Identification of a Met-Binding Peptide from a Phage Display Library

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Abstract

Purpose: Aberrant c-Met expression has been implicated in most types of human cancer. We are developing Met-directed imaging and therapeutic agents.

Experimental Design: To seek peptides that bind specifically to receptor Met, the Met-expressing cell lines ST14 and SK-LMS-1 were used for biopanning with a random peptide phage display library. Competition ELISA, fluorescence-activated cell sorting analysis, an internalization assay, and a cell proliferation assay were used to characterize a Met-binding peptide in vitro. To evaluate the utility of the peptide as a diagnostic agent in vivo, 125I-labeled peptide was injected i.v. into nude mice bearing s.c. xenografts of the Met-expressing and hepatocyte growth factor (HGF)/scatter factor-expressing SK-LMS-1/HGF, and total body scintigrams were obtained between 1 and 24 h postinjection.

Results: One Met-binding peptide (YLFSVHWPLKA), designated Met-pep1, reacts with Met on the cell surface and competes with HGF/scatter factor binding to Met in a dose-dependent manner. Met-pep1 is internalized by Met-expressing cells after receptor binding. Met-pep1 inhibits human leiomyosarcoma SK-LMS-1 cell proliferation in vitro. In SK-LMS-1 mouse xenografts, tumor-associated activity was imaged as early as 1 h postinjection and remained visible in some animals as late as 24 h postinjection.

Conclusions: Met-pep1 specifically interacts with Met; it is internalized by Met-expressing cells and inhibits tumor cell proliferation in vitro; it is a potential diagnostic agent for tumor imaging.
panning approach on intact cells. Specificity and affinity of this peptide were examined by a number of in vitro assays. Nuclear imaging of the radiolabeled peptide in mouse xenograft model exhibited tumor-associated activity, recommending this peptide as a promising candidate for future clinical applications.

Materials and Methods

Reagents and cell lines. The Ph.D.-12 Phage Display Peptide Library kit was purchased from New England Biolabs. NaI [480-630 MBq (13-17 mCi)/0.53 mmol/L] EDTA for 5 min at 37°C were spun down and washed thrice with PBS. Cell-bound phage were for the subsequent rounds) at RT with gentle agitation. The cells were spun down, and the depleted phage supernatant was transferred displayed phage (1.5 × 10^12 phage/mL) and were blocked with blocking buffer (PBS with 1% bovine serum albumin). Ph.D.-12 random peptide–displayed phage (1.5 × 10^11 plaque-forming unit) were incubated with 1.0 × 10^7 NIH 3T3 cells for 30 min at room temperature (RT). Cells were spun down, and the depleted phage supernatant was transferred to blocked S114 cells (1.0 × 10^6 cells) and incubated for 1 h (30 min for the subsequent rounds) at RT with gentle agitation. The cells were spun down and washed thrice with PBS. Cell-bound phage were eluted by incubation with 0.5 mL PBS containing 0.05% trypsin and 0.53 mmol/L EDTA for 5 min at 37°C. The eluted phage were amplified and precipitated according to New England Biolabs' instruction. The second round panning was carried out with SK-LMS-1/HGF cells, the third and fourth round were carried out with S114 and SK-LMS-1/HGF cells, respectively, to ensure the specificity of selected clones.

Phage capture ELISA and competitive ELISA. The specificity of individual peptide-display phage for Met binding was assessed by phage ELISA. Human Met-Fc fusion protein (R&D Systems) was coated at a final concentration of 0.4 μg/mL. Single plaques of phage were picked up randomly and amplified in 2 mL of ER2738 culture for 4.5 h. The culture supernatant was added to the coated plates and incubated at RT for 1 h, followed by horseradish peroxidase–conjugated anti-M13 IgG (Amersham), and the plates were incubated for 1 h at RT. After washing, TMB substrate (Pierce) was used according to manufacturer's instructions. For competitive ELISA, HGF samples (25 μg/mL) mixed with or without phage clones (2 × 10^11 plaque-forming unit per mL) at 2 × 10^12 plaque-forming unit per mL, with or without Met- pep1 (10, 100, 500 μm/L) were incubated on the Met-coated plates for 1.5 h at RT and detected by an in-house rabbit anti-HGF polyclonal antibody. Nonavid control peptide was tested at the concentration of 500 μm/L. Alkaline phosphatase–conjugated antirabbit IgG (Sigma) were added and phosphatase substrate CP-nitrophenyl phosphate (Kirkgaard & Perry Laboratories) was added for 30 min, and absorbance was measured at 405 nm.

DNA sequencing and peptide synthesis. Single-stranded phage DNA was prepared from ELISA-positive clones as described in the phage display peptide library manual. DNA sequencing was carried out in a 3700 DNA Analyzer (Applied Biosystems) following the manufacturer's instructions.

