic acid sequences of the present invention in yeast are known in the art and are commercially available. Such vectors are usually introduced in a yeast host cell via chemical or electroporation transformation methods well known in the art. Commercially available systems include, for example, the pYES™ (Invitrogen™ Corporation, Carlsbad Calif., USA) or the YEX™ (Clontech® Laboratories, Mountain View, Calif., USA) expression systems.

It will be appreciated that when expressed in eukaryotic expression systems such as those described above, the nucleic acid construct preferably includes a signal peptide encoding sequence such that the polypeptides produced from the first and second nucleic acid sequences are directed via the attached signal peptide into secretion pathways. For example, in mammalian, insect and yeast host cells, the expressed polypeptides can be secreted to the growth medium, while in plant expression systems the polypeptides can be secreted into the apoplast, or directed into a subcellular organelle.

A bacterial host can be transformed with the nucleic acid sequence via transformation methods well known in the art, including for example, chemical transformation (e.g., CaCl₂) or electroporation.

Numerous examples of bacterial expression systems which can be utilized to express the nucleic acid sequences of the present invention are known in the art. Commercially available bacterial expression systems include, but are not limited to, the pET™ expression system (Novagen®, EMB Biosciences, San Diego, Calif., USA), pSE™ expression system (Invitrogen™ Corporation, Carlsbad Calif., USA) or the pGEX™ expression system (Amersham Biosciences, Piscataway, N.J., USA).

As is further described in the Experimental Details section which follows, bacterial expression is particularly advantageous since the expressed polypeptides form substantially pure inclusion bodies readily amenable to recovery and purification of the expressed polypeptide.

Thus, this invention provides a preparation of bacterial-expressed inclusion bodies which are composed of over 30%, preferably over 50%, more preferably over 75%, most preferably over 90% by weight of the fusion protein or a mixture of fusion proteins of the present invention. The isolation of such inclusion bodies and the purification of the fusion protein(s) therefrom are described in detail in the Experimental Details section which follows. Bacterial expression of the fusion protein(s) can provide high quantities of pure and active forms of fusion proteins.

As is further described in the Experimental Details section which follows, the expressed fusion proteins form substantially pure inclusion bodies which are readily isolated via fractionation techniques well known in the art and purified via for example denaturing-renaturing steps.

The fusion proteins of the invention may be reassembled and refolded in the presence of a MHC-restricted peptide, which is either linked to, co-expressed with or mixed with other polypeptides of the invention and being capable of binding the single chain MHC class I polypeptide. As is further described in the examples section, this enables to generate a substantially pure MHC class I-antigenic peptide complex which can further be purified via size exclusion chromatography.

It will be appreciated that the CMV peptide used for refolding can be co-expressed along with (as an independent peptide) or be fused to the HLA-A2 chain of the MHC Class I molecule in the bacteria. In such a case the expressed fusion protein and polypeptide co-form inclusion bodies which can be isolated and utilized for MHC class I-antigenic peptide complex formation.

The following section provides specific examples for each of the various aspects of the invention described herein. These examples should not be regarded as limiting in any way, as the invention can be practiced in similar, yet somewhat different ways. These examples, however, teach one of ordinary skills in the art how to practice various alternatives and embodiments of the invention.


Experimental Details

Materials and Methods:

Cloning of Compound A

[0108] The sHLa-A2/SS1 (sFv) was constructed as previously described by linking the C-termine of sHLa-A2 to the N-termine of the SS1 sFv via a short linker ASGG (SEQ ID NO:4) (15). To construct the sHLa-A2/SS1 (sFv) with covalently bound MHC-restricted peptide, the MHC-restricted peptide was fused with the peptide linker GGGSGGGGGGGGGGGGGG (SEQ ID NO:6) to the N-termine of the sHLa-A2/SS1 (sFv) molecule by a PCR overlap extension reaction with the primers: 5'CMVcovLL (cassette) and 3'CMVcovLL (cassette) was performed by incubating the primers at 95°C for 2 min followed by 1 h incubation at room temperature. The ligation product was transformed to E.coli DH5a for plasmid amplification. Plasmid was purified by QIAGEN® Miniprep™, DNA isolation kit (Qiagen®, Inc., Valencia, Calif. USA) and samples were set for sequence analysis.

