**Selection of novel peptide mimics of the GD2 ganglioside from a constrained phage-displayed peptide library**

IRENA HORWACIK1, DOMINIK CZAPLICKI1, KATARZYNA TALAREK1, ALEKSANDRA KOWALCZYK1, ELZBIETA BOLESTA2, DANUTA KOZBOR2 and HANNA ROKITA1

1Faculty of Biochemistry, Biophysics and Biotechnology, The Jagiellonian University, 7 Gronostajowa St., 30-387 Krakow, Poland; 2Department of Immunology, Roswell Park Cancer Institute, Elm and Carlton Streets, Buffalo, NY 14263, USA

Received November 20, 2006; Accepted December 27, 2006

**Abstract.** Aberrant glycosylation is a universal feature of cancer cells. There are quantitative and qualitative changes in expression of gangliosides observed in tumors of a neuro-ectodermal origin such as neuroblastoma, melanoma and astrocytoma. The presence of large amounts of GD2 ganglioside on neuroblastoma cells, as compared to normal cells, opens the possibilities to use the tumor-associated carbohydrate antigen in diagnosis and immunotherapeutic approaches. In the quest for immunogens potentially capable of eliciting anti-GD2 ganglioside immune responses, we performed affinity purification of phage-displayed peptides from the LX-8 library (12-mer containing disulphide bridge). The library was screened with the biotinylated anti-GD2 ganglioside 14G2a mAb monoclonal antibody. Our goal was to isolate and characterize peptide mimics of GD2 ganglioside. Numerous individual phage clones that bound 14G2a mAb were identified with the application of immunoblotting technique in the phage pools yielded from the pannings. The phage-borne peptides were tested for their anti-GD2 ganglioside antibody binding ability using ELISA. Among these clones five different phage-displayed peptide sequences were identified. Moreover, we showed that the secondary structure of the peptides, stabilized by the disulfide bridging between cysteine residues at positions 2 and 11, was crucial for the binding of the peptides to 14G2a mAb. In a separate set of experiments, we observed a competition of the peptides, expressed on phages as well as in their synthetic form, with the nominal antigen GD2 ganglioside expressed on IMR-32 neuroblastoma cells for binding to 14G2a mAb. Based on the obtained results we concluded that all of these 5 peptides were mimics of the GD2 ganglioside.

**Introduction**

GD2 ganglioside belongs to a group of sialylated glycosphingolipids. The molecule has an amphipatic character and consists of an outer charged glycan portion with two sialic acid residues and a ceramide unit, fatty acyl sphingosine. Due to its hydrophobic nature, the sphingolipid tail embeds the molecule in the outer leaflet of the plasma membrane. Changes in ganglioside content (e.g. presence of O-acetylated terminal sialic acid residues at the position 7 or 9 of the glycan part, ceramide modifications) are a hallmark of certain types of tumors (1). Due to incomplete synthesis, precursors such as GM2, GM3, GD3 and GD2 gangliosides are accumulated in neuroblastoma, melanoma, small cell lung carcinoma, sarcoma, or B cell lymphoma (2). With reference to a vast number of research data, gangliosides have been related to tumorigenesis (3). An alteration in the ganglioside expression pattern has been linked to biological phenotype and clinical behavior that can be used to predict the prognosis of neuroblastoma patients (4,5). As for the GD2 ganglioside, high circulating levels of the tumor carbohydrate antigen were strongly related to more rapid disease progression and lower survival rates of advanced risk neuroblastoma patients (6). Additional experimental data from culture systems suggest that GD2 may be one of the modulating factors in tumor-host interactions. In particular, SH-SY-5Y neuroblastoma cells stably transfected with a construct encoding cDNA of GD3 synthase exhibited increased levels of both GD2 and GD3 gangliosides. This was accompanied by morphological changes and the tendency to form aggregates of the cells in cultures (7). Yoshida and colleagues showed that in the case of small cell lung cancer cells SK-LC-17, the observed rise in GD2 and GD3 ganglioside content led to an increase in proliferation and invasiveness. Moreover, the anti-GD2 monoclonal antibodies (220-51) were capable of diminishing the proliferation rates of the tumor cell line, in a mechanism that was shown to involve apoptosis. This indirect evidence indicates that the GD2 ganglioside may be involved in tumorigenesis (8). Another widely described phenomenon that may be added to the biological functions of gangliosides is suppression of immune cells by intact ganglioside molecules present on the tumor cells or shed into their microenvironment (9).
Many anti-GD2 ganglioside monoclonal antibodies have been identified (10). Overexpression of GD2 ganglioside distinguishes neuroblastoma from benign neural tumors (11), which has resulted in the application of antibodies targeting the carbohydrate in the diagnosis of localized and metastatic neuroblastoma and in the monitoring of therapy response and minimal residual disease (12,13). The fact that approximately 60% of children with neuroblastoma, which is the most common extracranial neoplasm of childhood, have high-risk tumors that can rarely be cured by conventional therapy, stresses the need for new treatment protocols to control minimal residual disease. As GD2 ganglioside is expressed strongly and quite uniformly on neuroblastoma cells and exhibits a restricted localization pattern in normal tissues, the tumor-associated carbohydrate antigen represents a good therapeutic target. Furthermore, GD2 ganglioside is not downregulated when bound by specific antibodies (14). Among the anti-GD2 ganglioside monoclonal antibodies available, two mouse antibodies 3F8 (IgG3), 14G2a (IgG2a) and one chimeric ch14.18 antibody were tested in clinical trials for their anti-tumor activity against neuroblastoma (15). Additionally, the etiology and immunogenicity of neuroblastoma encourage development of active immunotherapies. However, if such therapies are to be successful, they have to overcome self-tolerance to GD2 ganglioside and poor immunogenicity of the antigen in humans. Unlike peptides, most carbohydrates (due to their chemical nature) are thymus-independent antigens. They are not presented in the context of MHC molecules, stimulate primary B-cell responses, and do not require T cell help. They mainly induce short-lived IgM antibodies (10). Moreover, carbohydrate antigens are difficult and expensive to synthesize because of chirality and low chemical yield. Currently, clinically tested strategies to improve GD2 ganglioside immunogenicity include application of its conjugates with KLH (keyhole limpet hemocyanin) protein carrier and vaccination with substitute immunogens in the form of anti-idiotypic antibodies (in various adjuvant settings) (16).

