Ultrasound molecular imaging as a non-invasive companion diagnostic for netrin-1 interference therapy in breast cancer

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Abstract

In ultrasound molecular imaging (USMI), ligand-functionalized microbubbles (MBs) are used to visualize vascular endothelial targets. Netrin-1 is upregulated in 60% of metastatic breast cancers and promotes tumor progression. A novel netrin-1 interference therapy requires the assessment of netrin-1 expression prior to treatment. In this study, we studied netrin-1 as a target for USMI and its potential as a companion diagnostic in breast cancer models.

Methods: To verify netrin-1 expression and localization, an in vivo immuno-localization approach was applied, in which anti-netrin-1 antibody was injected into living mice 24 h before tumor collection, and revealed with secondary fluorescent antibody for immunofluorescence analysis. Netrin-1 interactions with the cell surface were studied by flow cytometry. Netrin-1-targeted MBs were prepared using MicroMarker Target-Ready (VisualSonics), and validated in in vitro binding assays in static conditions or in a flow chamber using purified netrin-1 protein or netrin-1-expressing cancer cells. In vivo USMI of netrin-1 was validated in nude mice bearing human netrin-1-positive SKBR7 tumors or weakly netrin-1-expressing MDA-MB-231 tumors using the Vevo 2100 small animal imaging device (VisualSonics). USMI feasibility was further tested in transgenic murine FVB/N Tg(MMTV/PyMT634Mul) (MMTV-PyMT) mammary tumors.

Results: Netrin-1 co-localized with endothelial CD31 in netrin-1-positive breast tumors. Netrin-1 binding to the surface of endothelial HUVEC and cancer cells was partially mediated by heparan sulfate proteoglycans. MBs targeted with humanized monoclonal anti-netrin-1 antibody bound to netrin-1-expressing cancer cells in static and dynamic conditions. USMI signal was significantly increased with anti-netrin-1 MBs in human SKBR7 breast tumors and transgenic murine MMTV-PyMT mammary tumors compared to signals recorded with either isotype control MBs or after blocking of netrin-1 with humanized monoclonal anti-netrin-1 antibody. In weakly netrin-1-expressing human tumors and normal mammary glands, no difference in imaging signal was observed with anti-netrin-1- and isotype control MBs. Ex vivo analysis confirmed netrin-1 expression in MMTV-PyMT tumors.

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Conclusions: These results show that USMI allowed reliable detection of netrin-1 on the endothelium of netrin-1-positive human and murine tumors. Significant differences in USMI signal for netrin-1 reflected the significant differences in netrin-1 mRNA & protein expression observed between different breast tumor models. The imaging approach was non-invasive and safe, and provided the netrin-1 expression status in near real-time. Thus, USMI of netrin-1 has the potential to become a companion diagnostic for the stratification of patients for netrin-1 interference therapy in future clinical trials.

Key words: ultrasound molecular imaging, netrin-1, companion diagnostic, breast cancer, targeted microbubbles

Introduction

Netrin-1 is a laminin-related protein involved in axon outgrowth and tumorigenesis [1–4]. Netrin-1 was shown to be overexpressed in 60% of metastatic breast cancers [5]. Following extensive preclinical studies [6,7], a novel therapeutic approach based on netrin-1 interference recently entered a Phase 1 clinical trial to evaluate safety, pharmacokinetics, and clinical activity of a humanized monoclonal antibody targeting netrin-1 (NET1-H-mAb, NP137, Netris Pharma, Lyon, France) in patients with advanced solid tumors (NCT02977195). A companion diagnostic is now required for patient stratification. Although netrin-1 is a secreted protein, its detection in blood serum samples is not feasible. To avoid invasive biopsy-based techniques, a non-invasive molecular imaging approach was developed. As netrin-1 is expressed on endothelial cells interacting among others with CD146 [8–11], we hypothesized that netrin-1-overexpressing tumors might present netrin-1 specifically on the tumor endothelium, providing the opportunity to develop an intravascular molecular imaging approach.

Ultrasound molecular imaging (USMI) can provide a non-invasive, cost-effective, image-based companion diagnostic for the novel netrin-1 interference therapy, as it combines the advantages of ultrasound, such as real-time anatomical imaging, with the capabilities of molecular imaging, such as high sensitivity and specificity in vivo. Anatomical and functional ultrasound imaging are integral parts of clinical breast cancer screening and diagnosis complementing mammography, guiding core needle biopsies, and detecting abdominal metastases for the management of breast cancer [12]. Though, ultrasound imaging is limited to soft tissues and cannot be applied to bone or lung metastases. Ultrasound imaging has become molecularly specific with the development of ultrasound contrast agents (UCAs) targeted with ligands such as antibodies or other proteins to detect expression of cancer-specific molecular markers on the vascular endothelium [13–15], such as CD276 (or B7-H3) in ovarian and breast cancer, or secreted frizzled related protein 2 in angiosarcoma [16–22], to allow cancer early detection, prediction of therapy response, and treatment follow-up [23–29]. Recent clinical demonstration of feasibility and safety of VEGFR2 imaging in breast, ovarian, and prostate cancer patients with the first clinical grade UCA (BR55) has opened the way for further clinical translations of USMI [30,31]. Our objective was to extend USMI to the novel cancer biomarker netrin-1 with the purpose of developing a unique, non-invasive approach to detect netrin-1 overexpression in breast cancer, eventually enabling patient-based treatment decisions.

Here we report the vascular presentation of netrin-1 by endothelial cells in vivo in netrin-1-positive breast tumor models and show the feasibility of ultrasound molecular imaging of netrin-1 to discriminate between strongly and weakly netrin-1-expressing tumors. We showed that (Figure 1): A) netrin-1 was presented by endothelial cells as assessed by in vivo immuno-localization; B) netrin-1-targeted microbubbles selectively bound to netrin-1 protein and netrin-1-presenting cells in vitro; C) in vivo imaging of netrin-1 revealed significantly enhanced signal with netrin-1-targeted MBs compared with isotype control MBs in nude mice bearing human strongly netrin-1-positive SKBR7 breast tumors and in transgenic mice developing spontaneous mammary adenocarcinoma (MMTV-PyMT), while there was no significant difference between targeted and control contrast agents in nude mice bearing weakly netrin-1-expressing human MDA-MB-231 breast cancer xenografts and in normal wild type mammary glands [5].