[0111] Expression Refolding and Purification of Compound A

[0112] Compound A was expressed in E.coli LB21 (ADE3) cells (Novagen®, Madison, Wis. USA) as inclusion bodies. Compound A was transformed to E.coli cells by heat shock, cells were plated on LBAMP plates and incubated over night at 37°C. Colonies were transferred to rich medium (super broth) supplemented with glucose, MgSO4, AMP and salts. The cells were grown to DO=2 (600nm) at 37°C, induced with IPTG (final concentration 1mM) and incubated for an additional 3 h at 37°C.

[0113] Inclusion bodies were purified from cell pellet by cell disruption with 0.2 mg/ml of lysozyme followed by the addition of 2.5% Triton X-100 (Ocylphenolpolyethylenelether), Roche Diagnostics GmbH, Roche Applied Science, Mannheim, Germany and 0.5M NaCl. The pellets of the inclusion bodies were collected by centrifugation at 13,000 rpm, 60 min at 4°C and washed three times with 20mM Tris buffer pH 7.4 containing 20mM EDTA. The isolated and purified inclusion bodies were solubilized in 6M Guanidine HCl pH 7.4, followed by reduction with 65-mM DTE. Solubilized and reduced inclusion bodies were refolded by a 1:100 dilution into a redox-shuffling buffer system containing 0.1-M Tris, 0.001M EDTA, 0.5-M Arginine, and 0.09-mM Oxidized Glutathione, pH 9, and incubation at 10°C for 24 h. After having been refolded, the protein was dialyzed against 150-mM Urea, 20-mM Tris, pH 8, followed by purification of the soluble Compound A by ionexchange chromatography on a Q-Sepharose® column (7.5 mm ID 60 cm) (Sigma-Aldrich, Inc., St. Louis, Mo. USA), applying a salt (NaCl) gradient. Peak fractions containing Compound A were then subjected to buffer exchange with PBS.

Cloning of Compound B

[0114] The sHLa-A2/SS1 (sFv) was constructed as previously described by linking the C-termine of sHLa-A2 to the N-termine of the SS1 sFv via a short linker ASGG (SEQ ID NO:15). To construct the sHLa-A2/SS1 (sFv) with covalently bound MHC-restricted peptide, the MHC-restricted peptide and the peptide linker GGSGGGGGGGGGGG (SEQ ID NO:8) were fused to the N-termine of the sHLa-A2/SS1 (sFv) molecule by a PCR overlap extension reaction with the primers 5'-Nde-2092BM: 5'GGAGCGTGTTTGGCGCCATATGCAATGAGCCAGCTT CCCTCCTGTTGGGCGAGGCTCCG- GGTGCGCGAGGTTCAGGCGGCG (GAATCCGACGTAATACGACTCACTATAGG) and 3'-VLSecSS1-5'GCGATTCTTCTGTTGGGCGAGGCTCCG- GGTGCGCGAGGTTCAGGCGGCG (GAATCCGACGTAATACGACTCACTATAGG). And the 3'VLSecSS1-5'GCGATTCTTCTGTTGGGCGAGGCTCCG- GGTGCGCGAGGTTCAGGCGGCG (GAATCCGACGTAATACGACTCACTATAGG) was introduced into the 209rev/sHLa-A2/SS1 (sFv) molecule was used as a template for the construction of Compound B. In this molecule the CMV peptide NLPVMVATV (SEQ ID NO: 4) was introduced into the 209rev/sHLa-A2/SS1 (sFv) sequence (exchanging the 209 peptide) by PCR reaction using the primers 5'-GAAAGCGTGTTTGGCGCCATATGCAATGAGCCAGCTTCCCTCCTGTTGGGCGAGGCTCCG-GGTGCGCGAGGTTCAGGCGGCG (GAATCCGACGTAATACGACTCACTATAGG) and 3'-VLSecSS1-5'GCGATTCTTCTGTTGGGCGAGGCTCCG-GGTGCGCGAGGTTCAGGCGGCG (GAATCCGACGTAATACGACTCACTATAGG).
The expression and purification protocols of Compound B were identical to the expression and purification protocols of Compound A.