Peptides representing the sequence of positive clone 30 (YLFSVH-WPPLKA) and nonavid peptide (ASYPFILVKLWHP) and their FITC and biotin conjugates were synthesized by Genemed Synthesis, Inc. For fluorescence-activated cell sorting analysis, DU145, MKN45, NIH 3T3, and PC3 cells (1.0 × 10^4) were detached with TrypLE Express (Invitrogen) and were incubated with Met- pep1-FITC or FITC-labeled nonavid peptide at a final concentration of 10 μg/mL for 15 min at 4°C. The cells were then washed twice and suspended in 500 μL PBS. Flow cytometry was done with a fluorescein-activated cell sorting Calibur cytometer and the CellQuest analysis program (Becton Dickinson).

Fluorescent images. Mixed MKN45 and NIH 3T3 cells were cultured in eight-chamber slides. After fixing with 10% formalin, the cells were blocked and incubated with Met- pep1-biotin or nonavid peptide-biotin (50 μg/mL) and a specific anti-Met murine monoclonal antibody, Met4 (20 μg/mL), for 1.5 h at RT. After washing, fluorescein (DTAF)-conjugated streptavidin (Jackson ImmunoResearch) and antiserum IgG–TRITC (Jackson ImmunoResearch) were added to cells and incubated for 1.5 h at RT. Nuclei were stained with 4′,6-diamidino-2-phenylindole.

Affinity measurement. The binding affinity of Met- pep1 to Met was measured by KinExA 3000. Serially diluted SK-LMS-1 cells (1:4 dilutions, starting from 16 × 10^4 cells per mL for curve 1 and 4 × 10^6 cells per mL for curve 2) were incubated with constant Met- pep1-FITC at 500 nmol/L for curve 1 or 100 nmol/L for curve 2 overnight until equilibrium was reached. Cells were removed by centrifugation. Unbound Met- pep1-FITC was retrieved by passing the supernatant fractions through Met-coated beads. Multiple curve analysis was done to determine optimal values for the equilibrium dissociation constant K_d by using the KinExA Pro program.

Internalization. S114 and NIH 3T3 cells were plated in eight-chamber slides. Met- pep1-biotin or nonavid peptide-biotin was added to the medium to a final concentration of 25 μm/L for 1 or 16 h. The cells were then washed and fixed with acetone/methanol (50:50) for 10 min. After blocking, the slides were incubated with Met4 at 20 μg/mL for 1.5 h at 37°C. After washing with PBS twice, the cells were incubated with a mixture of either antiserum FITC conjugate and streptavidin–rhodamine red conjugate or DTAF-streptavidin and anti-mouse-TRITC for 1.5 h at RT. 4′,6-Diamidino-2-phenylindole was used to stain nuclei. The slides were examined by confocal laser scanning microscopy.

Cell proliferation assay. SK-LMS-1 cells were seeded in 96-well plates (2,000 cells per well) and incubated for 24 h. After 12 h of serum starvation, Met- pep1 or nonavid control peptide (100 μm/L) was added to the medium and the cells were incubated for 4 days. The cultures were harvested and analyzed by flow cytometry. The percentage of cells in each cell cycle phase was determined using the ModFit program.

Fig. 1. Schematic representation of the experimental design.
added at different concentrations, with or without HGF/SF (100 ng/mL), and incubated for 12 h. Finally, [3H]thymidine was added for 4 h (0.5 mCi per well; PerkinElmer). [3H]Thymidine incorporation was measured using a scintillation counter (MicroBeta TriLux, PerkinElmer).

**Tumor xenographs and nuclear imaging with Met-pep1.** SK-LMS-1/HGF cells were used for xenograph induction. Female athymic nude (nu/nu) mice at 2 months of age received s.c. injections of the cell suspension (5 × 10^6 cells) in the posterior aspect of the right thigh. Tumors developed for 3 to 4 weeks, reaching 0.5 cm or more in greatest dimension by external caliper measurement before imaging. Mice were housed in small groups and given ad libitum access to mouse chow and drinking water under conditions approved by the Institutional Animal Care and Use Committee. For nuclear imaging experiments, the synthetic Met-avid peptide Met-pep1 or the nonavid control peptide was dissolved in 0.25 mol/L sodium phosphate buffer (pH 7), stored frozen in small aliquots, and thawed just before labeling. Protein radioiodination was done essentially as previously described (46).