Methods

Animal models

All procedures involving the use of laboratory animals were approved by the Institutional Administrative Panel on Laboratory Animal Care at Stanford University. For induction of subcutaneous tumors, female homozygous nude mice (NU/J, The Jackson laboratory) were used and 2×10⁶ cells of human strongly netrin-1-positive SKBR7 (N=9) or
weakly netrin-1-expressing MDA-MB-231 (N=9) breast cancer were subcutaneously inoculated in the hind limbs of mice at 4 weeks of age [5]. After 2 weeks, mice underwent USMI and in vivo immuno-localization of netrin-1. Female transgenic breast cancer-bearing (N=30) FVB/N-Tg(MMTV-PyMT)634Mul/J mice (MMTV-PyMT) and female, transgene-negative control littermates (N=15) were used at 8 weeks of age for USMI and in vivo immuno-localization of netrin-1. At this point of disease progression, female MMTV-PyMT mice were bearing invasive mammary carcinoma [21,32,33]. For netrin-1 and Fc receptor blocking experiments, mice were injected 100 µg of NET1-H-mAb 24 h prior to imaging.

Figure 1. Study design. (A) In vivo immuno-localization (IVIL) analysis of endothelial expression of netrin-1. Netrin-1 was labeled in vivo and NET1-H-mAb and CD31 were revealed ex vivo. (B) Binding of targeted MBs to purified netrin-1 protein and netrin-1-overexpressing cells was evaluated under static conditions (by upright incubation in 24-well plates incubated on a rocking platform) or dynamic conditions (by inverted incubation with circulating MBs in a flow chamber). (C) Mouse models of human and murine breast cancer were used for in vivo imaging of netrin-1. Here, the MMTV-PyMT mouse lying on its back shows five mammary glands on each lateral side of the body. Anti-netrin-1-MBs were intravenously injected and imaged with an ultrasound imaging probe. Signal of bound MBs was quantified using the destruction-replenishment method.
In vivo immuno-localization (IVIL) staining of netrin-1 in endothelium

For netrin-1 immunostaining, a previously described IgG localization technique, here called IVIL, was applied, as standard ex vivo immunofluorescence staining was prevented by the specificity of NET1-H-mAb to native state netrin-1 as opposed to fixed netrin-1, which undergoes conformational changes and antigen masking during tissue fixation [32]. Targeted and control antibodies show passive accumulation in tumor tissues due to the enhanced permeability and retention effect and Fc-mediated molecular interactions [34–39]. To assess the degree of targeted accumulation, fluorescence signal from isotype control antibody and NET1-H-mAb were compared. After USMI with netrin-1-targeted and isotype control contrast agents and 24 h prior to tumor or normal mammary gland collection, living nude mice bearing subcutaneous human SKBR7 and MDA-MB-231 tumors, or living FVB/N-Tg(MMTV-PyMT)634Mul/J (MMTV-PyMT) mice bearing invasive mammary carcinoma and wild type mice with normal mammary glands were intravenously injected with either 100 µg of primary NET1-H-mAb or 100 µg of human IgG isotype control antibody (NB5P-97043, Novus Biologicals, Littleton, CO, USA). To remove freely circulating antibody, cardiac perfusion was performed with PBS, tumor tissues or mammary glands were isolated, flash frozen, and sectioned at 15 µm thickness on a cryostat. Endothelial cells were labeled with primary rat anti-mouse CD31 antibody (550274, BD Biosciences), followed by secondary Alexa 488-coupled goat anti-rat IgG (A11006, Life Technologies). No additional primary antibody targeted to netrin-1 was applied. NET1-H-mAb was revealed with Alexa Fluor 594-coupled goat anti-human IgG (A11014, Life Technologies). Stained sections of a maximum number of tumor samples and normal mammary glands were imaged on a LSM 510 confocal microscope (Zeiss) on the same day to allow relative quantification of fluorescence staining. Fluorescence images were analyzed with FIJI (Fiji Is Just ImageJ) software [40]. Endothelial presence of netrin-1 was indicated by co-registration of fluorescence signals for netrin-1 (Alexa Fluor 594) and CD31 (Alexa Fluor 488).

HeLa model cell lines

Generation of HeLa model cell lines either netrin-1-negative (HeLa-Ctrl) or netrin-1-positive (HeLa-Net1) were described previously [6]. The self-inactivating HIV-1-derived vector encoding netrin-1 under control of Human Elongation Factor-1 Alpha (EF-1 Alpha) promoter was inserted by lentiviral transduction resulting in HeLa-Net1 cells. To create HeLa-Ctrl, a corresponding mock vector was used for lentiviral transduction.

Flow cytometry analysis of netrin-1 presentation on cell surfaces

Transduced netrin-1-positive (HeLa-Net1) and negative (HeLa-Ctrl) model cells and human umbilical vein endothelial cells (HUVEC) were used. Cell surface netrin-1 presentation was analyzed after heparin incubation, enzymatic digestion of heparan sulfate, and netrin-1 incubation via flow cytometry. For the heparin competition experiment, cells were seeded at 4×10^6 cells/100 cm² flask. One day after seeding, the medium was removed and the flask was rinsed with PBS prior to addition of fresh medium complemented with 50 µg/mL of heparin (Heparin Sodium Salt, A3004, AppliChem). Twelve hours later, cells were collected by mechanical removal using a cell scraper. For enzymatic digestion of heparan sulfate, cells were seeded at 1×10^6 cells/100 cm² flask. An enzyme cocktail composed of Heparinase III (H8891, Sigma), Heparinase I (H2519, Sigma) (both reconstituted in 20 mM Tris-HCl, pH 7.5, 0.1 mg/mL BSA, 4 mM CaCl₂, 150 mM NaCl buffer), Chondroitinase ABC (C2905, Sigma; reconstituted in 0.01% BSA in PBS), and β-Glucuronidase (G0251, Sigma; reconstituted in 0.1% BSA in PBS) was prepared. Cells were collected three days after seeding and were re-suspended in medium containing the enzyme cocktail for 1 h in the cell culture incubator with intermittent mixing. Treatment concentrations of the different enzymes were: Heparinase III, Heparinase I, and Chondroitinase ABC at 0.25 U/1×10^6 cells and β-Glucuronidase at 200 U/1×10^6 cells. For incubation with exogenous netrin-1, HeLa-Ctrl were seeded at 1×10^6 cells/100 cm² flask and HeLa-Net1 at 2×10^6 cells/100 cm² flask. The netrin-1-containing supernatant of HeLa-Net1 and the control supernatant of HeLa-Ctrl were collected 3 days after seeding and sterile-filtrated. HeLa-Ctrl cells were collected 3 days after seeding, and then re-suspended in HeLa-Net1 supernatant or HeLa-Ctrl control supernatant for 1 h in the cell culture incubator with intermittent mixing. For enzymatic digestion of heparan sulfate followed by incubation with exogenous netrin-1, HeLa-Ctrl cells were seeded at 1×10^6 cells/100 cm² flask. Cells were collected 3 days after seeding and re-suspended in medium containing the Heparinase III, Heparinase I, Chondroitinase ABC, and β-Glucuronidase enzyme cocktail for 1 h in the cell culture incubator with intermittent mixing. Next, the supernatant was removed by centrifugation and cells
were coated with different concentrations of netrin-1 in the cell culture incubator with intermittent mixing.

