Identification of VEGFR2-Binding Peptides Using High Throughput Bacterial Display Methods and Functional Assessment

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Abstract: The signal transduction pathway initiated by vascular endothelial growth factor-vascular endothelial growth factor receptor 2 (VEGF-VEGFR2) plays an important role in the angiogenesis of tumors. The effective antagonists of VEGFR2 would behave as potent drugs for the treatment of malignant cancers. In our study, specific binding peptides with high affinity to VEGFR2 were obtained through bacterial display technology. Conserved motif (FF/YEXWGVK) among those peptide sequences was discovered. The results of surface plasmon resonance (SPR) assay indicated that the dissociation constant ($K_D$) value of VRBP1 was 228.3 nM and this peptide competed with VEGF binding to VEGFR2. Particles conjugated with VRBP1 could recognize the human umbilical vein endothelial cells (HUVEC) which express VEGFR2 on the surface. Further therapeutic effect of VRBP1 was examined by \textit{in vivo} experiments. VRBP1 could result in a significant decrease in tumor size of H460 xenografts. The results from the immunohistochemical assay showed that CD31 positive signals in VRBP1-treated group were fewer than those in the control ones. These data highlighted the potential of VEGFR2-binding peptides as effective molecules for cancer diagnosis and therapy.

Keywords: Angiogenesis, bacterial surface display, cancer diagnosis, cancer therapy, peptide, vascular endothelial growth factor receptor 2 (VEGFR2).

INTRODUCTION

Angiogenesis is a normal process occurring during embryonic and post-embryonic development, reproductive cycle and wound repair. Now, many data have shown that angiogenesis plays a critical role in cancer development, especially in the rapid growth of solid tumors beyond 1-2 mm in diameter and metastatic tumors [1, 2]. Studies on cancer patients have shown a direct correlation between the density of tumor vessels and an adverse prognosis [3, 4], so the development of specific anti-angiogenic agents arises as an attractive approach for the treatment of cancer and other angiogenesis-dependent diseases [4]. Vascular endothelial growth factor (VEGF) as an endothelial cell-specific mitogen has been found to be the most important angiogenic factors in tumor angiogenesis [5-7]. VEGF functions by binding to one of its specific tyrosine kinase receptors: VEGFR1 (fms-like tyrosine kinase/Flt1) [8-10], VEGFR2 (a kinase insert domain-containing receptor/KDR) [10, 11] and VEGFR3 (Flt-4) [12, 13] to promote either physiologic or pathologic angiogenesis. Only homodimerization of VEGFR2 leads to a strong autophosphorylation and it exhibits a strong induction in angiogenesis [14, 15], while VEGFR1 participates in transduction of a weak angiogenic signal [14]. Furthermore, VEGFR2 is usually found to be aberrantly expressed or constitutively phosphorylated in tumors [16-18]. Recently, Sorafenib [18, 19], an anti-VEGFR2 antagonist was introduced to the market for cancer treatment. Although antibodies targeting the different cancer-related molecules have already been used in cancer therapy successfully [20], some targeting molecules such as peptides have presented numerous advantages over antibodies and may be used as substitutes for antibodies in future. For example, peptides have smaller sizes and better tissue penetration than antibodies, and coupling drugs or imaging agents with peptides is easier than that with antibodies [21, 22]. In addition, peptides represent alternatives to antibodies in terms of maximizing safety and minimizing production costs [21, 23].

Phage display technology or chemical synthesis has been successfully used to obtain the requisite peptides [24-26]. Several VEGFR2-binding peptides have been identified using phage display method [27, 28], but no consensus sequences were discovered and their affinity and specificity to VEGFR2 still need to be improved for the clinical use. Therefore, it is necessary to develop more specific binding peptides to VEGFR2 with high affinity for cancer diagnosis and therapy. In our study, we applied bacterial display
method to identify the specific binding peptides for extracellular domain of VEGFR2. Bacterial display method coupled with fluorescence-activated cell sorting (FACS) technology is a simpler, more effective, high throughput and more quantitative approach to discover and optimize peptides comparing with phage display methods [29-31]. In this study, a specifically VEGFR2 binding peptide with moderate affinity was identified. It not only inhibited proliferation of human umbilical vein endothelial cells (HUVEC) in vitro, which express VEGFR2 on their surface [32], but also reduced the tumor growth of H460 xenografts in vivo. Therefore, this peptide has the potential to be used as an effective molecule for cancer diagnosis and therapy.

