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Abstract. Molecular-targeted probes are emerging with applications for optical biopsy of cancer. An underexplored potential clinical use of these probes is to monitor residual cancer micrometastases that escape cytoreductive surgery and chemotherapy. Here, we show that leukocytes, or white blood cells, residing in nontumor tissues—as well as those infiltrating micrometastatic lesions—uptake cancer cell-targeted, activatable immunoconjugates nonspecifically, which limits the accuracy and resolution of micrometastasis recognition using these probes. Receiver operating characteristic analysis of freshly excised tissues from a mouse model of peritoneal carcinomatosis suggests that dual-color imaging, adding an immunostain for leukocytes, offers promise for enabling accurate recognition of single cancer cells. Our results indicate that leukocyte identification improves micrometastasis recognition sensitivity and specificity from 92 to 93%—for multicellular metastases >20 to 30 μm in size—to 98 to 99.9% for resolving metastases as small as a single cell. © 2014 Society of Photo-Optical Instrumentation Engineers (SPIE) [DOI: 10.1117/1.JBO.19.6.066006]

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Microscopic deposits of cancer cells missed during cytoreductive surgery frequently contain a subpopulation of cells with either intrinsic or acquired drug-resistance that escape follow-up chemotherapy and stimulate disease recurrence; for example, this is a key characteristic of ovarian cancer.1-4 A major challenge in oncology is the lack of imaging modalities capable of detecting or monitoring these residual microscopic tumors.5-7 Inspired by elegant reports of microscopic tumor visualization using intravital microscopy to study basic cancer biology,8 in vivo histopathology, and molecular imaging using microendoscopy9-10 as well as activatable fluorescent11-21 and photodynamic probes,22,23 we previously reported quantitative, repeated, and longitudinal monitoring of cancer micrometastases in vivo within select regions of the peritoneal cavity using a cancer cell-targeted, activatable, and photocytotoxic immunoconjugate integrated with fluorescence microendoscopy.24 The activatable immunoconjugate25,26 targets cancer cells overexpressing the epidermal growth factor receptor (EGFR) and serves dual functions as an imaging probe and a combinational therapeutic agent.24,27 enabling selective destruction of disseminated cancer micrometastases upon wide-field, near-infrared irradiation—a new mode of photoimmunotherapy.28–30 termed tumor-targeted, activatable photoimmunotherapy.24

The activatable immunoconjugate (Cet-BPD) is composed of cetuximab, an anti-EGFR monoclonal antibody (mAb) that targets cancer cells overexpressing EGFR, loaded with benzoporphyrin derivative (BPD) molecules, a U.S. Food and Drug Administration-approved photocytotoxic chromophore that undergoes electronic excited state quenching upon antibody conjugation.25 In this approach, both the photodynamic and fluorescent components of the immunoconjugate become de-quenched (activated) upon cellular internalization and processing.24,25 Because cancer cells overexpressing the target surface molecules take up the immunoconjugates more efficiently, this targeted activation occurs predominantly within tumors and enhances tumour selectivity (based on extensive imaging and phototoxicology studies comparing immunoconjugates with low- and high-quenching efficiencies).24 These studies, and the present study, were performed in a clinically motivated mouse model of human ovarian carcinomatosis previously established in our laboratory with gynecologic oncologists.31 In this orthotopic xenograft model, athymic Swiss female Nu/Nu mice (20 to 25 g and six- to eight-weeks old; Cox Breeding Laboratories, Cambridge, Massachusetts) are injected intraperitoneally (i.p.) with 16 × 106 NIH:OVCAR5 (human ovarian cancer) cells in 2 mL phosphate-buffered saline, leading to disseminated tumor nodules studing the peritoneal organs and surfaces. All animal experiments were conducted according to Massachusetts General Hospital Institutional Animal Care and Use Committee guidelines. Conjugates of BPD and mAb (Cet) were prepared as described previously24,25 and administered at 2 mg · kg−1 body weight BPD (1.4 mg cetuximab). All injections were done i.p. in 1 mL of sterile phosphate-buffered saline. The cetuximab dose (1.4 mg) is equivalent to ~180 mg · m−2 and near the clinical cetuximab dose of 250 to 400 mg · m−2.

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To examine the microscopic localization of activated immunoconjugates, we performed histopathologic examination of tissue biopsies coregistered with in vivo fluorescence microendoscopy images, and we performed confocal fluorescence microscopy of freshly excised tissues from the peritoneal cavity. Our prior results indicated 90% sensitivity and 90% specificity for tumors as small as 30 μm (a small cluster of cells) using monochromatic fluorescence detection of the activatable immunoconjugate. However, both immunofluorescence microscopy and histopathology suggested that a subpopulation of CD45+ (leukocyte common antigen or protein tyrosine phosphatase, receptor type, C) leukocytes uptake and activate the immunoconjugate and that this source of nonspecific probe activation limits the resolution of accurate micrometastasis recognition and quantitative monitoring post-treatment.