Another strategy that relies on surrogate antigens of GD2 gangliosides is the application of mimicking peptides, which can be isolated from phage-displayed peptide libraries. The filamentous bacteriophages infecting E. coli, of which M13, f1 and fd have been the most extensively studied, are a group of viruses that contain a circular single-stranded DNA genome encased in a long protein capsid. The filamentous phage structure and its mode of replication have made it a valuable tool for biological research. Due to the membrane-associated assembly process of the viruses, it is possible to display foreign peptides and even proteins on the surface of the phage particles, while the genetic material encoding each variant resides inside it. Therefore, by using a receptor, one can apply various in vitro selection techniques to purify binding phage clones from such libraries and analyze their interactions with the screening molecule. The phage display technique has numerous modifications and has been broadly applied, for example, in the production of recombinant antibodies, in the analysis of the interactions among proteins and their domains, in epitope mapping, and in binding motif and small ligand identification (17). One type of such libraries are peptide libraries, where short peptide sequences; random, linear, constrained or even expressed as a fragment of a larger protein scaffold; are expressed as fusion products on the surface of the virions. In many cases, where an antibody was used in screening, the isolated peptides were shown to interact with the antigen-binding site of antibodies recognizing native antigens in the form of proteins, carbohydrates, or nucleic acids. In addition, such peptides compete with the native antigen for binding to the antigen-binding site and therefore are named as their mimics (18,19). Mimetic peptides have been successfully tested in animal models and induce immune responses to carbohydrate compounds present on pathogens and tumor cells (20,21).

In the quest for antigens potentially capable of eliciting anti-GD2 ganglioside immune responses, we performed affinity purification of phage-displayed peptides from the LX-8 library (12-mer containing disulphide bridge) with biotinylated anti-GD2 ganglioside 14G2a mAb. We isolated and characterized five peptide mimics of GD2 ganglioside. We showed that the spatial structure of the isolated phage-displayed peptides was essential for their mimicking potential. We observed competition of the mimicking peptides with GD2 ganglioside, expressed on IMR-32 neuroblastoma cells, for the antigen-binding site of 14G2a mAb.

**Materials and methods**

*Materials.* The LX-8 phage-displayed peptide library was produced in the vector f88-4, as a fd-tet derivative (Entrpz Nucleotide accession no. AF218363). The library encoded 12 amino acid (aa) peptides fused to the N-terminus of the major coat protein pVIII. There were 2 fixed Cys residues (C) introduced in the 2nd and the 11th position of the encoded peptides (XCX,CX, random amino acids (aa) are marked as X). The peptides were followed by AAEGDD (22). The diversity of the library was 1.5x10^6 phage clones. K91Kan (K91) E. coli strain was the bacterial host for filamentous phages in this study. Its sex is Hfr Cavalli, and therefore deploys the F pilus, the attachment site for filamentous phage infection. The mini-kan hopper element was inserted in the lacZ gene using the vector βN1K1105 that confers kanamycin resistance on the cells (23). The hybridoma cell line secreting the mouse monoclonal antibody 14G2a was provided by Dr R. A. Reisfeld (The Scripps Institute, La Jolla, CA, USA). The antibody is of the IgG2a isotype, and binds to GD2 ganglioside (24). For all experiments, the antibody was purified from cell culture media using HI TRAP protein G columns according to the manufacturer's protocol (Amersham Biosciences AB, Uppsala, Sweden). The biotinylation of 14G2a mAb was performed using sulfo-NHS-LC biotin (Sigma, St. Louis, MO, USA) according to the protocol described in (25). Panning experiments and ELISAs were performed in 96-well microplates (MaxiSorp, NUNC A/S, Roskilde, Denmark). A horseradish peroxidase (HRP)-conjugated, anti-pVIII monoclonal antibody, developed in the mouse, was obtained from Amersham Biosciences. HRP-conjugated anti-mouse Ig polyclonal antibodies, developed in the rabbit, were purchased from Sigma. FITC-conjugated F(ab)2 fragments raised against murine IgG, IgM and IgA, developed in the goat, were from MP Biomedicals, Inc.
(Aurora, OH, USA). A human neuroblastoma cell line IMR-32 (CCL-127, ATCC) (26), expressing high levels of GD2 ganglioside (27), was cultured using E-MEM: 1 mM sodium pyruvate, 1% non-essential amino acid solution (all from Sigma), 10% FBS, 100 μg/ml penicillin, 0.25 μg/ml streptomycin and 0.085% amphotericin B (Invitrogen Ltd., Paisley, UK) at 37°C in a humidified 5% CO2 incubator. Synthetic peptides were from Sigma Genosys (The Woodlands, TX, USA). All other chemicals were purchased form Sigma unless stated otherwise.

Screening of the LX-8 library with 14G2a mAb. The major steps describing the panning experiments are described below. Phage clones binding to 14G2a mAb were isolated during 4 rounds of an ‘in-solution’ panning (phage clones #8, #65, #85 and #94) and 6 rounds of a ‘solid’ phase panning (a phage clone #D). For both types of pannings, in their first rounds 1x10⁹ phage particles (ph) (6.7x10⁹ equivalents of the library) per well were screened. To remove nonspecific binders, the library, diluted in TBS containing 2% BSA and 0.5% Tween-20 (TBS: 50 mM Tris-Cl, pH 7.5, 150 mM NaCl and bovine serum albumin (BSA) (Amresco Inc. Solon, OH, USA)] was preadsorbed for 4 h at room temperature (RT) in wells previously coated with streptavidin (SA, 1 μg per well in TBS), blocked with 2% BSA in TBS, and then washed 3 times with 0.5% Tween-20 in TBS.