MATERIALS AND METHODS

**Bacterial Strains, Cell Lines and their Growth Conditions, Plasmids and Reagents**

A library of peptides with 15 randomized amino acids (X_{15}) on eCPX scaffold including 2×10⁹ members used in this study was a gift from Patrick S. Daugherty in the Department of Chemical Engineering, University of California, Santa Barbara [33]. Recombinant human VEGF₁₆₅ (rhVEGF₁₆₅) was purchased from Beijing ZhongKeWuYuan Biotechnology (http://www.zkwysw.com). Recombinant human VEGFR1, VEGFR2 and VEGFR3 were purchased from Sino Biological Inc (http://www.sinobiological.com). Streptavidin R-phycoerythrin (SAPE) (Cat No. 1148354), Dynabeads MyoneTM Carboxylic Acid (Cat No. 65011) and MyOne streptavidin-coated magnetic microbeads (Cat No. 650.01) were bought from Invitrogen. IgG and human serum was purchased from Sigma. Fetal bovine serum (FBS), Dulbecco’s modified Eagle’s high glucose medium (DMEM/high), RPMI-1640 medium and Medium 199 (M199) were purchased from Thermo (Hyclone Corp., USA). Peptides in this article were synthesized by Shanghai Bioengineering Ltd. Bovine serum albumin (BSA), chloramphenicol (CM), penicillin/streptomycin and D- (+)-glucose were purchased from China National Medicine Corp. Superparamagnetic iron oxide particle (SPIO) was a gift from Dr. Yu Zhang (Southeast University, China) [34].

HLF (human lung fibroblast) cells were grown in DMEM/ high medium and human lung cancer cells (H460) were cultured in RPMI-1640 medium in a 5% CO₂ humidified incubator at 37 °C. The medium was supplemented with 10% FBS and 1% penicillin/streptomycin. H460 and HLF cells were kind gifts from Dr. Biliang Zhang (Guangzhou Institutes of Biomedicine and Health, CAS, China).

The HUVEC cells were separated and cultured according to the procedure described by Maruyama et al. [35, 36]. HUVEC cells were maintained in M199 in a 5% CO₂ humidified incubator at 37 °C. The medium was supplemented with 20% FBS and 1% penicillin/streptomycin. Tumor cell lines were authenticated by short tandem repeat (STR) analysis.

**Screening of Binding Peptides to VEGFR2 from a Random Library**

Biotinylation of VEGFR2 was performed using the FluoReporter mini-biotin-XX protein labeling kit (Invitrogen Corp., CA, USA, Cat No. F-6347) and following its instruction. A library of peptides with 15 randomized amino acids was used for screening the binding peptides to VEGFR2. Frozen aliquots of 2×10¹⁰ bacteria were thawed and cultivated overnight with shaking at 200 rpm in super optimal broth (SOB) at 37 °C with 34 mg/mL CM and 0.2% (w/v) D- (+)-glucose added. The next day, bacteria were sub cultured at 1:50 in lysogeny broth (LB) medium supplemented with 34 mg/mL CM for 2 h at 37 °C followed by induction with 0.02% (w/v) L- (–)-arabinose for 1 h at room temperature to ensure peptides expressed on the surface of bacteria [33]. Magnetic-activated cell sorting (MACS) was first performed with 100 nM biotinylated VEGFR2 to reduce the library size and ratio of beads: bacteria was 1:2, as previously described [29, 37]. Afterwards, the library was incubated with biotinylated VEGFR2 at various concentrations from 20 nM to 50 nM for 45 min at 4 °C with or without co-incubated with 20 μM human IgG. After washing with 1 mL phosphate-buffered saline (PBS, pH 7.4) for three times and 100 μL (3.3 nM) SAPE was added into the tube, bacteria clones with high fluorescence intensity, meaning that peptides expressed on the bacteria surface could bind with VEGFR2, were sorted. These sorted bacteria clones were used for the next round of screening.

A few rounds of FACS were performed and when the fluorescent signals of bacteria stopped increasing in the two sequential rounds, i.e., when the binding bacteria clones for VEGFR2 were fully enriched, the screening procedure for the binding peptides was terminated. Sorted bacteria with binding peptides were cultured on LB medium plate containing 1% agar and 34 mg/mL CM. After culturing for 16 h at 37 °C, monoclonal bacteria were picked and cultured in 5 mL SOB containing 34 μg/mL CM at 37 °C overnight. The sequences of the peptides showed on these clones were obtained by sending these bacteria to Sangon Biotech Co., Ltd. Shanghai, China for sequencing.

**Determination of Binding Activity for Peptides-Bacteria with VEGFR2**

The preliminary examination of peptides-bacteria binding properties was performed under different treatment conditions: tough washing and serum or IgG blocking during incubation process. Individual clones including 1×10⁵ bacteria induced by 0.02% (w/v) L-(–)-arabinose for 1 h at room temperature were incubated with 67 nM biotinylated VEGFR2 with or without 10% serum or 20 μM human IgG for 45 min at 4 °C. The bacteria were centrifuged at 3000×g for 5 min and washed with PBS. Sediments were resuspended in 100 μL SAPE and incubated for 20 min at 4 °C. Prior to analysis, bacteria were centrifuged at 3000×g for 5 min and resuspended in 1 mL PBS, with or without three extra washes. The binding of VEGFR2 was assessed via reading of fluorescent intensities of bacteria by flow cytometry.