Cells were labeled with NET1-H-mAb, with specificity for human and murine netrin-1, or humanized isotype control antibody (iso-mAb, Netris Pharma, Lyon, France) (2 µg antibody / 1×10^6 cells) [6]. For detection of heparan sulfate, mouse monoclonal anti-heparan sulfate (clone 10E4, Amsbio, Abingdon, UK) or the mouse monoclonal isotype control (clone MM-30, Abcam, Cambridge, UK) was used (1.5 µg antibody / 1×10^6 cells). Secondary PE-coupled anti-human IgG Fc (clone HP6017, BioLegend, San Diego, CA, USA) or Alexa Fluor 647-coupled anti-mouse IgG (A31571, Thermo Fisher, Waltham, MA, USA) was used (1 µg antibody / 1×10^6 cells). Cells were analyzed on a FACS Calibur (BD Biosciences, Franklin Lakes, NJ, USA). For analysis, debris and cell aggregates were excluded by gating on living single cells in FSCxSSC dot plots and 10,000 events were acquired. Fluorescence intensities were analyzed and fluorescence histograms were prepared with FSC Express 4 Flow Research Edition (De Novo Software, Glendale, CA, USA). Fluorescence signal ratios were calculated dividing marker intensities (in a.u.) by the respective isotype control intensity (in a.u.).

**In vitro MB binding assays on purified netrin-1 protein and cell lines**

To test the ability of netrin-1-targeted MBs to bind to recombinant netrin-1 protein, static and dynamic in vitro binding assays were performed as previously described [41,42].

**MB binding to recombinant netrin-1 protein:** To test the binding specificity of netrin-1-targeted MBs, a MB binding assay in 24-well plates under static binding conditions was performed as previously described [41]. Briefly, the wells of a 24-well plate were coated with different concentrations of netrin-1 protein (human recombinant, AdipoGen, San Diego, CA, USA). MBs were diluted to 1.2×10^6 MBs/mL in NaCl 0.9% + BSA 1% in H₂O and 350 µL of MB suspension was used per well. The plate was incubated in upright position on a rocking platform (25 rpm, 5 min) at room temperature so that MBs contacted with the coated well surface by liquid turbulences. Before microscopy (Zeiss Axiocam, Metamorph, Marly le Roi, France), unbound MBs were removed by incubation on a plate shaker (3x, 300 rpm, 10 s). MBs bound to netrin-1 were automatically counted by analyzing photomicrographs with MATLAB 2013 (The MathWorks, Natick, MA, USA) (see Supplementary Material).

**Effect of buffer on MB binding:** To test the effect of buffer on MB binding, a 96-well plate binding assay in inverted position was performed as previously described [41]. Briefly, a 96-well plate was coated at 4 µg/mL of netrin-1. MBs were diluted to 1.2×10^7 MBs/mL either in 1% BSA, 0.9% NaCl in H₂O or in 50% human plasma, 0.09% NaCl in H₂O. MBs were incubated in 96-well plates (350 µL of MB suspension per well) in upside-down position at 4 °C so that MBs got in contact with the coated well surface by buoyancy. Plates were returned to the right-side up position and photomicrographs were recorded with an inverted microscope (Nikon Eclipse Ti-S, Champigny sur Marne, France). This procedure allowed unbound MBs to float above the focal plane, eliminating the need for a separate washing step.

**Effect of temperature on MB binding:** To test the effect of temperature on MB binding, MBs were diluted to 1.2×10^7 MBs/mL in 100% plasma (human Na-Heparin, DivBioScience, Ulvenhout, The Netherlands) and incubated in a netrin-1-coated (4 µg/mL) 96-well plates in upside-down position either at room temperature or at 37 °C for 30 min. Plates were returned to the right-side up position and photomicrographs were recorded with an inverted microscope (Nikon Eclipse Ti-S).

**Effect of shear stress on MB binding:** To study the effect of shear stress on MB binding, a parallel plate flow chamber (gasket B, Glyco Tech, Maryland, USA) was used in combination with a 35 mm-Corning dish that was previously coated with netrin-1 at 4 µg/mL. To seal the chamber a vacuum pump was used (BioBlock, Fisher Scientific, Illkirch, France). The flow chamber had a flow path of 2.5 cm width, 0.254 mm height, and 10 mm length. MBs were diluted to 5×10^6 MBs/mL in 1% BSA, 0.9% NaCl in H₂O. Using a syringe pump (PHD ULTRA, Harvard Apparatus, Les Ulis, France), physiological shear stresses of 1 or 2 dynes/cm², which are equivalent to flow rates of 212 and 424 µL, were created to simulate shear stresses of tumor blood vessels according to previously described protocols [16,23,42]. MBs circulated for 5.5 min in the flow chamber. MB circulation and attachment were recorded via time-lapse video-microscopy (Leitz Laborlux S, Zeiss Axiocam 105 color). Attached MBs in video sequences were automatically counted using MATLAB 2013 (The MathWorks) (see Supplementary Material).

**MB binding on HeLa-Ctrl and HeLa-Net1 cells:** To verify binding of anti-netrin-1 MBs to HeLa-Ctrl and HeLa-Net1 cells, a binding assay in 96-well plate format in inverted position was performed. Cells were seeded at 1×10^4 cells per well. Three days after seeding, functionalized MBs (diluted to 1.2×10^7 MBs/mL in 50% human plasma, 0.9% NaCl in H₂O) were added to the 96-well plate (350 µL/well) and incubated in upside-down position at 4 °C for 30 min.
Plates were returned to the right-side-up position and photomicrographs were recorded with an inverted microscope (Nikon Eclipse Ti-S).