In the present study, we hypothesized that CD45 imaging can supplement activatable immunoconjugate imaging to enhance the accuracy and resolution of micrometastasis recognition. CD45+ cells, including macrophages residing in tissue, express Fc-γ-receptors that react with the Fc portion of MAbs to promote engulfment and phagocytosis as a mechanism of probe catabolism and elimination from the body. Here, we show that dual color imaging, adding an immunostain for CD45+ leukocytes—which can uptake activatable probes nonspecifically—can increase both the sensitivity and specificity to 98% for recognition of single cancer cells.

To test this concept of immune cell imaging in combination with activatable probes, we collected a series of confocal fluorescence image mosaics of freshly excised tissues from the mouse model of metastatic ovarian cancer prior to [day 0, no-tumor control; Fig. 1(a)] and at various days following

Fig. 1 (a) Confocal fluorescence image mosaics (left) and insets (right) of freshly excised tissue from control no-tumor mice. Human ovarian cancer cells (orange; antihuman cytokeratin 8 immunostain) are not found in these control mice. The locations of mouse leukocytes (cyan; antimouse CD45 immunostain) are shown for comparison with benzoporphyrin derivative (BPD) fluorescence (red; activated Cet-BPD immunoconjugates administered in vivo). Circles indicate leukocytes with activated BPD fluorescence (brightened 4x for visibility). (b) Confocal fluorescence image mosaics (left) and insets (right) of freshly excised tissue from epithelial ovarian cancer mice with disseminated, peritoneal micrometastatic disease. Note that leukocytes are an important cell population within the tumor microenvironment, are known to infiltrate cancer metastases, and often compose a large fraction of the tumor mass. Arrows indicate a micrometastasis composed of ovarian cancer cells (orange) as well as tumor-infiltrating leukocytes (cyan) with activated BPD fluorescence (red). Scale bars are 1 mm in (a) and (b) (left) and 100 μm in (a) and (b) (right).
tumor inoculation (days 3, 9, and 13; Fig. 1(b)) following administration of the Cet-BPD activatable immunoconjugate. BPD fluorescence corresponding to cancer cells and leukocytes is apparent in the resulting confocal image mosaics (Fig. 1). We designed two microscopic-resolution receiver operating characteristic (ROC) analyses using an antihuman cyto-keratin 8 MAb (CK8; clone LP3K, R&D Systems, Minneapolis, Minnesota) specific for human epithelial cancer cells23 as a gold standard marker: (1) a pixel-by-pixel analysis28 and (2) an object-by-object analysis (Fig. 2), where pixels are grouped into fluorescent objects and weighted as the object area normalized by the typical area of a single cell (diameter = 20 μm).

Following i.p. injection of Cet-BPD immunoconjugates (8 h post injection), freshly excised peritoneal wall specimens were immediately blocked (Dako Protein Block, Carpinteria, California), followed by immunofluorescence staining (10 to 16 μg/mL for each MAb), rinsed with fresh phosphate-buffered saline, and mounted on a coverslip bottom dish (MaTek, Ashland, Massachusetts). MAb-Alexa Fluor (AF) dye conjugates for ex vivo immunofluorescence staining were prepared using MAb-labeling kits (Life Technologies, Grand Island, New York): anti-human CK8 and anti-mouse CD45 (clone 30-F11, R&D Systems) were conjugated to AF647 and AF488, respectively. Imaging was performed with an Olympus FV1000 confocal microscope with a 10× objective. Excitation of BPD, AF488, and AF647 was carried out using 405-, 488-, and 635-nm lasers, respectively. Lasers were scanned sequentially to reduce channel crosstalk. The laser, photomultiplier tube detector, and pinhole settings were kept constant for each fluorophore. ROC analyses were performed using a custom, batch-processing Matlab (Mathworks, Natick, Massachusetts) routine. The tumor classification tests used objective intensity thresholds set to reject 99.5% of the background signal from the CK8 channel. That is, unstained control tissues were used to characterize and set quantitative thresholds to reject autofluorescence background (set to reject 99.5% of the non-Gaussian background intensity distribution25). For object selection in the CD45 channel (AF488, which overlaps strongly with autofluorescence), we developed a wavelet multiresolution decomposition (essentially band pass spatial frequency filtering) algorithm to select CD45+ cellular objects using the “wavedec2” and “wrcoef” functions with biorthogonal wavelets (bior3.7; Matlab Wavelet Toolbox). This routine calculates the absolute pixel values of image spatial frequencies band passed into detail level 4 and applies an intensity threshold (threshold level = 100) to select cellular objects.