**K₀ Measurement of Peptides-Bacteria to VEGFR2**

Biotinylated VEGFR2 at various concentrations of 0.5 nM, 1 nM, 5 nM, 15 nM, 40 nM and 60 nM was incubated with induced bacteria clones and the fluorescent intensities were recorded by flow cytometry. The relative binding affinity (K₀ value) was estimated by formula Fl-Fl_{eCPX}=Fl_{max}
receptors were removed and un-reacted moieties on the chip
response units for flow cell 2) were reached. Uncoupled
μL/min until the target immobilization levels (2,000
the activated surfaces of flow cell 2 at a flow rate of 10
μg/mL in 10 mM sodium acetate, pH 5.5) was injected over
according to the manufacturer's instructions. VEGFR2 (2-10
(EDC) and 0.1 M N-hydroxysuccinimide in water (NHS)
ethyl-3-(3-dimethylaminopropyl)-carbodiimide in water
Suzhou, China) was activated by a 1:1 solution of 0.4 M 1-
determined by SPR on a Biacore 3000 system (BIAcore AB,
VEGFR2
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indicates the fluorescent intensities of cCPX bacteria after
incubation with VEGFR2. Flmax represents the highest
fluorescent value of bacteria-VEGFR2 complex and [P] is
the concentration of VEGFR2.

Enzyme-Linked Immunosorbent Assay (ELISA) with
Fluorescein Isothiocyanate (FITC)-Conjugated Peptides

VEGFR2 (2 μg/mL in PBS buffer, pH 7.4) was coated on
96-well plates (50 μL/well) and kept at 4 °C overnight. These wells were washed with PBS and then blocked with
3% BSA in PBS with 0.05% Tween20, for 1 h at room
temperature. FITC-conjugated VRBP1 and control peptide
(CONP, GSSGSSGSGSSGSSGSGSSGSSG) at a
concentration of 20 μM in PBS were added and incubated at
room temperature for 30 min. After washing with PBS,
fluorescence intensity was measured by VICTOR X4
Multilabel Reader (PerkinElmer Life and Analytical
Sciences, Boston, MA, USA).

Examination of Peptides Binding Activity to HUVEC
Cells in vitro

These experiments were performed on both attached and
detached HUVEC cells. The fluorescence microscopy
experiment was carried out to examine the binding efficiency of
synthesized FITC-conjugated peptides with attached
HUVEC cells. HUVEC cells were seeded in 12-well plates
(Costa, corning incorporated) at a density of 1×105 cell/well
overnight and fixed with 4% paraformaldehyde at room
temperature for 10 min. After washing, cells were blocked with
a culture medium containing 5% BSA at 37 °C for 10
min and then incubated with FITC-conjugated peptides in
serum-free medium at 37 °C for 1 h. The plates were
visualized under a fluorescent microscope (Nikon Diaphot,
Nikon Corp., Tokyo, Japan). The binding activity between
synthesized FITC-conjugated peptides with detached
HUVEC cells was determined by flow cytometry. Cultured
(48 h post-seeding) HUVEC cells were harvested and
resuspended in tubes. 1×107 HUVEC cells were incubated with
0.5 nmol FITC-conjugated peptides for 45 min on an
inversion shaker at 4 °C. Cell suspensions were centrifuged at
300×g for 5 min at 4 °C and then washed three times with
PBS. The sediment was resuspended with 1 mL PBS and
immediately analyzed by FACS.

Measurement of K0 Values for Synthesized Peptides to
VEGFR2

Dissociation constants of synthesized peptide were
determined by SPR on a Biacore 3000 system (Biacore AB,
Uppsala, Sweden). The HRBio sensor chip (SJI Biomaterials,
Suzhou, China) was activated by a 1:1 solution of 0.4 M 1-
ethyl-3-(3-dimethylaminopropyl)-carbodiimide in water
(EDC) and 0.1 M N-hydroxysuccinimide in water (NHS)
according to the manufacturer’s instructions. VEGFR2 (2-10
μg/mL in 10 mM sodium acetate, pH 5.5) was injected over
the activated surfaces of flow cell 2 at a flow rate of 10
μL/min until the target immobilization levels (2,000
response units for flow cell 2) were reached. Uncoupled
receptors were removed and un-reacted moieties on the chip
were blocked with 1 M ethanolamine-HCl (pH 8.5). Flow
cell 1 was activated and blocked without exposure to
VEGFR2 and served as background control surfaces for the
binding interactions.

All kinetic binding experiments were performed at 25 °C
degassed PBS running buffer (pH 7.4). Various concentrations (0.2, 0.4, 0.5 and 1 μg/mL) of peptides were
flowed through the surface-immobilized VEGFR2 at 10
μL/min for 5 min and the binding events were monitored.
Following association, the dissociation process of peptide-receptor complexes was monitored for 7 min.