During the ‘in-solution’ panning the non-adsorbed phages were removed and incubated with the bionitylated 14G2a mAb at 1-, 3-, 10- and 30-nM concentrations overnight (O/N) at 4°C. Phage-antibody complexes were isolated on the SA-coated, BSA-blocked microplates for 15 min at RT with rocking. The reaction was stopped by the addition of TBS containing 10 nmol of biotin- and 2% BSA, for the last 5 min of incubation. Non-bound phages were removed during 6 washes with TBS containing 0.5% Tween-20. LB (30 μl) was added to each well, and infection of the binders with ~5x10⁷ cells of starved K91 in 10 μl of NAP buffer (80 mM NaCl and 50 mM NH₄H₂PO₄, pH 7.0) was carried out for 15 min at RT. Then, the tetracycline resistance genes were induced by the addition of tetracycline (tet) in LB to a final concentration of 0.2 μg/ml with shaking (45 min, 37°C, 250 rpm). In the next step, the antibiotic concentration was increased to a bacteriostatic level (15 μg/ml), while simultaneously the tac promoter, controlling the expression of the recombinant gene VIII, was induced by the addition of IPTG to a final concentration of 1 mM (Amresco). The amplification step was carried out with gentle shaking in a humidified container (42 h, 37°C). To collect the amplified phages, media were transferred to 1.5-ml Eppendorf tubes and centrifuged for 1 h at 1500 x g at 4°C. Supernatants were transferred to fresh tubes. The remaining bacteria were heat-killed (20 min, 70°C), and then removed by centrifugation (as above). The second, third and fourth rounds of panning were performed with the following modification; 40 μl of medium with amplified phages was used in these rounds of panning, and the incubation time of phage-antibody complexes in the SA-coated wells was decreased to 10 min. During the fourth round of panning, to increase the possibility of finding high-binding clones, the concentration of the bionitylated 14G2a mAb was decreased. The phage pools incubated in the preceding rounds with 10- and 30-nM antibody concentrations were panned using 14G2a mAb at 1- and 3-nM concentrations.

The ‘solid phase’ panning was performed as described above with the following modifications; the SA-coated, BSA-blocked wells were saturated with 35 μl of 100 nM bionitylated 14G2a mAb (20 min, RT). The remaining SA sites were blocked with 10 nmol of biotin (5 min, RT). Then, the wells were washed with TBS containing 0.5% Tween-20 (3 times), and the preadsorbed library was incubated in the wells (2 h, RT) with rocking. Prior to the infection of the K91 cells, the phages were eluted with 35 μl of a buffer containing 0.1 M HCl (pH 2.2 with glycine), 0.1% BSA (10 min, RT), and then neutralized with 7 μl of 1 M Tris-HCl (pH 9.0).

For both types of screening experiments, after each round, titering was performed to determine the efficiency of infection and percentage yield of the bound phages. The amplification yield was determined with agarose gel electrophoresis of the viral ssDNA. ELISAs were performed to estimate the enrichment of 14G2a binding clones in the amplified phage pools (28).

Immunoblotting. The first series of immunoblotting experiments was performed to isolate the 14G2a binding phage clones after the panning experiments. Phage particles (10¹⁰) in TBS containing 0.1% gelatin (from pools amplified during the last rounds of the pannings) were used to infect K91 cells (~10⁹ cells in LB) for 10 min at RT. After infection and tetracycline resistance gene induction (as above), bacteria were grown in the presence of 20 μg/ml tet (1 h, 250 rpm, 37°C), and then titered. In the next step, ~200 bacterial colonies were grown on NZY/agar plates with 100 μg/ml kanamycin (kan) and 20 μg/ml tet (O/N, 37°C). Replicas of the bacteria were prepared on nitrocellulose membranes (Schleicher and Schuller BioScience Inc., Dassel, Germany). The membranes with the transferred bacterial colonies were washed 3 times in TNT buffer (10 mM Tris-HCl, pH 7.5, 150 mM NaCl and 0.05% Tween-20), and then blocked with TNT containing 5% non-fat dry milk (Gostynin, Poland) with rocking (1 h, RT). Then, the membranes were incubated with 2 μg/ml of the 14G2a mAb dissolved in TNT containing 1% BSA (O/N, 4°C). The membranes were washed 3 times with TNT containing 0.1% BSA and incubated with the anti-mouse Ig antibodies, diluted 1:150,000 in TNT containing 1% BSA (1 h, RT, with rocking). After 3 washings (as above), the signal was developed with ECL Plus Western blot detection reagent (Amersham BioSciences, San Diego, CA, USA).

Finally, the membranes were exposed to film (CEA AB, Strängnäs, Sweden), and developed (Foma, Bohemia Ltd, Hradec Králové, Czech Republic). Selected bacterial colonies with phages binding to 14G2a mAb were grown in 2 ml of LB with 100 μg/ml kan and 20 μg/ml tet (4 h, 37°C) with shaking (250 rpm). The phage amplification yield was estimated by gel electrophoresis and titering. The second round of immunoblotting experiments was performed on the amplified phage clones to confirm their binding to 14G2a mAb.

Purification of phage particles by polyethylene glycol precipitation. Prior to ELISAs, sequencing and competition tests, phage clones were amplified and purified by double precipitation with polyethylene glycol (PEG). A small-scale
amplification of a single phage-infected bacterial colony was carried out in 2.5 ml of LB containing 15 μg/ml tet (O/N, 37°C, with shaking for 250 rpm). The next day, the media were centrifuged (20 min, 4°C, 1300 x g). Phages were precipitated twice by the addition of a 0.15 volume of PEG/NaCl solution (16.7% polyethylene glycol-8000; 3.3 M NaCl) to the supernatants, followed by thorough mixing and incubation on ice (4 h for the first incubation, O/N for the second incubation). For both precipitations phage particles were recovered by centrifugation (40 min, 4°C, 10,000 x g), pellets were dissolved in TBS followed by incubation on ice (1 h), and the non-dissolved material was discarded after the final centrifugation (1 min, 4°C, 16,100 x g).