**MB binding on breast cancer cells:** In the study of MB binding to breast cancer cells, the following cell lines were included: human SKBR7 and MDA-MB-231 (kindly provided by P. Mehlen, CRCL, Inserm U1052, Lyon, France). MB binding experiments on breast cancer cell lines were performed under dynamic conditions in flow chambers. Cells were seeded at 4.5×10^5 cells in 35-mm Corning dishes. Two days after seeding, MBs were diluted to 5×10^6 MBs/mL and circulated for 12 min in the parallel plate flow chamber at a shear stress of 1 dyne/cm². Flow chamber assays under the illuminating microscope were performed at a temperature of 25 to 26 °C as monitored by infrared thermometry. Circulating and attached MBs were recorded by time-lapse videomicroscopy and the image sequences were automatically analyzed. Cellular vesicles and MBs were distinguished based on the image analysis depicting cells prior and subsequent to incubation with MBs.

**In vivo imaging of netrin-1**

**Imaging protocol:** Tumors of MMTV-PyMT mice were imaged when they reached approximately 5 mm in diameter. These were compared with age-matched wild type mice with mammary glands of approximately 4 mm in diameter. MMTV-PyMT and wild type mice were depilated using a depilation cream (Nair™ Lotion With Aloe & Lanolin", Church & Dwight Co., Inc., Ewing, NJ, USA). SKBR7 tumors were imaged when they reached approximately 10 mm in diameter and MDA-MB-231 tumors when they reached approximately 8 mm in diameter. Nude mice lacking hair did not require depilation prior to imaging. During imaging, mice were under anesthesia at 2% isoflurane in oxygen at 2 L/min and kept warm using a heated stage and a heat lamp. For coupling of the ultrasound transducer, pre-warmed gel was applied on the skin of the regions to be imaged. The small-animal high resolution imaging system Vevo 2100 (VisualSonics, Toronto, CA) in combination with the MS-250 transducer (center frequency at 21 MHz, lateral resolution of 165 μm and axial resolution of 75 μm, focal length at 8 mm) that was immobilized on an adjustable stand was used for all acquisitions. Scanning through the tumor tissue in ultrasound B-mode, the 2D cross-section showing the tumor or gland at its biggest diameter was identified and selected for image acquisition. Shortly before imaging, MBs were functionalized with biotinylated NET1-H-mAb (MB_NETrin-1) or biotinylated human IgG isotype control antibody (MB_isotype) (NBP-1-96855, Novus Biologicals, Littleton, CO, USA) (50 μg antibody/1.6×10^6 MBs in one vial). The same batch of MBs was used for all *in vivo* imaging studies. A 27 G needle was used for bolus injection of a total of 1×10^8 MBs in 100 μL of PBS via a catheter (orange butterfly) into the tail vein. The catheter was flushed post injection with 50 μL of PBS to ensure the total MB dose was administered. Tumors or normal mammary glands were imaged 4-10 min post intravenous bolus injection of MBs according to previously described protocols [21,24,43]. To image several different tumors in the same mouse after a single MB injection, the transducer was moved from the first tumor to the next and a novel destruction-replenishment acquisition was performed. USMI was performed in contrast mode with an emission frequency of 21 MHz and a reception frequency range of 13-24 MHz, transmit power of 10%, mechanical index at 0.2, and dynamic range of 40 dB. The destruction-replenishment method was used for quantification of the targeted enhancement between pre-burst signal from free-circulating and bound MBs, and replenishment signal from only free-circulating MBs acquired immediately after a destructive pulse (Figure 1C) as previously described [21]. Between acquisitions with different types of MBs in the same mouse, a lag time of a minimum of 25 min was included for MB wash-out, which was verified by CEUS [44]. The order of injections with different MB types was randomized. B-mode was used for anatomical guidance to image the same position before and after blocking with NET1-H-mAb. USMI acquisitions with MB_NETrin-1 and MB_isotype prior to blocking were performed on day 1 of the imaging protocol. USMI acquisitions post blocking were performed on day 2; i.e. 24 h post injection of NET1-H-mAb for blocking.

**Imaging data analysis:** The imaging sequence was composed of 166 pre-burst frames (=12 s), a burst, and 130 replenishment frames (=10 s). To burst the MBs, a continuous high-power destructive pulse of 3.7 MPa (100% transmit power, mechanical index of 0.63, duration of 1 s) was applied. The pre-burst and replenishment frames were used to determine bound MB signal: short sequences between two respirations were manually selected; the pre-burst frame was selected right before applying the burst and the post-burst frame was selected a few seconds after burst when the circulating MBs reached the imaging frame, which was indicated by signal saturation. To avoid over- or underestimation of imaging signal in absolute values due to inter-tissue heterogeneity, the molecular imaging signals were calculated as the targeted enhancement ratio (T.E. (r)) in linear units (l.a.u.) using the VevoLab software.
T.E. (r) = (pre-burst signal in l.a.u.) / (post-burst signal in l.a.u.).

To avoid the presentation of data with artificially high background signal, the imaging data was then converted into percentages according to the following formula:

\[
\text{Molecular imaging signal in } \% = \left(1 - \left(\frac{1}{T.E. (r)}\right)\right) \times 100%.
\]

For visual presentation, image panels were provided that present tumors on anatomical B-mode images in grey (upper row in figure panels) and the respective contrast-enhanced ultrasound image in brown (lower row in figure panels) including the color-coded USMI signal. These images were exported from VevoCQ software, which allows visualization of the USMI signal distribution in the region of interest (green contour) as the differential targeted enhancement (d.T.E. in l.a.u.):

\[
d.T.E. = (\text{pre-burst signal}) - (\text{post-burst signal}).
\]

For comparison of molecular imaging signal between different tumor models, the differential USMI signal was calculated by subtracting the background signal (quantified with MBisotype) from the targeted molecular imaging signal (quantified with MBNetrin-1): \(\Delta(MB_{\text{Netrin-1}} - MB_{\text{isotype}})\) USMI signal (%). This was calculated for every tumor. Then, the individual tumor values were used to calculate the mean \(\Delta(MB_{\text{Netrin-1}} - MB_{\text{isotype}})\) USMI signal and SEM for each tumor model, and the statistical analysis was performed.