All the methods used to analyze the biochemical and biological properties of Compound B were identical to the methods used to analyze the activity of Compound A.

[0115] To construct the M1-cov/schLA-A2/SS1 (scFv) fusion protein the M1 58-66 peptide was fused to the N-terminus of schLA-A2/SS1 (scFv) fusion protein through a short 15 amino acid linker by overlapping PCR reaction with the 5'M1-linker primer: 5′GGAAGCGTIGGGC-GCATGCGACCTTTGATGTT-TACCCGGGCG AGGAAGCTGCTGGGCGAGGT-TCAGGACCGCTTTCTGATTGATGAGATCCAGCCTACTCAGGAA TCCGAGACTGTTGGAATCTTCATGTTTAATCTCCAAACTTTTGT3′ (SEQ ID NO: 14).

Flow Cytometry

[0118] Cells were incubated with Compound A (60 min at 4°C in 100 μl, 10 μg/ml), washed and incubated with the anti-HLA-A2 MAb BB7.2 (60 min at 4°C, 10 μl/ml). The cells were washed and incubated with anti-mouse FITC (60 min at 4°C, 10 μl/ml) that served as a secondary antibody. The cells were subsequently washed and analyzed by a FACs caliber flow cytometer (Becton-Dickinson, San Jose, Calif. USA).

Enzyme Linked Immunosorbent Assay

[0119] Immunoplates (Falcon®, Becton-Dickinson Labware, Franklin Lakes, N.J. USA) were coated with 10 μg/ml of purified recombinant mesothelin (O/N at 4°C). The plates were blocked with PBS containing 2% skim milk and then incubated with various concentrations of Compound A (60 min at RT) and washed three times with PBS. Binding was detected using the anti-HLA-conformation-dependent antibody W6/32 (60 min, RT, 1 μg/ml). The plates were washed three times with PBS and incubated with anti-mouse IgG-peroxidase (60 min, RT, 1 μg/ml). The reaction was developed using TMB (DAKO) and terminated by the addition of 100 μl H2O2. Anti-mesothelin antibody (K1) was used as a positive control. The immunoplates were analyzed by ELISA reader using 450 nm filter (Anthos 2001®, Anthos Labtech, Salzburg, Austria).

Cytotoxicity Assays

[0120] Cytotoxicity was determined by S35-methionine release assays. Target cells were cultured in culture plates in RPMI 10% FCS Methionine free for 2 h, followed by incubation overnight with 15 μc/ml of S35-methionine (NEN). The target cells were harvested by trypsinization and washed twice with 40 ml RPMI 10% FCS. The target cells were plated in 96-well plates (5·103 cells per well) in RPMI+10% FCS and incubated overnight at 37°C, 5% CO2. Target cells were incubated with different concentrations of Compound A fusion proteins for 2 h, effector CTL cells were added at different target: effector ratios and the plates were incubated for 8-12 h at 37°C, 5% CO2. Following incubation, S35-methionine release from target cells was measured in a 25 μ sample of the culture supernatant. All assays were performed in triplicate, lysis was calculated directly: ([experimental release–spontaneous release]/[maximum release–spontaneous release])·100. Spontaneous release was measured as S35-methionine released from target cells in the absence of effector cells, and maximum release was measured as S35-methionine released from target cells lysed by 0.5% NaOH.

Cell Lines

[0121] A431 and A431K5 cells (epidermoid carcinoma) were maintained in RPMI medium containing 10% FCS, L-glutamine and penicillin/streptomycin. The A431K5 cell line is a human epidermoid carcinoma A431 cell line stably transfected with Mesothelin, the transfected cells were maintained with 700 μg/ml G418 (Gibco-BRL®, Invitrogen Inc., Carlsbad, Calif. USA).

[0122] CTL’s with specificity for CMV pp 65 epitope (NLVPMVATV (SEQ ID NO:4)) were kindly provided by Dr Ditmar Zeln (Charitee, Berlin). The CTL’s were expanded by incubation with peptide pulsed, radiated (4000rad) PBMC’s from a healthy HLA-A2 positive donor and were maintained in AIMV medium+8.9% FCS+50 μM-2-mercaptoethanol+ penicillin/streptomycin 1:105 U/L.