Examination of Competence between VEGF and
Peptides Binding to VEGFR2 by SPR Assay

The HRBio sensor chip was activated by a 1:1 solution of
EDC/NHS. The activated surfaces of flow cell 2 were passed
over by 10 μg/mL VEGFR2 at a rate of 10 μL/min until the
2000 response units (RU) were reached. After the
immobilization of VEGFR2, binding peptides at a
concentration of 10 μg/mL were injected at a rate of 10
μL/min followed by 3 μg/mL VEGF passed over at a rate of
10 μL/min. In another chip, the 3 μg/mL VEGF was injected
followed by 10 μg/mL binding peptides passed over. The RU
was monitored in real-time and presented in a sensorgram of
RU versus time.

Competitive Binding Assay of VEGF to VEGFR2 with
ELISA

VEGFR2 was coated onto 96-well plates as described
above. FITC-conjugated VRBP1 at a concentration of 20 μM
in PBS was mixed with VEGF at different concentrations (0,
0.05, 0.1, 0.2, 0.4, 0.8, 1 and 2 ng/mL) and added to wells.
After 30 min of incubation at room temperature, the wells
were washed and FITC-conjugated VRBP1 bound to VEGFR2
was measured by VICTOR X4 Multilabel Reader.

MTT Assay

The blocking activity of peptides to VEGFR2 was
evaluated by methyl thiazolyltetrazolium (MTT) assay. HUVEC cells were plated into 96-well tissue culture plates
(Costa, corning Corp., USA) at a density of 8000 cells per
well and cultured overnight. After that, cells were serum
starved for another 24 h. In the serum free medium,
rhVEGF165 (30 ng/mL) alone or rhVEGF165 together with
peptides ranging from 5 μg/mL to 50 μg/mL was introduced
to wells for 48 h. Twenty μL MTT (5 mg/mL) was added
to each well and incubated for another 4 h. The supernatant
was removed followed by adding 150 μL dimethyl sulfoxide
(DMSO) in order to dissolve the formed water-insoluble
formazan crystals. Reading of optical density (OD) at 490
nm was obtained from VICTOR X4 Multilabel Reader.

Conjugation Experiments of Peptides with Dynabeads or
Superparamagnetic Iron Oxide Particles (SPIO)

Dynabeads MyoneTM Carboxylic Acid with an average
diameter of 1.06 μm and SPIO with an average diameter
of 20 nm were used to perform the peptides conjugation
experiment. A mixture of EDC (20 mg/mL) and NHS
(1.2 mg/mL) in 0.1 M 2-(N-morpholino)-ethanesulfonic acid
times, mice were sacrificed and tumor tissues were put into 0.9% physiological saline was injected to tumor sites. Mice, 100 μL VRBP1 (400 μg/mL), CONP (400 μg/mL) or saline. One day after tumor xenograft was transplanted to the right leg of each mouse. One day later, mice were divided into 2 groups, with four mice in each group.

Evaluation of the Targeting Effect of Peptides to VEGFR2 by Trapping Experiments

HUVEC cells, HLF cells (as a control cell which expressed VEGFR2 at a low level) were plated into 24-well cell culture plate at a concentration of 3×10⁴ cells per well. Ten μL Dynabeads coupled with different peptides were re-suspended in 200 μL PBS and added into wells. During the incubation process, the plate was shaken every 30 seconds for 10 min. Afterwards, each well was washed 3 times with PBS. The targeting effects of peptides to VEGFR2 were evaluated by comparing the amount of cells with different peptide treatments under the inverted microscope (Nikon Diaphot, Nikon Corp., Tokyo, Japan).

Evaluation of Targeting Effect of Peptides to VEGFR2 by MRI (Magnetic Resonance Imaging) Detection

HUVEC cells and HLF cells, at a density of 0.5×10⁶ cells per well in a 6-well plate, were washed and trypsinized after incubation with SPIO at a final Fe concentration of 0, 19, 38, and 76 μg/mL for 2 h. Cells were collected by centrifugation at 300×g for 5 min. The obtained cells were suspended in 1% agar in 500 μL PBS and the mixtures were injected into the NMR tubes and detected as follows. T2 weighted images were acquired using multi-slice multi-echo (MSME) protocol on a 11.7 T Bruker micro 2.5 micro-MRI system with the following parameters: repetition time (TR) = 5000 ms, echo time (TE) = 80 ms, imaging matrix = 128×128, slice thickness = 1.5 mm and field of vision (FOV) = 2.20 cm × 2.20 cm.

Construction of Tumor Mice Models with Subcutaneous Xenograft

The nu/nu mice (female, 3-4 weeks old) were obtained from Nanjing Sikerui Biological Technology Co., Ltd. Mice were maintained under super pathogen-free conditions and housed in barrier facilities on a 12h light/dark cycle, with food and water ad libitum. The mice were maintained as per the principles and guidelines of the ethical committee for animal care in accordance with the China National Law on animal care and use. Cultured tumor cells (H460) were implanted subcutaneously (sc) into the flanks of mice. When tumor volume reached 2 cm³, mice were sacrificed and the tumor mass was divided into small pieces. A small tumor tissue about 2 mm³ was transplanted subcutaneously into the right leg of each mouse. One day later, mice were divided randomly into 2 groups, with four mice in each group.