**Netrin-1 expression analysis by q-RT-PCR**

Total RNA was extracted from biopsies using the TRizol-Reagent (Ambion, Life Technologies) and 1 µg was reverse-transcribed using the iScript cDNA Synthesis kit (BioRad). Real-time quantitative RT-PCR was performed with a LightCycler 2.0 apparatus (Roche) using LightCycler® TaqMan® Master kit (Roche, Basel, Switzerland). Reaction conditions for all optimal amplifications, as well as primers selection were determined as already described [6]. The ubiquitously expressed human or murine hypoxanthine phosphoribosyl transferase (HPRT) gene was used as an internal loading control. The following primers were used: murine netrin-1 forward - 5’-GCAAGCTGAAGATGAACATGA-3’, reverse - 5’-CTTTGTCGGCCTTCAGGAT-3’; murine HPRT forward - 5’-TTCTTCTAGACCGTTTTT-3’, reverse - 5’-CCTGGTTCATCATCGCTAATC-3’; human netrin-1 forward - 5’-TGACCTTGATTTATTTTGCAATCC-3’, reverse - 5’-CGAGCAAGACGTTCAGTCTCCT-3’.

**Statistical analysis**

The number of samples or replicates per experimental condition or group is indicated as N. Standard errors (STD) or standard errors of the mean (SEM) are shown in graphs as indicated in the figure legends. Differences between two groups were tested by Student’s t-test. Comparisons of more than two groups were performed using ANOVA followed by a Tukey-Kramer post-hoc test. Statistical p-values are indicated in the graphs. The absence of statistically significant p-values indicated that differences were not significant. Statistical analyses are specifically indicated in every figure legend. Details about the receiver operating characteristic (ROC) curve are described in Supplementary Material.

**Results**

**In vivo immuno-localization of netrin-1**

First, we assessed whether netrin-1 was localized on vascular endothelial cells of netrin-1-positive tumors and was an accessible molecular target for USMI. IVIL and immunofluorescence showed that the anti-netrin-1 antibody signal was present in epithelial and endothelial layers of human SKBR7 tumors (Figure 2). Comparison of fluorescence intensities of NET1-H-mAb and isotype control that were co-localized with CD31 showed a significantly greater extent of immunofluorescence with NET1-H-mAb in netrin-1-positive SKBR7 (NET1-H-mAb: fluorescence intensity of 51.54 ± 4.2 a.u.; isotype control: 38.6 ± 3.4 a.u.; p=0.035), while there was no difference between NET1-H-mAb and isotype control in weakly netrin-1-expressing human MDA-MB-231 (NET1-H-mAb: fluorescence intensity of 28.29 ± 2.5 a.u.; isotype control: 34.17 ± 4.3 a.u.; n.s.) (Figure 2). Q-PCR confirmed netrin-1 expression in SKBR7 tumor tissues while netrin-1 was barely detectable in MDA-MB-231 tumors (Figure S1). Thus, SKBR7 breast tumors with epithelial netrin-1 expression showed NET1-H-mAb staining in the tumor endothelium, suggesting an association between netrin-1 expression in tumor cells and endothelial presentation of netrin-1.

**Netrin-1 on cell surfaces**

Flow cytometry analysis confirmed netrin-1 presentation on the cell surface of HeLa-Net1 model cells while netrin-1 was not detected by Western blot and flow cytometry analysis of HeLa-Ctrl cells (Table 1, Figure S2 and Figure S3A). Incubation of HeLa-Net1 cells with heparin reduced netrin-1 presentation by 69.5% (Table 1 and Figure S3B). The use of an enzyme cocktail degrading heparan sulfates
resulted in depletion of heparan sulfate by 95.4% and netrin-1 by 55.9% in HeLa-Net1 cells (Table 1 and Figure S3C). Incubation of HeLa-Ctrl cells with exogenous netrin-1 resulted in strong presentation shown by a 49-fold rise in signal intensity (Table 1 and Figure S3D). The enzymatic digestion of heparan sulfate and netrin-1 addition to HeLa-Ctrl cells induced a decrease of heparan sulfate by 76.2%, and of netrin-1 by 37.4% (Table 1 and Figure S3D). HeLa-Net1 cells revealed a lower heparan sulfate level than HeLa-Ctrl cells (50.9% of HeLa-Ctrl), but the heparan sulfate signal of HeLa-Ctrl cells dropped once netrin-1 was added (Table 1 and Figure S3C-D). HUVEC endothelial cells did not naturally present netrin-1, but addition of exogenous netrin-1 resulted in presentation with a 2.5-fold higher signal compared with untreated HUVEC cells (Table 1 and Figure S3E). On HUVEC cells, heparin incubation again prevented netrin-1 presentation (Table 1). To summarize, netrin-1 was detected on cell surfaces of cancer and endothelial cells using the anti-netrin-1 antibody and netrin-1 presentation was at least partially mediated by heparan sulfate proteoglycans.

Table 1. Netrin-1 interaction on cell surfaces studied in HeLa-Ctrl and HeLa-Net1 model cells, and HUVEC endothelial cells studied by flow cytometry. Mean fluorescence intensities were converted into signal ratios by division of marker intensity by the respective isotype control. Cell surface presentation of netrin-1, the effect of heparin addition (12 h), enzymatic digestion of cell surface glycans (3 d), addition of exogenous netrin-1 (1 h), a combination of enzymatic digestion of glycans and subsequent netrin-1 addition, and a combination of heparin pre-treatment and addition of exogenous netrin-1 were tested. For enzymatic treatment, a cocktail containing Heparinase I, Heparinase III, Chondroitinase ABC and Glucorinidase was used. Netrin-1 was produced by HeLa-Net1 cells. N=1.