For large scale amplification of selected phage clones, 10⁵ ph in 1 ml of 0.1% gelatin/TBS were mixed with an equal volume of K91 cells (5x10⁹) in a 50-ml tube (10 min, RT). The tetracycline resistance genes were induced (as above), and the samples were incubated with shaking (30 min, 37°C, 250 rpm). Then the tet concentration was increased to 20 μg/ml, and the incubation was continued (O/N). On the following day, the media were centrifuged (10 min, 4°C, 6,200 x g), and PEG/NaCl was added (0.15 volume), followed by thorough mixing and an incubation step (O/N, 4°C). To recover the precipitated phage particles, the samples were centrifuged for 40 min at 4°C at 6,200 x g. The phage pellets were dissolved in 1 ml of TBS, and incubated (30 min, 37°C) with gentle shaking. Then, the samples were transferred to 2-ml tubes, and the non-dissolved material was removed by centrifugation (10 min, 4°C, 14,500 x g). Again, the supernatants were transferred to fresh tubes, and the second phage precipitation was performed on ice (3 h), followed by centrifugation (15 min, 4°C, 16,100 x g). The viral pellets were dissolved in 1 ml of TBS, and then incubated on ice (1 h). Finally, the non-dissolved material was removed by centrifugation (10 min, 4°C, 16,100 x g), and the supernatants were transferred to fresh tubes.

The phage concentrations were measured by scanning diluted samples from 240 to 320 nm using a UV spectrophotometer (Pharmacia BioSciences Biochrom Ltd., Cambridge, UK) and gel electrophoresis along with samples of known concentrations.

**Phage ELISAs under reducing conditions** were performed to determine the role in the peptide binding played by the disulfide bridging between peptide cysteine residues in the positions 2 and 11. The phage clones representing 5 different peptide sequences (phage #8, #85, #94, #65 and #D) and a phage #0 (a phage clone which did not bind to the 14G2a mAb in the immunoblotting experiments) were dissolved in TBS, and immobilized in microplates (O/N, 4°C). The next day, the non-bound phages were removed, and the wells were blocked with TBS containing 2% BSA (30 min, 37°C). Then, the wells were washed 3 times with TBS containing 0.5% Tween-20. Modification of the peptide disulphide bonds was introduced in two steps (both performed for 30 min at 4°C). In the first step, the appropriate wells were subjected to reducing conditions through the addition of 35 μl of 100 mM dithiothreitol in TBS (DTT+/NEM-) or 35 μl of 15 mM DTT, 100 mM NEM in TBS (DDT+/NEM+). In the second step, cysteine-blocking was introduced through the addition of 35 μl of 100 mM N-ethylmaleimide in TBS (DTT+/NEM+) or 35 μl of 15 mM DTT, 100 mM NEM in TBS (DDT+/NEM+). Controls were used including phage clones treated with 35 μl of TBS (DDT+/NEM-), BSA conjugate of non-constrained #47 peptide (29) or BSA alone. Then, the contents of the wells were removed, and 35 μl of the 14G2a mAb (1 μg/ml) in TBS containing 5 mM DTT, 2% BSA and 0.1% Tween-20 was added to the DDT+/NEM- and DDT+/NEM+ wells, while the mAb solution without DTT was applied to the DDT+/NEM- and DDT+/NEM+ wells. Following 1 h of incubation (RT, with rocking), and a washing step (6 times with TBS containing 0.5% Tween-20), the HRP-conjugated anti-mouse Ig antibodies were used to detect the bound 14G2a mAb (1 h, 37°C). The secondary antibodies were diluted 1:4,000 in TBS containing 2% BSA and 0.1% Tween-20. After the final washing step (as above), the signal was developed with the TMB substrate, and read with a spectrophotometer at 450 nm. Each determination was performed in duplicate.

**Sequencing.** The phage clones were sequenced using the primer 5' GCCAATAGTAGCACCACCAAGATA 3' (DNA Sequencing and Oligonucleotide Synthesis Facility, Institute of Biochemistry and Biophysics, Polish Academy of Science, Warsaw, Poland), and the sequences of the peptide coding region of the recombinant fusion pVIII gene protein were analyzed using Chromas (version 1.45). Peptide sequence analysis was performed with ClustalW, a command line interface of Clustal. Clustal is a general purpose multiple sequence alignment program for DNA and proteins. An amino acid identity matrix was used to generate the alignment and a branching diagram for the five isolated peptide sequences (30).

**Competition tests.** Phage particles (10¹²) of the clones expressing the 5 different peptides (and the phage #0) were diluted in PBS containing 2% BSA, and then incubated with 200 ng of 14G2a mAb (O/N, 4°C). The next day the samples were mixed with 1x10⁶ IMR-32 neuroblastoma cells and incubated for 45 min on ice. Next, the samples were washed twice with ice-cold PBS containing 2% FCS and 0.1% NaN₃, followed by centrifugation (300 x g for 5 min at 4°C). The binding of 14G2a mAb to the neuroblastoma cells was
detected with the FITC-conjugated (Fab')2 fragments binding mouse IgG, IgM and IgA, dissolved 1:40 in PBS containing 2% FCS and 0.1% NaN3 (45 min, 4˚C). After the washing step, the cells were analyzed using a flow cytometer (FACSScan, BD Biosciences, Erembodegen, Belgium). Based on the signal from negative controls (without 14G2a mAb), the pools of cells that bound 14G2a mAb were determined. The fluorescence signal in the samples (mean fluorescence intensity, MFI) was extracted from the data, and % of inhibition of binding of 14G2a mAb to the IMR-32 cells by each phage clone was calculated using the formula % inhibition = (1 - MFIpeptide/MFImax) x 100%. MFIpeptide was the mean fluorescence intensity value of samples incubated with a given peptide-expressing phage clone, and MFImax was the mean fluorescence intensity value of samples incubated with the phage clone #0. The measurements were performed in duplicate, and the experiments were repeated three times.

Synthetic peptides in dilutions ranging from 0.1-1000 μM were incubated with 200 ng of 14G2a mAb in PBS containing 2% BSA (O/N, 4˚C), then used in a competition assay on IMR-32 neuroblastoma cells, and analyzed using flow cytometry technique (as described above). A sample with non-binding peptide #Con (HEDIISLWDQSLK), a peptide from HIV-1 envelope protein was included as a specificity control (31). Based on the signal from negative controls (without 14G2a mAb), the pools of cells that bound 14G2a mAb were determined. The fluorescence signal in the samples (MFI) was extracted from the data, and % of inhibition of binding of 14G2a mAb to the IMR-32 cells by each synthetic peptide was calculated using the formula % inhibition = (1 - MFIpeptide/MFImax) x 100%. MFIpeptide was the mean fluorescence intensity value of samples incubated with a given concentration of the synthetic peptide tested, and MFImax was the mean fluorescence intensity value of samples incubated with 14G2a mAb alone. The IC50 was defined as the concentration of the given peptide that reduced the binding of 14G2a mAb to IMR-32 cells by 50%. The mean IC50 value was calculated based on the results from three independent experiments.