Results:

Construction of Compound A

[0123] A construct encoding a single-chain MHC molecule composed of the β2 microglobulin gene fused to the α1, α2 and α3 of the HLA-A2 gene via a short peptide linker (15 amino acids) was fused to the scFv SS1 which targets mesothelin (FIG. 1A). This construct was analyzed in detail for its biochemical and biological activity and was found to be functional in vitro and in vivo (15). To construct a fusion protein with covalently linked peptide a 9 amino acids peptide derived from the CMV pp 65 protein (NLVPMVATV (SEQ ID NO:4)) was fused to the N-terminus of the schLA-A2/SS1 (scFv) fusion protein via 20 amino acids linker GGGGGGGSGGGGGGGGGGGS (SEQ ID NO:6) (FIG. 1B). Compound A was constructed in two steps: First a covalent fusion protein termed M1/schLA-A2/SS1 (scFv) was constructed by overlap extension PCR. In this construct the influenza M158-66 peptide GILGFVFTL (SEQ ID NO:21) and a 15 amino acid linker were fused to the N-terminus of the schLA-A2/SS1 (scFv) fusion protein. In this construct, a new, unique restriction site (BanHI) was inserted to the linker sequence by a silent mutation. In the second step PB3 plasmid containing the M1/schLA-A2/SS1 (scFv) full sequence was digested with Ndel and BamHI restriction enzymes. This digestion produced two fragments. One fragment contains the peptide and part of the linker sequence, and the second fragment contains the plasmid, part of the linker and the schLA-A2/SS1 (scFv) sequence. The fragment which contains the plasmid, part of the linker and the schLA-A2/SS1 (scFv) sequence was then ligated to dsDNA primer that codes the CMV pp 65 peptide sequence and an extension of the
linker sequence (FIG. 1B). The new plasmid was transformed to E. coli DH5α cells and positive colonies were sent to DNA sequencing (FIG. 2).

Expression and Purification of Compound A

[0124] Compound A was expressed in E. coli BL21 cells and, upon induction with isopropyl β-D-thiogalactoside, large amounts of recombinant protein accumulated in intracellular inclusion bodies. SDS/PAGE analysis of isolated and purified inclusion bodies revealed that Compound A with the correct size constituted 80-90% of the total inclusion bodies mass (FIG. 3A). The isolated solubilized inclusion bodies were reduced and refolded in vitro in a redox-shuffling buffer. Monomeric soluble fusion proteins (Compound A) were purified by ion-exchange chromatography on Q-sepharose®. SDS/PAGE analysis of Compound A revealed a highly purified monomeric molecule with the expected size of 72 KDa (FIG. 3B).

Biological Activity of the Compound A

ELISA

[0125] To test the binding ability of purified Compound A to its target antigen, the recombinant mesothelin was immobilized to immuno-plates. The binding of Compound A was monitored by using conformation sensitive mAb W6/32, this antibody recognizes MHC molecules that are folded correctly with a peptide in its groove. As shown in FIG. 4, the binding of Compound A to recombinant mesothelin was dose-dependent. This suggests that the two functional domains of Compound A, the scFv (SS1) domain and the peptide/sclHLA-A2 domain are folded correctly. Moreover, the scFv (SS1) domain of the fusion protein is in active form and can specifically bind mesothelin.

Flow Cytometry Analysis (FACS)