The Treatment of Mice with Peptides

VRBP1 and CONP were dissolved in 0.9% physiological saline. One day after tumor xenograft was transplanted to mice, 100 μL VRBP1 (400 μg/mL), CONP (400 μg/mL) or 0.9% physiological saline was injected to tumor sites subcutaneously every other day. After being injected for 11 times, mice were sacrificed and tumor tissues were put into 4% paraformaldehyde for further experiments. During treatment, the tumor size was measured and recorded in two dimensions, and the volume was calculated using the formula: tumor volume (mm³) = 1/2 × length (mm) × width (mm).

Immunohistochemistry and Immunofluorescence-Like Assay

For immunohistochemistry assay, the fixed tumor tissue samples were embedded in paraffin and sectioned into slices as thin as 5 μm with a microtome. Tissue sectioned at 5 μm were deparaffinized in xylene, rehydrated through graded alcohol series, and subjected to immunohistochemistry assay with a monoclonal rabbit anti-mice CD31 antibody. Antigen retrieval was done in citric acid (pH 6.0) at 95 °C for 20 min. Endogenous peroxidase activity was blocked with 3% hydrogen peroxide for 10 min, and nonspecific binding sites were blocked with 5% BSA for 30 min at 37 °C. Primary antibody to CD31 was diluted at 1:100 in PBS. Next day, goat anti-rabbit immunoglobulin antibodies labeled with horseradish peroxidase were diluted at 1:500 in PBS and incubated with sections for 45 min at 37 °C. Sections were incubated with diaminobenzidine chromogen substrate solution for 10 min and counterstained with hematoxylin. After that, sections were visualized with a Nikon Eclipse 55i microscope equipped with a Nikon DS-5M camera, using Image-Pro Plus acquisition software. 100×, 200× and 400× images were acquired.

For immunofluorescence-like assay, tissue sections from mice xenograft were incubated with 30 μM FITC labeled peptides for 45 min at room temperature after blocking with 1% BSA for 30 min. After washing with PBS three times, tumor sections were observed under fluorescence microscope.

Intratumormicrovessel Density (MVD) Accessment

MVD was assessed by light microscopic analysis for neovascular hot spots. Areas with neovascularization at the highest density were selected by scanning the tumor sections at low magnification (40× and 100× total magnification). Brown-staining microvessel counts were made on a Nikon Diaphot, Nikon Corp., Tokyo, Japan). P-value < 0.05 was considered to be statistically significant.

RESULTS

Bacteria Clones Binding to VEGFR2 were Identified from a Fully Random Bacterial Peptide Library

To obtain the peptide sequences that bound specifically to VEGFR2, bacteria displaying a fully random 15-mer
peptide library ($X_{15}$) were incubated with VEGFR2. Bacterial libraries resulting from each cycle of screening were analyzed with flow cytometry to track enrichment. The sorting processes were carried out for 8 cycles including one round MACS (M1), and seven rounds of FACS selections. Initially, in the first two rounds of FACS selections, the concentrations of VEGFR2 were 50 nM (R1) and 20 nM (R2), and percentages of bacteria in the sorting gate were 25.1% and 38.7% (Fig. 1A), respectively. Subsequently, the sorting strategy was changed by increasing the stringency of sorting. Following the M1, the concentration of VEGFR2 in the following screening cycles (F1 to F7, Fig. 1B) was decreased from 50 nM to 20 nM and 20 μM human IgG was added to the screening system. In the end, 100 clones were sent for sequencing and only seven independent binding peptides sequences were obtained (clone1-clone7). A consensus sequence of FF/YEXWGVK (Fig. 2A) was presented in most of the clones. The preliminary examination of the affinity and specificity of those peptides clones which were on the surface of bacteria for VEGFR2 was performed and in this in situ test, clone 1 and clone 2 kept higher fluorescent signals even under serum blocking and tough washing treatment compared with other clones (Fig. 2B), which meant that these two clones might have higher binding affinity and specificity to VEGFR2. The apparent $K_D$ values of bacteria clone 1 and clone 2 to VEGFR2 were 287.7 ± 0.988 nM and 312 ± 0.994 nM, respectively (Fig. S1).

Fig. (1). The enrichment of peptide-fluorescent bacteria binding with VEGFR2. (A) One round of MACS with concentration of VEGFR2 at 100 nM (M1) and two rounds of FACS sorting with concentrations of VEGFR2 at 50 nM (R1) and 20 nM (R2), respectively. (B) Seven rounds of FACS sorting with 20 nM VEGFR2 and 20 μM human IgG (F1-F7).


VEGFR2 Binding Bacteria Clones with Higher Affinity and Specificity Were Identified from Focused Library

In order to get higher affinity peptides and make them more suitable for application in clinic, a focused library was designed and constructed. On the N-terminus of enhanced CPX (eCPX) [33], a focused library was constructed with the form ZZZZZFF/YEXWGVKZZZZZ in which Z stood for biased amino acids except for F, L, M, V, and I. PCR was performed with CGTAGCTGCCAGTCGTGGCAGNVSN VSNVSNVSNVSNVSTYTT/TAYGARNSTAGGGNTR AARNVSNVSNVSNVSGAAGCCGTCTGGGCGA GTC as the forward primer, GCCGTGAAATCTTCAGTC as the reverse primer and pB33eCPX-SApep [38] as a template. The PCR product was digested with SfiI and ligated into a similarly digested pB33CPX vector. The library size was 5 × 10^6.