Results
Isolation of phage-displayed peptides binding the anti-GD2 ganglioside antibody 14G2a. We performed biopanning experiments of the LX-8 phage display peptide library, encoding constrained 12 aa peptides, using the anti-GD2 ganglioside mouse monoclonal antibody 14G2a. Two types of panning experiments were designed. One was carried out as a 4-round ‘in-solution’ biopanning, in which the phage-displayed peptide library was incubated with 30-, 10-, 3- and 1-nM concentrations of the screening antibodies for the first 3 rounds, and the 3- and 1-nM 14G2a mAb during the final round. The second experiment was performed as a ‘solid phase’ biopanning experiment (6 rounds) in microplate wells coated with the 100-nM biotinylated-screening molecule immobilized via streptavidin. Both protocols resulted in isolation and amplification of pools of phages enriched for 14G2a binders, as verified by ELISA performed after each round (data not shown).

Immunoblotting, applied in the next step, allowed us to verify the presence of the 14G2a binding phages in the pools generated after both types of panning. Replicas of bacteria infected with the phages on nitrocellulose membranes were screened with 14G2a mAb. Then, the binding of the antibody was detected with the secondary HRP-conjugated antibodies, and the membranes were developed with electrochemiluminescence technique. The experiments confirmed that the phage pools obtained from the biopanning experiments were indeed enriched in the 14G2a-binding clones. Furthermore, analyses of the results led to the identification of numerous phage-infected bacterial colonies, whose replicas yielded a strong signal from the binding of 14G2a mAb (Fig. 1A). We chose 60 colonies that were subsequently re-screened in the
second round of immunoblotting (Fig. 1B). Fifty-five of the phage clones again yielded a strong positive signal of 14G2a mAb binding, as compared to a randomly chosen phage clone #0. The phage #0 was picked in the first round of immunoblotting and used as a specificity control since it did not exhibit binding to the antibody (data not shown).

Determination of the phage-displayed peptide sequences. To further characterize and determine the relative affinity of the isolated phage clones to the GD2-specific antibody, we performed ELISA on all of the 55 isolated clones. Amplified and PEG-purified viral particles (10^{10}) were captured via their displayed peptides by the biotinylated 14G2a mAb immobilized to streptavidin, and then were detected with the HRP-conjugated anti-phage pVIII antibody. Each value was a ratio of the mean values of OD 450-nm readings of the respective phage clone and the phage clone #0.

Table I. The phage clones isolated with immunoblotting and their binding reactivity to 14G2a mAb as determined with ELISA.

<table>
<thead>
<tr>
<th>Phage clone</th>
<th>Fold rise of ELISA value</th>
<th>Phage clone</th>
<th>Fold rise of ELISA value</th>
<th>Phage clone</th>
<th>Fold rise of ELISA value</th>
</tr>
</thead>
<tbody>
<tr>
<td>1-b</td>
<td>1.90</td>
<td>25-a</td>
<td>4.90</td>
<td>85-ct</td>
<td>17.70</td>
</tr>
<tr>
<td>2-b</td>
<td>1.80</td>
<td>26-c</td>
<td>5.00</td>
<td>86-ct</td>
<td>11.45</td>
</tr>
<tr>
<td>3-a</td>
<td>5.60</td>
<td>27-c</td>
<td>1.20</td>
<td>94-at</td>
<td>3.90</td>
</tr>
<tr>
<td>4-a</td>
<td>4.30</td>
<td>28-e</td>
<td>1.20</td>
<td>B-2</td>
<td>13.40</td>
</tr>
<tr>
<td>5-a</td>
<td>1.20</td>
<td>2-bt</td>
<td>2.30</td>
<td>B-4</td>
<td>8.40</td>
</tr>
<tr>
<td>6-a</td>
<td>0.90</td>
<td>4-bt</td>
<td>5.20</td>
<td>F-1</td>
<td>12.60</td>
</tr>
<tr>
<td>7-c</td>
<td>4.00</td>
<td>6-bt</td>
<td>2.10</td>
<td>F-2</td>
<td>12.10</td>
</tr>
<tr>
<td>8-a</td>
<td>1.10</td>
<td>8-bt</td>
<td>11.20</td>
<td>F-4</td>
<td>13.00</td>
</tr>
<tr>
<td>9-c</td>
<td>1.70</td>
<td>15-at</td>
<td>3.20</td>
<td>F-5'</td>
<td>14.20</td>
</tr>
<tr>
<td>10-a</td>
<td>7.90</td>
<td>19-at</td>
<td>5.20</td>
<td>F-5&quot;</td>
<td>14.70</td>
</tr>
<tr>
<td>12-c</td>
<td>4.90</td>
<td>22-at</td>
<td>5.40</td>
<td>G-1</td>
<td>9.60</td>
</tr>
<tr>
<td>17-a</td>
<td>4.10</td>
<td>24-at</td>
<td>1.60</td>
<td>G-2</td>
<td>22.70</td>
</tr>
<tr>
<td>18-b</td>
<td>4.40</td>
<td>25-bt</td>
<td>3.70</td>
<td>G-3</td>
<td>15.70</td>
</tr>
<tr>
<td>19-b</td>
<td>2.00</td>
<td>52-et</td>
<td>3.40</td>
<td>H-1</td>
<td>14.30</td>
</tr>
<tr>
<td>20-a</td>
<td>1.70</td>
<td>55-bt</td>
<td>1.10</td>
<td>H-2</td>
<td>17.00</td>
</tr>
<tr>
<td>21-c</td>
<td>4.40</td>
<td>56-bt</td>
<td>3.90</td>
<td>H-3</td>
<td>18.40</td>
</tr>
<tr>
<td>22-b</td>
<td>3.70</td>
<td>59-at</td>
<td>1.70</td>
<td>H-4</td>
<td>16.40</td>
</tr>
<tr>
<td>23-c</td>
<td>1.70</td>
<td>65-bt</td>
<td>4.50</td>
<td></td>
<td></td>
</tr>
<tr>
<td>24-b</td>
<td>4.00</td>
<td>75-at</td>
<td>17.00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*aFor ELISA the isolated and amplified phage clones were captured via their displayed peptides by the biotinylated 14G2a mAb immobilized to streptavidin, and then were detected with the HRP-conjugated anti-phage pVIII antibody. Each value was a ratio of the mean values of OD 450-nm readings of the respective phage clone and the phage clone #0.