| Netrin-1-positive and -negative model cell lines |  |
|---|---|
| Cell line | Marker | Signal ratio |
| HeLa-Ctrl | Netrin-1 | 0.9 |
| HeLa-Net1 | Netrin-1 | 2.6 |
| **Heparin** |  |
| Cell line | Marker | Signal ratio |
| HeLa-Net1 | Netrin-1 | Heparin |
| Heparin sulfate | 135.8 | 2.5 |
| 97.2 |
| **Enzymatic degradation of cell surface glycans** |  |
| Cell line | Marker | Signal ratio |
| HeLa-Net1 | Netrin-1 | Heparin sulfate |
| 11.8 | 5.2 |
| 28.1 | 1.3 |
| **Addition of Netrin-1 and enzymatic digestion of cell surface glycans** |  |
| Cell line | Marker | Signal ratio |
| HeLa-Ctrl | Netrin-1 | Heparin sulfate |
| 0.9 | 44.1 | 27.6 |
| 55.2 | 34.8 | 8.3 |
| **Netrin-1 presentation on HUVEC endothelial cells** |  |
| Cell line | Marker | Signal ratio |
| HUVEC | Netrin-1 | Heparin & Netrin-1 |
| 1.1 | 2.7 | 1.4 |
Figure 3. MB binding in vitro on purified netrin-1 protein and on netrin-1-overexpressing cells. (A) Anti-netrin-1-MBs binding specificity showed significant differences between MB concentration at 4 µg/mL and all other conditions (one-way ANOVA (F(5,12) = 12.55, p=0.0002 followed by Tukey-Kramer post-hoc test). N=3. (B) Anti-netrin-1-MB binding in presence of human plasma. Binding ratios (numbers of MBs binding to BSA vs. number of MBs binding to netrin-1 protein) are depicted in dashed grey boxes. N=3. (C) Anti-netrin-1-MB binding at body temperature. N=3. (D) Anti-netrin-1-MB binding under shear stress. N=3. (E) Anti-netrin-1-MB binding on HeLa-Ctrl and HeLa-Net1 cells under static conditions in presence of plasma. N=3. (F) Anti-netrin-1-MB binding on human strongly netrin-1-expressing SKBR7 and weakly netrin-1-expressing MDA-MB-231 breast cancer cells under shear stress (1 dyne/cm²). N=2. Except for (A), two-group comparisons were performed with Student’s t-test. Error bars present SEM; scale bars indicate 10 µm.

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Preparation of netrin-1-targeted MBs and validation of MB binding in vitro

Next, a netrin-1-targeted UCA was created for USMI (Figure S4 and Figure S5A-D) and validated in in vitro binding assays. Netrin-1-targeted MBs bound to netrin-1 in a dose-dependent manner (Figure 3A). Netrin-1 blocking with NET1-H-mAb prior to incubation with anti-netrin-1-MBs significantly reduced binding (Figure 3A). The use of plasma buffer and a rise in temperature to 37 °C improved the ratio of MB binding to BSA vs. binding to netrin-1 protein (Figure 3B-C). Under dynamic conditions in flow chamber assays, MBs bound to netrin-1 at a shear stress of 2 dynes/cm² and essentially did not bind to BSA (Figure 3D). MB binding was significantly higher on HeLa-Net1 cells than on HeLa-Ctrl cells in static binding assays (Figure 3E). In agreement with q-RT-PCR analysis of netrin-1 expression (Figure S6), MB binding in flow chamber assays was significantly higher on netrin-1-positive human SKBR7 cells than on barely netrin-1-expressing human MDA-MB-231 cells (Figure 3F) [5]. In short, the UCA was able to bind to recombinant netrin-1 and to tumor cells expressing high levels of netrin-1.

In vivo imaging of netrin-1 in engrafted breast tumors of human and murine origin

To validate the use of MBs in vivo, breast tumors were imaged with NET1-H-mAb-functionalized MBs (MBNetrin-1) and human isotype control MBs (MBIsotype). Netrin-1-positive human SKBR7 breast tumors showed a significantly higher molecular imaging signal with MBNetrin-1 compared with MBIsotype (26.4% ± 3.3% vs. 17.8% ± 3.4% vs. 3.8% ± 4.8% signal intensity; \( p=0.044 \)) (Figure 5A). After blocking with NET1-H-mAb, MBNetrin-1 and MBIsotype showed similar intensities: MBNetrin-1 at 33.7% ± 3.4% and MBIsotype at 26.4% ± 3.3%. In normal glands, there was no significant difference between signals of both MB types (Figure 5B). USMI signal detected with MBIsotype in breast tumors was significantly higher than the signal observed in normal mammary glands (32.0% ± 2.9% for breast cancer vs. 9.0% ± 2.2% for normal glands; \( p=2.72 \times 10^{-8} \)) indicating heterogeneity in unspecific background signal (Figure 5A-B). The differential USMI signal, which normalizes the molecular imaging signal to the non-specific background signal, showed a 2.7-fold higher imaging signal in MMTV-PyMT tumors compared with normal mammary glands (14.7% ± 3.1% vs. 5.5% ± 3.9% signal intensity; \( p=0.0003 \)) suggesting that MMTV-PyMT tumors were netrin-1-positive (Figure 5C).

Q-RT-PCR confirmed netrin-1 overexpression in MMTV-PyMT breast tumors (26.8 ± 16.6-fold expression relative to HPRT in tumors in contrast to a 1.8 ± 0.4-fold expression in normal breast glands; \( p=0.024 \)) (Figure 6A). IVIL and immunofluorescence analysis showed specific labeling of the CD31-positive MMTV-PyMT breast tumor endothelium with anti-netrin-1 antibody compared to isotype control
antibody (fluorescence intensity of 50.3 ± 3.5 a.u. vs. 37.6 ± 1.8 a.u.; \( p = 4.8 \times 10^{-3} \)) (Figure 6B). In normal breast glands, NET1-H-mAb and isotype control antibody showed no significant differences in fluorescence intensity (39.7 ± 3.5 a.u. vs. 41.3 ± 4.8 a.u., respectively) (Figure 6B). Endothelial localization of netrin-1 in all netrin-1-positive tumor models was further confirmed by co-localization with CD146, an endothelial receptor with affinity for netrin-1 (Figure S8) [11]. Taken together, in vivo imaging results for MMTV-PyMT and wild type mice were confirmed by ex vivo analysis of netrin-1 presentation.