Sorting experiments for VEGFR2 binding peptides were performed with 20 nM biotinylated VEGFR2 and 20 μM human IgG. After 4 rounds of sorting, a series of binding clones out of 30 random ones containing the core sequences were obtained (Fig. 2C). The preliminary examination of the affinity and specificity of those peptides-bacteria for VEGFR2 was performed with methods similar as part of ‘Determination of binding activity for peptides-bacteria with VEGFR2’ in materials and methods section. Clone F5 and clone F11 (Fig. 2C) seemed to have relatively higher fluorescent signals compared with other clones which meant that these two clones might have high binding affinity to VEGFR2 (data not shown). The peptide displayed on clone F11 (YDGNSFYEMWGVKPTES) was named as VRBP1 and that on clone F5 (RSRNAFFELWGVKRETPG) was named as VRBP2. VRBP1, VRBP2, consensus sequence FF/YEXWGVK (named as VRBCP) and CONP were synthesized with/without FITC conjugated at their N termini. The KD value of synthesized VRBP1 to VEGFR2 was 228.3 nM measured by SPR assay. And the results of ELISA indicated that VRBP1 could specifically bind to VEGFR2 (Fig. 2D).

VRBP1 could Recognize VEGFR2 on HUVEC Cells and Tumor Vessels

To investigate physiochemical properties of peptides sorted from bacterial library, the VEGFR2 recognizing
experiment was performed with VRBP1, VRBP2, VRBPC and CONP. Fig. 3A showed that HUVEC cells emitted strong green fluorescent signals after incubation with FITC-conjugated VRBP1 and VRBP2, while weaker fluorescent signals on HUVEC cells incubated with VRBPC were observed. The ignorable fluorescent signals were observed on HUVEC cells when incubated with CONP.

Suspended HUVEC cells were also applied to examine the binding properties of peptides. As shown in Fig. 3B, the percentage of the HUVEC cells binding with fluorescent labeled VRBP1 was 34.1%. HUVEC cells incubated with FITC-VRBP2 also showed reasonable high fluorescent signals, which indicated that peptides obtained from focus library could recognize HUVEC cells effectively. The percentage of the VRBPC-bound HUVEC cells was 23.2%, while that with CONP was 5%.

The binding efficiency and specificity of VRBP1 to VEGFR2 were further confirmed by the immunofluorescence-
like assay with tumor sections. More green fluorescent signals were observed on the slides incubated with VRBP1, while there were ignorable signals shown up on the slides with CONP (Fig. 3C).

**VRBP1-Beads could Specifically Attach to HUVEC Cells**

Since peptide VRBP1 could bind with VEGFR2 with higher affinity, the examination of the targeting function of VRBP1 was performed. The carboxylic beads coupled with VRBP1 and CONP were incubated with the HUVEC and HLF cell lines respectively at 4 °C for 10 min. After washing for 3 times with PBS, huge amount of the beads conjugated with VRBP1 on the surface was observed to be accumulated on the surface of HUVEC cells and very few on HLF cells (Fig. 4A). Furthermore, very few of the beads conjugated with CONP were observed on both HLF and HUVEC cells (Fig. 4A). These results indicated that the VRBP1 could be used as the targeting molecules for tumor vessels in which VEGFR2 was over-expressed.

**VRBP1 could Increase the Contrast Effect of MRI Examination**

For verifying that the VRBP1 could work as the guiding molecule of imaging agents for cancer diagnosis, we conjugated VRBP1 and CONP with SPIO and incubated those particles with cells. The changes of magnetic signals on cells were detected with MRI machine. SPIO can produce a predominant T2 relaxation effect. SPIO-VRBP1 and SPIO-CONP at various concentrations were internalized by HUVEC cells in vitro and monitored by MRI. As shown in Fig. 4B, the T2-weighted signal intensity decreased with the increase of the iron concentration. The T2-weighted images of HUVEC cells incubated with SPIO-VRBP1 appeared to be much darker than those of incubated with SPIO-CONP. Furthermore, the T2-weighted images of the HLF cells incubated with SPIO-VRBP1 showed non-changeable signals. The MRI results suggested that VRBP1 could increase the particles uptake efficiency of HUVEC cells when particles were conjugated and VRBP1 might work as effective guiding molecules for blood vessels and tumors which over-expressed VEGFR2.