Table II. The sequences of the 12 aa peptides isolated from the LX-8 phage-displayed peptide library through biopanning with 14G2a mAb.

<table>
<thead>
<tr>
<th>Peptide</th>
<th>Identified sequence</th>
<th>Number of clones</th>
</tr>
</thead>
<tbody>
<tr>
<td>#8</td>
<td>NCDLLTGPMLCV</td>
<td>12</td>
</tr>
<tr>
<td>#65</td>
<td>SCQSTRMDPNCW</td>
<td>3</td>
</tr>
<tr>
<td>#85</td>
<td>VCNPLTGAALLCS</td>
<td>6</td>
</tr>
<tr>
<td>#94</td>
<td>RCPNPMEPPRCF</td>
<td>4</td>
</tr>
<tr>
<td>#D</td>
<td>GCDALSGHLLCS</td>
<td>14</td>
</tr>
</tbody>
</table>

Total number of sequenced clones 39

second round of immunoblotting (Fig. 1B). Fifty-five of the phage clones again yielded a strong positive signal of 14G2a mAb binding, as compared to a randomly chosen phage clone #0. The phage #0 was picked in the first round of immunoblotting and used as a specificity control since it did not exhibit binding to the antibody (data not shown).

Determination of the phage-displayed peptide sequences. To further characterize and determine the relative affinity of the isolated phage clones to the GD2-specific antibody, we performed ELISA on all of the 55 isolated clones. Amplified and PEG-purified viral particles (10^{10}) were captured via their displayed peptides by the biotinylated 14G2a mAb immobilized in microplate wells to streptavidin, and then detected with the HRP-conjugated anti-phage pVIII antibody (Table I). In comparison to phage #0, among all the screened clones, 39 exhibited a signal that was ~2-fold higher. All these phage clones were scored as specific binders to the 14G2a.

The phage clones that yielded signals which were insignificant compared to the control phage #0, were removed from further analyses (16 clones). Based on the results of the ELISAs, all 39 phage clones were selected for DNA sequencing. Analysis of the sequencing data yielded 5 different 12 aa peptide sequences, which were collected in Table II. All of the identified sequences have cysteine residues at positions 2 and 11. Analysis of the sequence of the #0 phage revealed a stop codon at the first position in the peptide encoding sequence, suggesting that the phage #0 expressed no peptide on the coat surface.

Analysis of the sequence similarities among the isolated peptides was used to estimate the relative contribution of different amino acids of non-sibling peptides to their binding ability to 14G2a mAb. Alignment of the sequences allowed for indication of the degree of identity between the verified sequences. One group of peptides is represented by sequences #8, #85 and #D. A motif of 3 aa residues Leu-Thr-Gly were found in peptides #8 and #85, while #D contained Ser in place of Thr. In addition, the peptides #85 and #D
The role of displayed peptide disulfide bonds in the binding to 14G2a mAb. In the following experiments, we aimed to determine the role of the peptide secondary structure, stabilized by the disulfide bonds formed between the cysteine residues at positions 2 and 11, in the binding of the phage-displayed peptides to the anti-GD2 antibody. Therefore, the phage clones representing all identified sequences were treated with dithiothreitol (DDT), and N-ethylmaleimide (NEM), which modify the peptides. DTT was used to reduce disulfide bonds, while NEM was used to prevent the reforming of these bonds, since it blocks cysteine through reaction with sulphhydryl residues. The immobilized phage clones were subjected to DDT and NEM treatment alone or in combinations (DDT+/NEM-, DDT+/NEM+, DDT-/NEM+ or DDT-/NEM-). Subsequently, the phages were incubated with 14G2a mAb, and the anti-GD2 antibody binding was detected with the HRP-conjugated anti-mouse Ig antibodies, and developed with the TMB substrate. As shown in Fig. 3, we observed a significant decrease in affinities of all the isolated peptides to 14G2a mAb in cysteine modifying conditions (the presented data were normalized to the signal obtained for the phage clone #0). Changes were seen for the wells coated with the phage clones expressing all 5 identified sequences, which were treated with DDT alone, as well as DDT and NEM. No such changes in the binding of the 14G2a mAb were measured for the non-treated phage-coated wells as well as the wells treated only with NEM. Specificity controls were included to ensure that the observed effects were indeed due to modifications of the secondary structure of the peptides stabilized with the disulphide bond. Additional wells were coated with non-constrained 15 aa GD2 ganglioside mimicking peptide #47, and subjected to ELISA in reducing conditions, as above. No changes in the 14G2a binding signal were detected in these wells (data not shown). The obtained results confirm that the effect of the reduction of the binding of 14G2a mAb to the phage clones representing all 5 identified sequences was due to DTT- and NEM-induced changes in the secondary structure of the peptides. To conclude, the experiments showed that the disulfide bond is necessary for the anti-GD2 antibody to bind to the phage particles expressing the identified peptides.

The phage-expressed peptides compete with IMR-32 neuroblastoma cells for binding of the anti-GD2 14G2a mAb. To further characterize the phenomenon of mimicry of the phage-expressed peptides to GD2 ganglioside, we performed a competition assay on a human neuroblastoma cell line IMR-32, which abundantly expresses GD2 ganglioside. 14G2a mAb (200 ng) was incubated overnight with 10^12 ph particles expressing the identified peptides, and then was used to stain the GD2 ganglioside present on the IMR-32 cells. The binding of the antibody to the cells was detected with the FITC-conjugated secondary antibodies, and was analyzed with flow cytometry technique.