Figure 4. Assessment of USMI for netrin-1 detection using anti-netrin-1-MBs (MBNetrin-1) and isotype control MBs (MBisotype). (A) Netrin-1-positive human subcutaneous SKBR7 breast tumors in nude mice were imaged with both MB types prior to and post blocking with NET1-H-mAb (N=9). (B) Weakly netrin-1-expressing human subcutaneous MDA-MB-231 breast tumors were imaged with both MB types (N=9). Image panels present ultrasound B-mode images in grey (upper row) and the respective contrast-enhanced ultrasound image in brown (lower row) including the color-coded USMI signal distribution in the region of interest (green contour). Scale bars indicate 1 mm; error bars indicate SEM. (C) For comparison of molecular imaging signal between SKBR7 tumors and MDA-MB-231 tumors, the differential USMI signal was calculated by subtracting the background signal (quantified with MBisotype) from the targeted molecular imaging signal (quantified with MBNetrin-1): \( \Delta \) (MBNetrin-1-MBisotype) USMI signal (%). Results show a significantly enhanced signal in SKBR7 tumors compared to MDA-MB-231 tumors. SKBR7 (N=9), MDA-MB-231 (N=9); Error bars indicate SEM. In all figures, two-group comparisons were performed with Student’s t-test.
Figure 5. Feasibility of USMI of netrin-1 in the murine transgenic MMTV-PyMT breast cancer model. (A) USMI signal intensities obtained with anti-netrin-1-MBs (MB\textsubscript{Netrin-1}) and isotype control MBs (MB\textsubscript{Isotype}) prior to and post blocking of netrin-1 with NET1-H-mAb in MMTV breast cancer (N=30). (B) As a control, normal mammary glands of wild type mice were imaged with both types of MBs (N=15). Image panel presents ultrasound B-mode images in grey (upper row) and the respective contrast-enhanced ultrasound image in brown (lower row) including the color-coded USMI dTE signal distribution ((pre-burst) – (post-burst)) in the region of interest (green contour). Scale bars indicate 1 mm; error bars indicate SEM. (C) For comparison of molecular imaging signal between transgenic murine tumors and normal mammary glands, the differential USMI signal (Δ(MB\textsubscript{Netrin-1} - MB\textsubscript{Isotype}) USMI signal (%)) was calculated and showed a significantly enhanced signal in MMTV-PyMT tumors compared to normal glands. MMTV-PyMT breast cancer (N=30), normal mammary glands (N=15); error bars indicate SEM. In all figures, two-group comparisons were performed with Student’s t-test.
Figure 6. Ex vivo analysis of netrin-1 expression in the MMTV-PyMT breast cancer model. Q-RT-PCR analyses (A) and in vivo immuno-localization (IVIL) (B) of netrin-1 expression confirm the strong expression of netrin-1 in MMTV-PyMT tumors and significantly weaker expression of netrin-1 in normal mammary glands. N=13 tumors (of two mice) per group of MMTV-PyMT and N=7 mammary glands (of one mouse) per group of normal glands in IVIL; N=4 for q-RT-PCR; error bars present SEM; IVIL scale bars indicate 20 µm. In all figures, two-group comparisons were performed with Student’s t-test. Finally, a receiver operating characteristic (ROC) curve was estimated based on the MMTV-PyMT tumor vs. normal gland USMI signal in % derived from netrin-1-targeted MBs only (and not from the difference between netrin-1-targeted and isotype control MBs). The datasets contained 45 imaging samples thus enabling a statistically robust estimation of the curve. The area under the curve (AUC) and confidence interval (CI) showed that USMI based on MB$_{Netrin-1}$ signal alone allowed differentiating between normal mammary glands and MMTV-PyMT breast tumors (AUC: 0.98; 95% CI: 0.95-1) with high diagnostic accuracy (Figure S9). At a threshold level as low as 33.3%, normal mammary glands of wild type mice and breast tumors in MMTV-PyMT mice were predicted to be diagnosed with a sensitivity of 90% (95% CI: 80–100) and a specificity of 100% (95% CI: 98–100). These observations confirm the potential diagnostic value of USMI for the detection of netrin-1 in tumors.
Discussion

**In vivo immuno-localization of netrin-1**

Netrin-1 expression by endothelial model cells remains controversial [8,45]. To understand whether netrin-1 is presented on the endothelium of netrin-1-overexpressing tumors, we applied IVIL and showed netrin-1 expression in the tumor epithelium and co-localization with endothelial CD31 and CD146 in human SKBR7 breast tumors. However, the results did not allow conclusions on the origin of netrin-1—whether it was secreted by epithelial tumor cells and diffused to the endothelium or whether the tumor induced netrin-1 expression in the endothelium. Alternatively, a third explanation for co-localization of netrin-1 and CD31 could be a structurally impaired endothelial lining interspersed with netrin-1-overexpressing tumor cells [46].

**Netrin-1 on cell surfaces**

Netrin-1 is a secreted ligand containing a heparin-binding domain, and heparan sulfate proteoglycans act as flow sensors on the luminal side of endothelial cells [1,47]. Thus, heparan sulfate proteoglycan-mediated netrin-1 presentation on cell surfaces was studied. The data confirmed removal of netrin-1 from the cell surface after heparin addition or depletion of heparan sulfate proteoglycans. Further studies are required to understand whether other netrin-1-binding molecules mediate netrin-1 presentation on cell surfaces [45]. The incubation of cells with netrin-1 protein showed that netrin-1 binding to the cell surface was independent of the origin of netrin-1, i.e., whether it was an autocrine or exocrine production. Netrin-1 binding on cell surfaces interfered with the detection of heparan sulfate, suggesting direct or indirect netrin-1 binding to the anti-heparan sulfate antibody epitope. Our results suggest netrin-1 secretion, diffusion, and immobilization on cell surfaces of epithelial and endothelial cells. Taken together, the endothelial localization of netrin-1 as shown in the in vivo immuno-localization study, and the reliable detection of cell surface netrin-1 with NET1-H-mAb on endothelial cells, provided our rationale for USMI with netrin-1-targeted UCAs.