**VRBP1 could Competitively Bind to VEGFR2 with VEGF Examined by SPR Method and ELISA**

The original SPR readout for binding of 3 μg/mL VEGF with VEGFR2 was 200.8 RU (Fig. 5A, red line). When 10 μg/mL VRBP1 was introduced into the binding system before VEGF was added to the channel, the SPR readout of VEGF-VEGFR2 binding was decreased by 40 RU (Fig. 5A). If 3 μg/mL VEGF was introduced into the reaction channel first before the addition of 10 μg/mL VRBP1, the SPR readout of VRBP1-VEGFR2 binding was decreased by around 34 RU (from 77.9 RU to 46.1 RU) (Fig. 5B). The results might imply that VRBP1 bound with VEGFR2 at the same or adjacent position where VEGF bound with VEGFR2, or at other locales which could interfere the binding of VEGF to VEGFR2. The results of ELISA experiment confirmed that VRBP1 could antagonize the binding of VEGF to VEGFR2 (Fig. S2).

**VRBP1 Specifically Inhibited the Proliferation of Vascular Endothelial Cells in vitro**

The proliferation of endothelial cell proliferation is dependent on VEGF. Our data showed that the selected peptide VRBP1 could block VEGF binding to VEGFR2, therefore, it was necessary to examine the effect of VRBP1
Fig. (5). VRBP1 blocked VEGF binding to its receptor VEGFR2 analyzed by SPR experiment. (A) VRBP1 at concentration of 10 μg/mL was introduced into the SPR reaction channel followed the binding of VEGF to VEGFR2. (B) VEGF at concentration of 3 μg/mL was introduced into the SPR reaction channel followed the binding of VRBP1 to VEGFR2.

Fig. (6). VRBP1 could inhibit proliferation of HUVEC cells induced by VEGF at higher concentration. HUVEC cells treated with 30 ng/mL VEGF were separately co-incubated with CONP, VRBP1, and VRBP2 at various concentrations. Data were presented as mean ± S.D. of three independent experiments. Significance indicated by: *P<0.05 and **P<0.01 in comparison to VEGF treated HUVEC cells by Student’s t-test; †P<0.05 and ‡P<0.01 in comparison to corresponding concentration of CONP co-incubated HUVEC cells by Student’s t-test.
and VRBP2 on the proliferation of HUVEC cells which over-expressed VEGFR2. As shown in Fig. 6, VRBP1 significantly suppressed the mitogenic response of rhVEGF 165 to HUVEC cells at a concentration of 50 μg/mL. The less inhibitory effect on proliferation of HUVEC cells was observed by VRBP2 treatment. In order to verify the growth inhibition of VRBP1 that we observed in HUVEC cells, we performed similar experiments on HLF cells. The proliferation of HLF cells was not affected by VRBP1 (Fig. S3), indicating that the growth inhibitory effects of VRBP1 to HUVEC cells were not due to its potential cytotoxicity and the effect of peptides on HUVEC cells proliferation might be due to the blockade of the VEGF-VEGFR2 pathway by peptides.

**VRBP1 Suppressed Tumor Growth and Angiogenesis in vivo**

To examine the effect of VRBP1 on tumor growth and angiogenesis in vivo, nu/nu mice with H460 tumor xenograft were used. One day after tumor xenograft was transplanted to mice, 100 μL VRBP1 (400 μg/mL) or CONP (400 μg/mL) or physiological saline was injected to tumor sites. The growth of tumors was assessed every other day. Our results (Fig. 7A) indicated that VRBP1 could decrease the growth of tumor after 15 days compared with CONP and physiological saline group. The average tumor size in VRBP1 treated mice was around 75.47 mm³, compared with 172.61 mm³ in CONP group and 220.84 mm³ in physiological saline group at 21 days post injection (Fig. 7B), which indicated VRBP1 could decrease tumor size significantly compared to CONP. The effects of VRBP1 on angiogenesis were examined by immunohistochemical experiment. Serial sections from the solid tumors of nude mice with different treatments were incubated with CD31 antibody. The immunohistochemical results showed that there were more CD31 positive vessels presented in the physiological saline and CONP-treated tumor xenograft slides than those in the VRBP1 group (Fig. 7C). Furthermore, sections from VRBP1 treated mice had a mean microvessel count of 221 per 200× field. For those sections from physiological saline and CONP treated mice, the corresponding value was 576 and 506 per 200× field (Fig. 7D).