[0126] To test the binding ability of Compound A to mesothelin-expressing cell lines, FACS analysis was made. As a model, target cells that are HLA-A2 negative were used, thus the reactivity of an anti-HLA-A2 mAb can be used to measure the binding of Compound A to cells that express mesothelin on their surface. This model of mesothelin-positive, HLA-A2-negative cells represents the extreme case in which the tumor cells lose its HLA expression. Therefore for the FACS analysis, HLA-A2 negative A431K5 cells were used, which are human epidermoid carcinoma A431 cells that were stably transfected with mesothelin. The parental A431 human epidermoid carcinoma cells which are mesothelin-negative and HLA-A2-negative are used as negative control. The binding of Compound A to the target cells was monitored with anti-HLA-A2 mAb B87.2 as primary antibody followed by a FITC labeled secondary antibody. A mesothelin anti-mAb K1 was used to test the expression levels of mesothelin. As shown in FIG. 5A, A431K5 cells express high levels of mesothelin, whereas the parental A431 cells do not express the target antigen. The cell lines A431 and A431K5 were also tested for the expression of HLA-A2 using HLA-A2 specific antibody (BB7.2), both cell lines were HLA-A2 negative. However, when A431K5 cells were pre-incubated with Compound A, they were positively stained with the HLA-A2 specific antibody BB7.2 (FIG. 5B), Antigen-negative A413 cells were not affected. The specific binding of Compound A to A431K5 but not to A431 cells further indicates that the binding is exclusively dependent on the interaction of the targeting scFv domain of the fusion with mesothelin and that the fusion protein can bind its target antigen as natively expressed on the surface of cells.

Cytotoxicity Assay

[0127] To test the ability of Compound A to mediate the killing of HLA-A2-negative mesothelin-positive cells by HLA-A2-restricted CMV pp 65 NLPMVATV (SEQ ID NO:4) specific CTLs, S9-Methionine release assay was performed using HLA-A2-negative mesothelin-transfected A431K5 cells, and the HLA-A2-negative mesothelin-negative A431 parental cells. To determine the killing potential of the CMV specific CTLs, cytotoxicity assay was performed using HLA-A2-positive JY cells that were radiolabeled with Met52 and labeled with the CMV peptide NLPMVATV (SEQ ID NO:4). The average specific killing of the JY cells by the CMV specific CTLs was 47% with an E:T ratio of 10:1 (data not shown). As shown in FIG. 6A, Compound A effectively mediated the killing of the A431K5 cells (mesothelin-positive HLA-A2-negative). Specific killing could reach 66% in comparison to peptide-loaded JY cells. Thus, killing with the fusion protein was even more efficient compared to peptide-pulsed antigen presenting cells which represent optimal targets. However, when the target A431K5 cells were incubated with the CMV specific CTLs alone without preincubation with Compound A or when the target cells were A431 mesothelin-negative cells with or without preincubation with Compound A, no cytotoxic activity was observed. Next, a titration experiment was performed to determine the potency of Compound A, as shown in FIG. 6B, the killing of mesothelin-positive A431K5 cells was dose-dependent with an IC50 of 0.5-1 μg/ml.

[0128] These results indicate that the killing of mesothelin-positive HLA-A2-negative A431K5 cell was specific and controlled by the recognition of mesothelin by the targeting domain of Compound A (scFv/SS1) and the specificity of the CMV CTLs to the peptide/sclHLA-A2 domain.

Biological Activity of Compound B

Flow Cytometry Analysis (FACS)

[0129] To test the binding ability of Compound B to mesothelin-expressing cell lines a flow cytometry analysis was used. As a model, target cells that are HLA-A2 negative were used, thus, the reactivity of the anti-HLA-A2 mAb will indicate the binding of Compound B to the cell surface antigen. A431K5 cells which are human epidermoid carcinoma A431 cells that are stably transfected with mesothelin and are HLA-A2 negative. As controls the parental A431 human epidermoid carcinoma cells were used which are mesothelin-negative and HLA-A2 negative. The binding of Compound B to the target cells was monitored by anti-HLA-A2 mAb B87.2 as primary antibody followed by a FITC labeled secondary antibody. To test the expression levels of mesothelin and as positive control a commercial anti-mesothelin mAb K1 was used. As shown in FIG. 10A-B A431K5 cells express high levels of mesothelin, whereas the parental A431 cells do not express mesothelin. The cell lines A431 and A431K5 were also tested for their expression of HLA-A2 using HLA-A2 specific antibody (BB7.2), both cell lines were HLA-A2 negative. However, when A431K5 cells were pre-incubated with Compound B, they were positively stained with the HLA-A2 specific antibody BB7.2 whereas control A431 cells
were not stained (FIG. 10 C-D). The specific binding of Compound B to A431K5 cells but not to A431 cells indicate that binding is exclusively depended on the interaction of the targeting scFv domain with mesothelin.