**In vivo imaging of netrin-1**

The in vivo USMI study indicated the feasibility of netrin-1 imaging in murine blood vessels supplying human SKBR7 breast tumors in nude mice, and corroborated in vitro results of MB binding on breast cancer cell lines. The data confirmed the hypothesis of a correlation between epithelial and endothelial netrin-1 presentation in tumors. Our in vitro and in vivo data show that netrin-1, although described as a secreted and diffusible protein, bound to the cell surface and/or extracellular matrix. This observation is in line with previous reports about USMI of secreted frizzled related protein 2, interleukin 16, and GPIIb/IIIa in tumors, and confirms the feasibility of USMI in settings where free-circulating proteins can potentially saturate MBs and prevent immobilization [22,48,49]. Inter- and intra-tumoral heterogeneity with regard to vascular perfusion rates and functionality of the vascular network are crucial factors for USMI with micrometer-sized contrast agents that are injected and limited to the blood vessels [50,51]. Thus, different tumors might require differently timed injections and imaging protocols to ensure that MBs reach the tumor before and after the destructive pulse, a concept that cannot be translated into practice. Alternatively, one might individually assess tumor heterogeneity and use netrin-1-targeted and isotype control contrast agents to determine background signals and assess the amount of molecular imaging signal in every tumor model. Comparison between MB_{Netrin-1} and MB_{isotype} enabled the reliable detection of netrin-1-positive subcutaneous tumors and confirmed IVIL results. The transgenic MMTV-PyMT mouse model reflects naturally occurring tumor development and neoangiogenesis, critical in USMI, and is more physiologically accurate to human disease than subcutaneously engrafted tumor models, providing clinically relevant insight into imaging of netrin-1 [52]. USMI with both MB_{Netrin-1} and MB_{isotype} enables the reliable detection of netrin-1 expression in MMTV-PyMT breast tumors, and netrin-1 absence in normal mammary glands as confirmed by IVIL analysis. Furthermore, the use of MB_{isotype} revealed that the intensity of background signal differs strongly between the SKBR7 and MDA-MB-231 models so that their netrin-1 molecular imaging signals cannot be directly compared with each other and require the subtraction of background signal. A future challenge will be to reduce baseline signal and amplify signal from bound MBs to facilitate the distinction between netrin-1-positive and negative tumors using netrin-1-targeted MBs only. USMI signal can be normalized by vascularity, which is thought to have an impact on background signal, after estimation of relative blood volume, vessel density, and vessel perfusion [53,54]. The use of three-dimensional USMI might enable more complete assessment of tumor heterogeneity compared to the two-dimensional imaging data of this study, but further technological development is required for a routine use in preclinical or clinical settings [27].

In addition to the characterization of tumor heterogeneity, contrast agents and imaging strategies
can be further developed to overcome tissue-dependent signal variances and produce imaging signals that are solely dependent on the degree of netrin-1 expression: (1) Contrast agents can be prepared by directly integrating/conjugating the targeting moiety to the MB shell, eliminating the need for an immunogenic avidin-biotin bridge. Instead of the whole antibody, which might cause Fc-mediated inadvertent non-specific interactions [55], an antibody Fab fragment or novel targeting peptide has to be prepared to provide a clinically translatable targeted contrast agent that reduces background signal in the tissue. Our blocking studies using NET1-H-mAb IgG, which was expected to bind to netrin-1 and Fc receptors, did not reveal a change in MBI isotype imaging signal. Therefore, it was assumed that Fc-mediated binding of NET1-H-mAb did not cause background signal. (2) In this study, MBs were used at a high concentration compared to other studies to increase the sensitivity of our imaging approach for the secreted ligand netrin-1, which was described to be membrane-associated but also free-circulating in the blood. Having shown the feasibility of USMI of netrin-1, a novel study is conceivable in which lower concentrations of MBs can be employed to reduce the level of background signal and facilitate the comparison of heterogeneous tumor models [56]. (3) The imaging protocol based on the destruction-replenishment method has become the standard technique in pre-clinical studies of ultrasound molecular imaging [24]. Though, MB destruction causes cavitation, which potentially harms the blood vessels and affects the comparison of pre- and post-destructive acquisitions [57]. Alternative strategies have been proposed, such as late-phase signal enhancement, which is used in clinical studies [30], and in which images are acquired 30 min after microbubble injection when free-circulating microbubbles are degraded, or techniques based on the microbubble dwell time [58], but they require optimization to allow for fast acquisitions before they can be routinely used.

The diagnostic potential of USMI of netrin-1 was further assessed with ROC curves, which were analyzed with the MBI isotype signals of the transgenic MMTV-PyMT tumors and normal mammary glands. ROC curves indicated that USMI of netrin-1 allowed distinction between normal mammary tissue and transgenic tumor endothelium, thus underlining the diagnostic power of the technique.

**Conclusion**

To the best of our knowledge, this is the first study proposing USMI as a companion diagnostic for a molecularly targeted therapy, thereby promoting novel targets and applications for USMI. Compared to clinical diagnostics such as immunohistochemistry, USMI is non-invasive, cost efficient, rapid, and can be performed at the point-of-care. In conclusion, after confirmation of netrin-1 presentation on the vascular endothelium of netrin-1-expressing tumors, USMI was successfully developed as a companion diagnostic for this secreted tumor marker. Technical improvements and further pre-clinical therapy response monitoring studies will boost this powerful imaging approach for personalized medicine. The development of USMI for relevant targets can significantly improve the quality of patient management.

**Abbreviations**

AUC: area under curve; CI: confidence interval; IVIL: *in vivo* immuno-localization; MB: microbubble; MFI: mean fluorescence intensity; MMTV-PyMT: mouse mammary tumor virus polyoma virus middle T antigen; ROC: receiver operating characteristic; SEM: standard error of mean; STD: standard deviation; UCA: ultrasound contrast agent; USMI: ultrasound molecular imaging.

**Supplementary Material**

Supplementary figures and tables. http://www.thno.org/v08p5126s1.pdf

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Author contributions

J.W. planned the study, conducted the experiments, analyzed the data, the prepared the figures, and wrote the paper. K.E.W. assisted with experiments and data analysis, and edited the manuscript. J.-G.D. planned experiments, prepared reagents, conducted experiments, analyzed data, and assisted with manuscript writing. R.M.-P. performed experiments. B.G. assisted with experiments and manuscript writing. S.J., J.N., and D.G. performed experiments. P.M. provided funding, and edited the manuscript. J.K.W. helped with planning of experiments and interpretation of data, edited the manuscript, and provided funding. F.P. helped with planning of experiments and interpretation of data, provided funding, performed statistical analyses, and assisted with manuscript writing.

Competing Interests

D. Goldschneider is an employee of Netris Pharma. J.G. Delcros is the inventor of the anti-netrin-1 antibody (patent EP289399A1). J.G. Delcros and B. Gbert are consultants for Netris Pharma. P. Mehlen is a shareholder of Netris Pharma. The authors J. Wischhusen, K. E. Wilson, R. Molina-Peña, S. Jiang, J. Ngo, and J. K. Willmann, and F. Padilla declare no potential conflicts of interest.

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