Inhibition of binding of 14G2a mAb to the GD2 ganglioside expressing neuroblastoma IMR-32 cells was observed for all of the phage clones screened (Fig. 4). The competition of the peptide-expressing phage clones for 14G2a mAb resulted in a significant decrease in both the number of positively stained cells, as well as the mean fluorescence intensity values, compared to positive control samples, i.e. cells incubated with the 14G2a mAb. Moreover, pre-incubation of the phage clone #0 with the anti-GD2 ganglioside mAb had no effect on the binding of the antibody to IMR-32 cells. The above findings suggest that the observed decrease of the binding of the 14G2a mAb to the GD2 ganglioside present on the cell surface of IMR-32 cells

Figure 2. Dendrogram of five peptide sequences identified as GD2 mimics. Sequence analysis was performed with the ClustalW multiple sequence alignment program with amino acid identity matrix. The analysis revealed two clusters: (#85, #D, #8) and (#65, #94). Identical positions in the sequence are printed in bold type. Motifs of sequence similarity have been identified and highlighted.

Figure 3. Phage ELISA under reducing conditions. Immobilized phage clones representing 5 different peptide sequences (phage #8, #65, #85, #94 and #D) isolated from the LX-8 peptide library were subjected to reducing conditions (with dithiothreitol, DDT+/NEM-), cysteine-blocking conditions (with N-ethylmaleimide, DTT-/NEM+) or both (DTT+/NEM+). A control without the cysteine modifying compounds was also included (DTT-/NEM-). The obtained values were normalized to data obtained for the non-binding phage #0. Each value is a mean ± SE from three individual experiments.
Figure 4. Results of competition test. Phage particles (10^12) expressing the 5 different peptides binding to 14G2a mAb were incubated overnight with 200 ng of the antibody, and then were used in a competition assay on the GD2 ganglioside-positive IMR-32 neuroblastoma cells. The binding of 14G2a mAb to the cells was detected with the secondary FITC-conjugated (Fab')_2 fragments using flow cytometry technique. Three peptides that exhibited the highest inhibition; the non-binding phage #0; PC, positive control (no phage present); and NC, negative control (without 14G2a mAb) are shown. Data shown are a representative of three individual experiments.

Figure 5. The binding of the synthetic peptides to 14G2a mAb was verified using the competition assay. The synthetic peptides (#8, #65, #85, #94 and #D) were incubated overnight with 200 ng of the 14G2a mAb, and then used in the competition assay on the IMR-32 cells. The binding of 14G2a mAb was detected with the secondary FITC-conjugated antibodies using flow cytometry. A sample with a non-binding peptide was included as a specificity control (#Con peptide). Data shown represent means ± SE from three independent experiments.
was due to the presence of phage-displayed peptides. Based on the signal from negative controls, the pools of IMR-32 cells that specifically bound the 14G2a mAb were determined. In these pools, mean fluorescence intensity was measured, and then used for the calculation of % of inhibition. Among the 5 phage clones tested, the phages expressing peptides #65, #85 and 94 showed the highest % of inhibition; 76.8% (±5.1) for the phage #65; 82.9% (±4.1) for the phage #85; and 82.9% (±1.1) for the phage #94 (Fig. 4). The values calculated for the last 2 tested phages were significantly lower; 30.6% (±7.4) and 45.9% (±4.8), for the phages #8, and #D, respectively (data not shown).

Synthetic peptides are mimics of GD2 ganglioside. Based on the results of the above experiments, synthetic peptides encoding the 5 identified sequences were tested in the competition assay on IMR-32 cells. This test allowed us to verify the results observed with phage-displayed peptides. The concentrations of the peptides that were used ranged from 0.1-1000 μM. Indeed, all the synthetic peptides prevented the 14G2a mAb from binding to IMR-32 cells. The observed effects were dose dependent. A sample with a non-binding peptide was included as a specificity control (#Con peptide). For this peptide, only slight inhibition was observed for concentrations of the peptide >100 μM. The results suggest that the observed effects of inhibition of binding of the anti-GD2 mAb to IMR-32 cells were specific for the synthetic peptides isolated from the LX-8 library (Fig. 5). IC₅₀ values were calculated for the GD2-mimicking peptides. All 4 peptides isolated with the ‘in solution’ panning protocol had comparable equal abilities to compete for peptide #8. The IC₅₀ value determined for peptide #D was ~8 times lower (165.8±60.2 μM). To conclude, we state that the observed mimicry of the GD2 ganglioside by the peptides was related both to the presence of the specific amino acids and the overall conformation of the peptides stabilized with disulphide bridge. More information on the involvement of the specific amino acid residues in the binding of the peptides to the GD2-specific antibody may be gained from replacement analyses of the sequencing data. As the identified peptide sequences was used to estimate the relative binding contributions to a receptor of different amino acids of non-sibling peptides (34). However, since only 5 peptide sequences were identified among 39 sequenced clones, such detailed analysis cannot be performed. Yet, our isolated peptides showed some sequence identities. The presence of the defined motif Leu-Thr/Ser-Gly did not seem to be crucial for binding to 14G2a mAb, as peptides #65 and #94 lacked the motif as well as Leu at position 9, compared to peptides #8, #85 and #D. Yet, similar 14G2a binding affinities were obtained for the peptides #65, #85 and #94, both presented on the phage surface as well as in the synthetic form. Notably, the secondary structure of the peptides, stabilized by the disulfide bridging between cysteine residues at positions 2 and 11, was shown pivotal for the 14G2a binding abilities of all isolated peptides. Thus, it may be that the observed mimicry of the GD2 ganglioside by the peptides was mimics of GD2 ganglioside, since they were shown to specifically compete for the binding of 14G2a mAb with its native antigen, the GD2 ganglioside present on IMR-32 neuroblastoma cells.

Discussion

In the current study, we focused our research to isolate constrained peptides from phage-displayed peptide library LX-8 with 14G2a mAb, and to characterize their ability to mimic GD2 ganglioside, which is one of the most prominent neuroblastoma antigens. The performed biopanning experiments led to the isolation of affinity-enriched phage pools for the screening molecule. Two rounds of immunoblotting were successfully applied to eliminate irrelevant binders that represented e.g. plastic, BSA as well as Fc binding clones as such phage clones could have been amplified, despite the pre-adsorption step preceding each panning round. Additionally, prior to the decoding of the peptide sequences from the viral ssDNA, application of ELISA allowed us to estimate the strength of 14G2a binding of all 55 clones isolated through plaque assay, and to discriminate between weak and strong binders. As a result, 16 weak binding clones were excluded from further analyses. Five varying peptide sequences were identified through analyses of the sequencing data. As the identified peptide sequences were isolated from phage pools obtained through panning carried out according to two protocols, and with different 14G2a mAb concentrations, it was impossible to establish a correlation between the measured 14G2a binding strength and the frequency of the isolated clones. Such a correlation has been reported in some experiments, while other data have suggested only a weak link between the binding affinity of the phage clones and its selection frequency (32,33). The fact that only one sequence (#D) was identified with the 6-round ‘solid phase’ protocol may be explained by the overgrowth of certain clones caused by factors other than affinity (e.g. biological fitness).