[0130] To analyze the binding of Compound B in comparison to the Compound A fusion, a FACS analysis was performed using both molecules in similar conditions and concentrations. As shown in FIGS. 10E-F only the mesothelin-positive cells A431K5 were positively stained with HEL-A2-specific Ab when pre-incubated with both fusion proteins, however, Compound A exhibited better binding.

Cytotoxicity Assay

[0131] To test the ability of Compound B to mediate the killing of HEL-A2-negative mesothelin-positive cells by HEL-A2-restricted CMV NLVPMVATV (SEQ ID NO:4) specific CTLs, performed S35-Methionine release assay using HEL-A2-negative mesothelin-transfected A431K5 cells was performed, and the HEL-A2-negative mesothelin-negative A431 parental cells. To determine the killing potential of the CMV-specific CTLs cytotoxicity assay was performed, using HEL-A2-positive JY cells that were radiolabeled with MetS35 and labeled with the CMV peptide NLVPMVATV (SEQ ID NO:4). The average specific killing of the JY cells by the CMV-specific CTLs was around 45-50% using an E/T ratio of 10:1 (data not shown). As shown in FIG. 11A, Compound B effectively mediated the killing of the A431K5 cells (mesothelin-positive HEL-A2-negative), this specific killing was 66% in comparison with peptide-loaded JY cells (~150% compared to JY cells). However, when the target A431K5 cells were incubated with the CMV-specific CTLs alone without preincubation with Compound B or when the target cells were mesothelin-negative (A431 cells), with or without preincubation with Compound B, no cytotoxic activity was observed. Titration experiments which determined the potency of the fusion protein, shown in FIG. 11B indicated that the killing of mesothelin-positive A431K5 cells was dose-dependent. To compare the cytotoxic activity of Compound B and Compound A fusion proteins, which differ in the length of peptide used to covalently attach the antigenic peptide to the β2-microglobulin, a S35-Methionine release assay was performed using similar conditions for both fusion proteins. As shown in FIG. 12, both molecules efficiently and specifically mediated the killing of A431K5 cells, but not A431 cells. When relatively high concentrations of fusion proteins (Compound B and Compound A) were used, the killing activity of both molecules was similar. However, when low concentrations of fusion proteins were used the cytotoxic activity of Compound A was superior probably due to better stability and positioning of the CMV peptide in the MHC peptide-binding groove due to the longer linker.

Flow Cytometry (FACS)

[0132] The M1-cov/scHL-A2/SS1 (scFv) fusion protein was constructed by overlap extension PCR reaction in which the Influenza M1 58-66 peptide and a 15 amino acid linker GGGGSGGGGSGGGGGS were fused to the N-terminus of the scHL-A2/SS1(scFv) fusion protein (FIG. 13). The PCR product was ligated to TA-cloning vector (p-GEM, Promega), transformed to E.coli DH5α cells. Positive colonies were selected and the insert was isolated using EcoRI and NdeI. The insert was ligated to PRB expression vector and transformed to E.coli DH5α cells. Positive colonies were sent to DNA sequencing (FIG. 14).

Expression and Purification of Compound B

[0133] The M1-cov/scHL-A2/SS1 (scFv) fusion protein was expressed in E.coli BL21 cells and, upon induction with isopropyl β-D-thiogalactoside, large amounts of recombinant protein accumulated in intracellular inclusion bodies. SDS-PAGE analysis of isolated and purified inclusion bodies revealed that the M1-cov/scHL-A2/SS1 (scFv) fusion protein with the correct size constituted 80-90% of the total inclusion bodies mass (FIG. 15A). The isolated solubilized inclusion bodies were mesothelin-transfected in vitro with a reduct-shuffling buffer. Monomeric soluble fusion proteins (M1-cov/scHL-A2/SS1 (scFv)) were purified by ion-exchange chromatography on Q-Sepharose®. SDS-PAGE analysis of the M1-cov/scHL-A2/SS1 (scFv) fusion proteins revealed a highly purified monomeric molecule with the expected size of 72 KDa (FIG. 15B).