Animals were imaged, and scintigams were analyzed by established methods (21, 46). In brief, each mouse received 125I-labeled Met-pep1 (77 μCi, 0.1 mL) or control nonavid peptide (124 μCi, 0.1 mL) i.v. by tail vein injection under light inhalational anesthesia. Just before each imaging session, each mouse was given up to 13 mg/kg xylazine and 87 mg/kg ketamine s.c. in the interscapular region. Serial posterior whole-body gamma camera images of each mouse were acquired overnight; Fig. 3C). Cells were removed by centrifugation. The percentage of Met-pep1-FITC free in the supernatant was measured by passing the supernatant through Met-coated beads. Multiple curve analysis was done to determine optimal values for the equilibrium dissociation constant K_d, which was 64.2 nmol/L, calculated by n-curve analysis using the KinExA Pro program.

**Met-pep1 internalizes Met-expressing cells via receptor binding.** We examined the internalization of Met-pep1 by S114 cells.

### Table 1. Summary of Met-binding clones

<table>
<thead>
<tr>
<th>Phage clone</th>
<th>Peptide sequence</th>
<th>% Competition</th>
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<tr>
<td>2</td>
<td>QHKTSTIGGHHLEP</td>
<td>28.6</td>
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<tr>
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<td>TLPSPLALLTVH</td>
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<tr>
<td>30</td>
<td>YLFSVHWPPLKA</td>
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</table>

**Results**

**Identification of peptide binders against human Met using phage display.** A schematic representation of the experimental design is shown in Fig. 1. A 12-mer random peptide phage display library was screened with Met-expressing S114 and SK-LMS-1/HGF cells alternately to seek Met-binding peptides. NIH 3T3 cells were used for negative selection to remove nonspecific binders. After four rounds of biopanning, 30 phage clones were picked randomly for screening, all of which were found reactive against Met recombinant protein. They were then tested by competitive ELISA with or without HGF for Met binding. Three clones with the ability to compete with HGF in a dose-dependent manner for Met binding were sequenced (Fig. 2A; Table 1). Clone 30 (YLFSVHWPPLKA) displayed the highest competitive ability (59.3% inhibition as summarized in Table 1). The synthetic peptide of clone 30 (designated Met-pep1) also showed the competition ability to HGF for Met-binding in a dose-dependent fashion (Fig. 2B). At the concentration of 100 and 500 μmol/L, Met-pep1 competed 30% and 50% of HGF at 25 μg/mL for Met-binding. The nonavid peptide showed no competitive effect on HGF/Met binding (Fig. 2B).

**Met-pep1 binds Met on a live or fixed cell surface.** The specific binding activity of Met-pep1 to Met on live Met-expressing cells was determined by fluorescence-activated cell sorting analysis. FITC-conjugated Met-pep1 specifically binds Met-expressing DU145, MKN45, and PC3 cells (Fig. 3A). Very low level of binding was observed in NIH 3T3 cells. FITC-labeled nonavid peptide did not show any binding in all cells.

Since MKN45 and NIH 3T3 cells display different morphologies in culture flasks, we mixed these two cell lines when plating them in eight-chamber slides. The colocalization of Met-pep1 and an anti-Met-specific monoclonal antibody Met4 in fluorescent staining indicated that Met-pep1 binds specifically to Met on formalin-fixed MKN45 cells, but not on NIH 3T3 cells (Fig. 3B).

To determine the affinity of Met-pep1, serially diluted SK-LMS-1 cells (1:4 dilution, starting from 16 × 10^6 cells per mL or 4 × 10^6 cells per mL) were incubated with constant Met-pep1-FITC (500 nmol/L, diamonds; 100 nmol/L, closed circles) until equilibrium was reached overnight. Cells were removed by centrifugation. The percentage of Met-pep1-FITC free in the supernatant was measured by passing the supernatant through Met-coated beads. Multiple curve analysis was done to determine optimal values for the equilibrium dissociation constant K_d, which was 64.2 nmol/L, calculated by n-curve analysis using the KinExA Pro program.

**Met-pep1 internalizes Met-expressing cells via receptor binding.** We examined the internalization of Met-pep1 by S114 cells.

**Fig. 2. Identification of Met-binding peptides. A, phage competitive ELISA.** HGF samples (25 μg/mL) mixed with or without phage clones were incubated on the Met-coated plates and detected by a rabbit anti-HGF polyclonal antibody. Column, A_{405} mean of duplicate samples, bars, SD.
through endocytosis (Fig. 4). After incubation of S114 cells and NIH 3T3 cells with biotin-conjugated Met-pep1 for 1 and 16 h, cells were fixed and stained with fluorescein-conjugated or rhodamine red-conjugated streptavidin. Anti-Met mAb Met4 was used as a positive control for immunostaining. The green and red colors were found colocalized in S114 cells, suggesting that Met-pep1 was internalized by S114 cells via receptor binding. Internalization of Met-pep1 into S114 cells could be detected as early as 1 h, remained visible at 16 h. No internalization of Met-pep1 was found in NIH 3T3 cells. No internalization of the nonavid control peptide on both cell lines was detected (data not shown).