Analysis of the sequence similarities between the isolated peptide sequences was used to estimate the relative binding contributions to a receptor of different amino acids of non-sibling peptides (34). However, since only 5 peptide sequences were identified among 39 sequenced clones, such detailed analysis cannot be performed. Yet, our isolated peptides showed some sequence identities. The presence of the defined motif Leu-Thr/Ser-Gly did not seem to be crucial for binding to 14G2a mAb, as peptides #65 and #94 lacked the motif as well as Leu at position 9, compared to peptides #8, #85 and #D. Yet, similar 14G2a binding affinities were obtained for the peptides #65, #85 and #94, both presented on the phage surface as well as in the synthetic form. Notably, the secondary structure of the peptides, stabilized by the disulfide bridging between cysteine residues at positions 2 and 11, was shown pivotal for the 14G2a binding abilities of all isolated peptides. Thus, it may be that the observed mimicry of the GD2 ganglioside by the peptides was related both to the presence of the specific amino acids and the overall conformation of the peptides stabilized with disulphide bridge. More information on the involvement of the specific amino acid residues in the binding of the peptides to the GD2-specific antibody may be gained from replacement and truncation experiments. Additionally, such analyses would allow further optimization of the obtained sequences (35). Other peptide mimicking GD2 gangliosides have already been published. In comparison, our 5 peptides are of different length, have no considerable sequence identity, and have no obvious similarities in motif. Linear 15 aa peptides were isolated with 14G2a mAb (29). Forster-Waldl and co-researchers reported phage clones expressing cyclic peptides build by 10 amino acids, which were looped by the addition of 2 flanking cysteine residues, and were isolated from the phage-displayed library based on pIII fusions, and with the anti-GD2 ganglioside antibody ch14.18. However, the exact role of the disulphide bridge has not been investigated for their peptides (36).

The therapeutic potential of peptides derived from phage-displayed peptide libraries relies on the fact that they can structurally mimic the native antigen interactions with the antigen-binding site of the screening antibody. The final proof that all of the isolated phage-displayed peptides interacted with the GD2 ganglioside-binding site of the 14G2a mAb was obtained from competition assays. Here, we measured quantitative differences in the inhibition of the
isolated with the ‘in solution’ panning protocol showed peptides transferred to the non-phage milieu were still on the results of the inhibition of binding of 14G2a to IMR-14. Moreover, we confirmed specific interactions of 14G2a mAb specific due to the presence of the displayed peptides. Inhibition, we concluded that the observed effects were the phage clone #0, which expresses no peptide, yielded no representing peptides #8 and #D performed, respectively, the GD2-positive neuroblastoma cells. The phage clones side, expressed on IMR-32 neuroblastoma cells. With reference to the calculated percentage of inhibition, the phage clones displaying peptides #94, #65 and #85 were shown to almost completely inhibit the binding of the 14G2a mAb to the GD2-positive neuroblastoma cells. The phage clones representing peptides #8 and #D performed, respectively, three and two times less efficiently in the inhibition assay. As the phage clone #0, which expresses no peptide, yielded no inhibition, we concluded that the observed effects were specific due to the presence of the displayed peptides. Moreover, we confirmed specific interactions of 14G2a mAb binding with the antigen-binding site during the inhibition tests carried out with chemically synthesized peptides. Based on the results of the inhibition of binding of 14G2a to IMR-32 cells by synthetic peptides, we concluded that all five peptides transferred to the non-phage milieu were still mimicking GD2 ganglioside. Peptides #8, #65, #85 and #94 isolated with the ‘in solution’ panning protocol showed similar IC50 values, while peptide #D showed significantly higher IC50 values. The reasons for these were not investigated. We hypothesized that analysis of the peptides in a monovalent system overcame the affinities of multiply displayed peptides and effects of the phage on the peptide structure, as it might occur on the phage surface.

The mimicry of peptides isolated from phage libraries can be evaluated on various levels. These include a structural mimicry related to the 3D conformation, an immunochemical mimicry based on the patterns of binding to an antibody, and a functional mimicry that allows induction of immune responses specific for a nominal antigen (37). Based on the available research data it is apparent that these levels may not necessarily be interrelated. For instance, Willers and colleagues found that peptide mimics of GD3 ganglioside did not elicit any detectable immune responses in mice, probably due to the presence of common motifs between the reported GD3 mimics, and mouse or human proteins (38). On the contrary, peptide mimics of Lewis Y induced specific anti-carbohydrate antibodies in mice (39). Furthermore, the generated immune responses could be further manipulated with various delivery schedules, i.e. prime/boost strategies with peptides and carbohydrate, or a DNA vaccination approach (21,40). The presence of anti-GD2 ganglioside responses has already been investigated with application of mimicking peptides (29,41). Indeed, mutual relationships between the observed mimicry in binding to 14G2a mAb and the functional ability to induce GD2 ganglioside-specific immune responses must be established if the GD2 mimicking peptides isolated from the LX-8 phage-displayed peptide library with 14G2a mAb are mimics of the tumor-associated GD2 ganglioside. These results warrant further consideration and study of the molecules as possible novel immunogens to overcome tolerance to the GD2 ganglioside, and to induce specific immune responses that may be clinically important for the treatment of patients bearing GD2 expressing tumors, especially, to control the minimal residual disease.

Acknowledgements

This study was supported by grant no. 3P05A 00124 from the Polish Ministry of Scientific Research and Information Technology. We are grateful to Dr Ralph A. Reisfeld for the 14G2a-producing hybridoma cells. We thank Dr Malgorzata Bzowska for help with the flow cytometry analysis.

References