ELISA

[0134] To test the binding ability of the purified M1-cov/scHL-A2/SS1 (scFv) fusion protein to its target antigen, recombinant mesothelin was immobilized to immunoplates. The binding of M1-cov/scHL-A2/SS1 (scFv) fusion protein was monitored by using conformation sensitive mAb W6/32, this anti body recognizes MHC molecules that are folded correctly with a peptide in its groove. As shown in FIG. 16 the binding of the M1-cov/scHL-A2/SS1 (scFv) fusion protein to recombinant mesothelin was dose-dependent. This suggests that the two functional domains of M1-cov/scHL-A2/SS1 (scFv) fusion protein, the scFv (SS1) domain and the M1-cov/scHL-A2 domain are folded correctly. Moreover, the scFv (SS1) domain of the fusion protein is in active form and can specifically bind mesothelin.
indicates that the binding is exclusively dependent on the interaction of the targeting domain (scFv(SS1)) with mesothelin.

Cytotoxicity Assay

To test the ability of the M1-cov/scHLA-A2/SS1 (scFv) fusion protein to mediate the killing of HLA-A2-negative mesothelin-positive cells by HLA-A2-restrictive M158-66 specific CTLs, S55-Methionine release assay using HLA-A2-negative mesothelin-transfected A431K5 cells was performed. As shown in FIG. 18, M1-cov/scHLA-A2/SS1 (scFv) fusion protein did not mediate the lysis of A431K5 cells (mesothelin-positive HLA-A2-negative). However, the scHLA-A2/SS1(scFv) bearing the M158-66 peptide in its groove mediated the killing of mesothelin-positive target cells by the HLA-A2-restricted M158-66 specific CTLs.

Discussion:

This study demonstrates the ability to target covalently linked peptide/scMHC/scFv fusion protein to tumor cells that can render HLA-A2-negative cells susceptible to lysis by the relevant HLA-A2-restricted CTLs. As previously shown by Lev et al., and Oved et al. (15,21), this strategy has two major advantages. First, it takes advantage of the use of recombinant Ab fragments that can localize on those malignant cells that express a tumor marker, usually associated with the transformed phenotype (such as growth factor receptors and/or differentiation antigens), with a relatively high degree of specificity. Second, this strategy has the ability to recruit a particular population of highly reactive cytotoxic T-cells specific to a preselected, highly antigenic peptide epitope present in the targeted MHC-peptide complex, such as viral-specific T-cell epitopes. This platform approach generates multiple molecules with many tumor-specific scFv fragments that target various tumor specific antigens, combined with the ability to target many types of MHC-peptide complexes carrying single, preselected, and highly antigenic peptides derived from tumor, viral, or bacterial T-cell epitopes. These examples present a strategy one step further by fusing the 9 amino acid peptide linked by a short linker (20AA) to the previously reported scHLA-A2/scFv fusion protein (15AA), and by so doing, stabilizing the peptide in the MHC groove prolonging the general stability of the fusion proteins. As a model for this new generation of fusion proteins, the present invention relates to construction of a fusion protein in which the CMV pp 65 derived (NLVPVMATV) fused to the N-terminus of the scHLA-A2/SS1 (scFv) molecule and its biochemical and biological characteristics. It is shown that the two domains of the new fusion protein can refold in vitro to form correctly folded molecules with the peptide within the HLA-A2 groove and an active targeting domain (scFv) that can specifically bind its target antigen. Moreover, this fusion protein had successfully mediated the lysis of HLA-A2-negative mesothelin-positive tumor cells by HLA-A2-restricted CTLs.

Tumor progression is often associated with the secretion of immune-suppressive factors and/or the down-regulation of MHC class I antigen-presentation functions (2). Even when a specific CTL response is demonstrated in patients, this response is low because the anti-tumor CTL population is rare, very infrequent, and in some cases the CTLs are not functional or anergic (26). Moreover, it is well-established that the number of MHC-peptide complexes on the surface of tumor cells that present a particular tumor-associated peptide is low (27). As shown herein, the new strategy overcome these problems. First, the tumor cells are coated with MHC-peptide complexes independent of their endogenous MHC expression. Second, the use of tumor specific antigens that are usually part of the tumor phenotype (such as growth factor receptors and differentiation antigens) prevent the down regulation of those antigens and prolong the efficiency of the treatment. Third and most important, the effector domain of the fusion protein the MHC-peptide complex can recruit specific populations of CTLs depending on the peptide harboring the MHC groove.