**DISCUSSION**

In our study, we successfully screened and selected the specific binding peptides for VEGFR2 with bacterial display method. Selected peptides in two different forms, peptides-bacteria and free peptides were characterized through examining their specificity and affinity to VEGFR2. VRBP1 obtained from focused library had higher affinity (K_D value was 228.3 nM) to VEGFR2 compared with the peptide (MAKGDFFERWGVKEM) (K_D value was 574.7 nM) from random library by the SPR assay (data not shown), which indicated that focused library with conserved sequences could promote to get better binding peptides for VEGFR2. Although the K_D value of peptide VRBP1 was not higher than that of antibodies, it was still in the acceptable range for the research and clinical usage. It could be understood since the molecular weight of peptides is smaller than that of antibodies, the force of hydrophobic between peptides and proteins may be weaker than that between antibodies and proteins [39]. Although the affinity between VRBP1 and VEGFR2 was less strong than that between VEGF and VEGFR2 [40], the competitive binding between VRBP1 and VEGF to VEGFR2 was observed from SPR assay and
confirmed by ELISA experiment. These results implied that VRBP1 might bind to VEGF or VEGFR2 on the same or adjacent position as VEGF, which need to be verified in future. Through blocking the interaction between VEGF and VEGFR2, VRBP1 could defer the growth velocity of HUVEC cells, which meant that VRBP1 had the ability to block the function of VEGF on promoting the proliferation of HUVEC cells. Moreover, the tumor size and number of newly generated tumor vessels in the tumor xenograft models were reduced after the mice were injected with VRBP1. The results from these functional assays further proved that VRBP1 and VEGF competitively interacted with VEGFR2. Furthermore, we presented that VRBP1 could not only be used as therapeutic drugs for cancer patients potentially through inhibiting the growth of new vessels for cancers, but also for locale diagnosis of cancers after conjugating with SPIO.

Besides, peptides obtained from our study which could be used for cancer diagnosis and therapy, two robust tools were developed for the biotechnological research: 1. Affinity evaluation could be performed directly with the peptides on the surface of bacteria; 2. SPR method might substitute ELISA to examine the competitive relationship between various molecules.

Normally, isothermal titration calorimetry (ITC) [41], microscale thermophoresis (MST) [42, 43] and SPR [44] were often used to measure the $K_D$ value of one molecule to the responding molecule. However, the successful measurement of these methods needs sophisticated machines and skillful technicians. In the preliminary section of the whole study, especially when the measurement objectives were very diverse, these methods were not suitable considering the cost and time. In our display system, peptides were displayed on the surface of bacteria, which could be considered as organism matrix. Provided that the expression level of any selected peptides on the bacteria surface was similar, when peptides-bacteria incubated with different concentrations of targeted fluorescent proteins, the fluorescent signals of bacteria had positive correlation with the binding rate between peptides and proteins. When we firstly designed this assay, the only thing expected was to get the relative comparative data about binding efficiency among different peptides with target proteins. Interestingly, from our data, the $K_D$ values for selected peptides-bacteria were in the same mathematical level compared with those of same selected synthesized peptides from SPR measurement. It might prompt us that we could use this novel method to evaluate the $K_D$ values of peptides to proteins.

As for the second useful tool developed from our study, we could use SPR method to substitute ELISA to examine the competitive relationship between different molecules to the same target. Antibodies and luminescent materials are the essential composites for ELISA experiments which implied that proteins/peptides need to be either labeled or used as luminescent antibodies in the system [45]. It is well known that SPR could be used to measure $K_D$ values between two different molecules without any labels [46]. In theory, it could be used to examine the relationships between the two molecules when they could bind with the same target. However, there is no such report till now. In our study, we introduced the second molecule into the reaction channel when the first molecule finished interacting with the target protein without starting the washing procedure. The decreased reaction unit (SPR readout) presented between the second molecule and target proteins meant that these two molecules had competitive relationship binding to the target molecule. In our study, results from ELISA experiment and functional assays related to the peptides verified the validity of this method.

**CONCLUSION**

Taken together, in our study, peptides obtained from bacterial display methods could serve as powerful tools for both cancer diagnosis and therapy. Further verification of application for those peptides in clinic needs to be done in future. Besides that, two biotechnological tools developed would benefit the research in this area. We do hope that results from our study would have profound scientific and applicable value in both research and clinical fields.

**SUPPLEMENTARY MATERIAL**

Supplementary material is available on the publisher’s web site along with the published article.

**CONFLICT OF INTEREST**

The authors confirm that this article content has no conflict of interest.

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**LIST OF ABBREVIATIONS**

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tbody>
<tr>
<td>BSA</td>
<td>bovine serum albumin</td>
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<tr>
<td>CM</td>
<td>chloramphenicol</td>
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<tr>
<td>DMEM/high</td>
<td>Dulbecco’s modified Eagle’s high glucose medium</td>
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<tr>
<td>DMSO</td>
<td>dimethyl sulfoxide</td>
</tr>
<tr>
<td>EDC</td>
<td>1-ethyl-3-(3-dimethylaminopropyl)-carbodiimide</td>
</tr>
<tr>
<td>ELISA</td>
<td>enzyme-linked immunosorbent assay</td>
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<tr>
<td>FACS</td>
<td>fluorescence-activated cell sorting</td>
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<tr>
<td>FBS</td>
<td>fetal bovine serum</td>
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<tr>
<td>FITC</td>
<td>fluorescein isothiocyanate</td>
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<td>HLF</td>
<td>human lung fibroblast</td>
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<tr>
<td>HUVEC</td>
<td>human umbilical vein endothelial cells</td>
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<tr>
<td>ITC</td>
<td>isothermal titration calorimetry</td>
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<tr>
<td>$K_D$</td>
<td>dissociation constant</td>
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<tr>
<td>LB</td>
<td>lysogeny broth</td>
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References