Met-pep1 specifically inhibits the proliferation of human leiomyosarcoma cells. The exposure of Met-expressing cells to HGF/SF can elicit a variety of cellular responses including proliferation. We tested the inhibitory activity of Met-pep1 in HGF-mediated cell proliferation assay using SK-LMS cells. The cells were stimulated with HGF for 12 h before [3H]thymidine was added; then [3H]thymidine incorporation was measured. The results showed that 100 μmol/L of Met-pep1 clearly inhibited cell proliferation when the cells were treated with 100 ng/mL of HGF (Fig. 5). The \( P \) value was 0.017, calculated from Student's \( t \) test, indicating the differences were significant.

**In vivo nuclear imaging with Met-pep1.** As an additional functional test of Met-pep1 and as the first step in evaluating it as a potential *in vivo* diagnostic agent, we have conducted preliminary nuclear imaging experiments with radiiodinated Met-pep1 in nude mice bearing xenografts of the autocrine Met-HGF/SF–expressing human leiomyosarcoma cell line SK-LMS-1/HGF (Fig. 6). Five female athymic nude mice bearing s.c. xenografts of human leiomyosarcoma SK-LMS-1/HGF in the posterior aspect of the right thigh were each injected i.v. with 77 μCi \( {^{125}}\text{I} \)-Met-pep1, whereas an additional five xenograft-bearing mice were each injected i.v. with 124 μCi \( {^{125}}\text{I} \)-labeled nonavid control peptide. Sequential posterior total body nuclear images of each group of mice were obtained 1 to 24 h postinjection (Fig. 6A). Tumor-associated activity was imaged as early as 1 h after injection of \( {^{125}}\text{I} \)-Met-pep1 and remained visible in some animals as late as 24 h postinjection. Activity was highest in the gastric region (g) at all imaging time
points, whereas thyroid (t) activity increased over time. Activity was also evident in the nasopharyngeal (n) region and in the urinary bladder (b). The activity in the lower midline at 4 and 7 h postinjection represents the urinary bladder (b). By quantitative region-of-interest analysis (Fig. 6B), mean tumor-associated radioactivity accounted for 4% to 5% of the estimated injected activity of $^{125}$I-Met-pep1 at 1 and 4 h and declined to ~3% by 7 h postinjection. In contrast, mean tumor-associated radioactivity for host mice injected with $^{125}$I-labeled nonavid control peptide was only about one-third of the respective Met-pep1 values at all postinjection time points.

**Discussion**

There has been an exponential growth in the development of radiolabeled peptides for diagnostic and therapeutic applications in oncology. Peptide ligands, which have the advantages of low immunogenicity, easy incorporation into certain delivery vectors, and being readily diffusible, are being pursued as targeting moieties for the selective delivery of radionuclides,
cytokines, chemical drugs, or therapeutic genes to tumors (47). Faster blood clearance, rapid tissue penetration, and higher target-to-background ratios have been attained for peptides relative to macromolecular compounds in nuclear imaging studies. Radiolabeled receptor-binding peptides have become an accepted new class of radiopharmaceuticals (48, 49).

In this paper, we have shown through multiple approaches that Met-pep1 binds specifically to the extracellular domain of human Met, as expressed on the surface of human cancer cells in vitro and in vivo, and our data indicate that it must bind near the HGF/SF ligand binding site. Met-pep1 is internalized after its binding to Met, as evidenced both by direct imaging in vitro and by the liberation of free iodide in vivo.

To begin evaluating the diagnostic potential of Met-pep1, we conducted preliminary nuclear imaging experiments with radiiodinated Met-pep1 in nude mice bearing xenografts of the autocrine Met-HGF/SF–expressing human leiomyosarcoma cell line SK-LMS-1/HGF. We previously used this model system to evaluate full-length murine anti-Met mAbs (Met3 and Met5) as potential radiopharmaceuticals (21, 22, 46), as well as a human Fab fragment that specifically recognizes Met (27). Consistent with the general observation that peptides display a lower affinity for a given antigen, the tumor-associated radioactivity values are lower than we reported for the radiolabeled full-length murine anti-Met mAbs and human Fab (21, 22, 27, 46), but nonetheless are higher than the levels we found for either an irrelevant full-length mAb in the same model (21) or for a nonavid control dodecameric peptide (Fig. 6B).

In conclusion, due to its specific binding to Met-expressing tumor cells in vitro and in vivo together with internalization, Met-pep1 represents a promising candidate useful for clinical Met-directed diagnostic and therapeutic applications. Coupling cytotoxic drugs to macromolecular carriers has been shown to be a promising approach for efficient drug targeting. To further develop the use of this peptide in drug delivery, conjugation of toxins to Met-pep1 and their activities in vitro assays are now under way in our laboratory.

Acknowledgments

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References

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