It is appreciated that certain features of the invention, which are, for clarity, described in the context of separate embodiments, may also be provided in combination in a single embodiment. Conversely, various features of the invention, which are, for brevity, described in the context of a single embodiment, may also be provided separately or in any suitable subcombination. Although the invention has been described in conjunction with specific embodiments thereof, it is evident that many alternatives, modifications and variations will be apparent to those skilled in the art. Accordingly, it is intended to embrace all such alternatives, modifications and variations that fall within the spirit and broad scope of the claims. All publications, patents and patent applications mentioned in this specification are herein incorporated in their entirety by reference into the specification, to the same extent as if each individual publication, patent or patent application was specifically and individually indicated to be incorporated herein by reference. In addition, citation or identification of any reference in this application shall not be construed as an admission that such reference is available as prior art to the present invention.

REFERENCES CITED


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met asp leu leu ser leu thr ser glu asp ser ala val tyr phe cys
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What is claimed is:

1. A fusion protein comprising consecutive amino acids which, beginning at the amino terminus of the protein, correspond to consecutive amino acids present in (i) a cytomegalovirus human MHC-restricted peptide, (ii) a first peptide linker, (iii) a human β-2 microglobulin, (iv) a second peptide linker, (v) a HLA-A2 chain of a human MHC class I molecule, (vi) a third peptide linker, (vii) a variable region from a heavy chain of a scFv fragment of an antibody, and (viii) a variable region from a light chain of such scFv fragment, wherein the consecutive amino acids which correspond to (vii) and (viii) are bound together directly by a peptide bond or by consecutive amino acids which correspond to a fourth peptide linker and the scFv fragment is derived from an antibody which specifically binds to mesothelin.

2. The fusion protein of claim 1, wherein the first peptide linker has the amino acid sequence GGGGSGGGGSGGGGSGGGGS (SEQ ID NO:6).
3. The fusion protein of claim 1, wherein the second peptide linker has the amino acid sequence GGGGSGGGGSGGGGS (SEQ ID NO:8).

4. The fusion protein of claim 1, wherein the third peptide linker has the amino acid sequence ASGG (SEQ ID NO:10).

5. The fusion protein of claim 1, wherein the fourth peptide linker has the amino acid sequence GVOGSGGGGSGGGGS (SEQ ID NO:19).

6. The fusion protein of claim 1, wherein the cytomegalovirus human MHC-restricted peptide has the amino acid sequence NLVPMVATV (SEQ ID NO:4).

7. The fusion protein of claim 1, wherein the sequence of the consecutive amino acids corresponding to (vii), followed by the fourth peptide linker, followed by (viii) is set forth in SEQ ID NO:12.

8. The fusion protein of claim 1, wherein the consecutive amino acids have the amino acid sequence set forth in SEQ ID NO:2.

9. A composition comprising the fusion protein of claim 1 and a carrier.

10. The composition of claim 9 wherein the fusion protein is present in the composition in a therapeutically effective amount and the carrier is a pharmaceutically acceptable carrier.

11. A nucleic acid construct comprising a nucleic acid sequence encoding the fusion protein of claim 1.

12. The nucleic acid construct of claim 11, wherein said nucleic acid sequence is as set forth in SEQ ID NO:1.

13. A vector comprising the nucleic acid construct of claim 11.

14. An expression vector comprising the nucleic acid construct of claim 11 and a promoter operatively linked thereto.

15. A transformed cell comprising the vector of claim 14.

16. An isolated preparation of bacterially-expressed inclusion bodies comprising over 30 percent by weight of the fusion protein of claim 1.

17. A process for producing a fusion protein comprising culturing the transformed cell of claim 15, so that the fusion protein is expressed, and recovering the fusion protein so expressed.

18. A method of killing a tumor cell which expresses mesothelin on its surface, the method comprising contacting the tumor cell with the fusion protein of claim 1 in an amount effective to initiate a CTL-mediated immune response against the tumor cell so as to thereby kill the tumor cell.

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