In the first ROC analysis to assess the cancer cell-selectivity of Cet-BPD activation, we performed pixel-by-pixel tumor classification based on a Cet-BPD fluorescence intensity threshold from which we determined Cet-BPD sensitivity (fraction of cancer cell pixels correctly identified) and specificity (1—fraction of cancer cell pixels incorrectly identified). The optimal BPD intensity threshold attained 92 to 93% sensitivity and specificity (ROC area under the curve = 0.96) for micrometastases as small as 20 to 30 μm [Figs. 3(a) and 3(b)], as reported previously.24 For this analysis, we applied a balanced 21 μm dilate, fill and erode binary image operations using the “bwconvhull” and “imfill” functions (Matlab) to connect intercellular spaces of multicellular tumor objects. Size filtering of fluorescent objects in each channel was performed using the “bwareaopn” function. In addition, we calculated a penalty for size filtering potential BPD+ objects (above the 99.5% background rejection intensity threshold) below the 20 to 30 μm limit. The penalty is calculated as the percentage of BPD+ objects or pixels removed by the size filter, indicating that a substantial number of sub-20-μm objects are missed by this analysis [Fig. 3(c)]. Thus, size-filtering objects below the 20 to 30 μm limit provides an optimized algorithm for analyzing monochromatic fluorescence imaging of multicellular micrometastatic nodules, but there is great potential for dual-color fluorescence imaging—combining cancer cell-targeted activatable immunoconjugate with anti-CD45 MAb imaging—to further increase the accuracy and resolution of the smallest (i.e., single cell) micrometastases.

To explore the potential of dual-color imaging, we next added a third classification (false positives due to leukocytes residing in nontumor tissue, or cancer cell pixels incorrectly identified as tumor corresponding to CD45+ false positive pixels). The CD45+ false positive classification has no effect on the pixel-by-pixel ROC analysis due to the relatively small number of CD45+ nontumor pixels, which suggests that an object-by-object analysis is more appropriate. As described above, we weighted objects to their cellular mass (units of number of pixels correctly identified as tumor or correctly identified as nontumor) with false positive and false negative objects.

A noteworthy complication for object-by-object ROC analysis (with object assignment among three or more classes) is the potential to fragment objects during multiplicative binary operations.
masking operations. That is, binary masks calculated by intensity thresholding the raw image channels are normally multi-
plied to select colocalized objects and assign them to specific classes. However, this approach can split objects in complex images involving multiple channels and classifications. To prevent object fragmentation and to avoid splitting objects among different classifications, we used additive masking operations. This was accomplished using the “bwconncomp,” “regionprops,” “labelmatrix,” “find,” and “ismember” functions (Matlab) to assign contiguous BPD+ objects (following a 13 micron, balanced dilate, fill and erode binary image operation) to tumor, nontumor leukocytes, or nontumor (and nonleukocyte) classifications. Binary mask addition sets pixels within overlapping objects to 2, such that the mean intensity of the object is > 1.

Objects with mean intensity > 1 can then be found (using the “find” and “ismember” functions) and then the entire object can be assigned to the corresponding class (e.g., true positive). Similarly, objects with a mean intensity equal to 1 can be assigned to the opposing classification (e.g., false positive) without object fragmentation into multiple classifications. This strategy is less sensitive to chromatic aberrations and other sources of pixel color shift, as well as decreasing sensitivity to out-of-focus shadowing effects surrounding multilayer tumor objects.

The free parameters for these analyses (the wavelet threshold and the dilate-fill-erode distance) were tuned manually on a pair of test images (one nontumor control and one tumor biopsy), and then the entire set of image mosaics (eight tissue biopsies) were batch processed.

The results of the weighted object-by-object ROC analysis were calculated with (+CD45) and without (−CD45) the leuko-
cyte channel using the optimal BPD intensity threshold identi-
cified by the pixel-by-pixel ROC analysis (optimal BPD intensity threshold = 69 on a 12-bit scale). Dual-color Cet-BPD and CD45 imaging had a marked impact on the false positive rate, improving the specificity of micrometastasis recognition [Figs. 2(d) and 2(e)]. The dual-color, object-by-object ROC analysis indicates 99.9% sensitivity and 98.3% specificity for micrometastases as small as a single cell [no size filtering, 0% penalty; Figs. 3(d) to 3(f)]. Note that almost all of the unweighted objects are < 20 μm in size; therefore, the penalty approaches 100% [Fig. 3(c)]. In contrast, weighting the objects by the number of cells they contain decreases the penalty for size filtering small objects. However, size filtering is not needed and the penalty is zero when CD45 imaging is implemented [Figs. 3(d) to 3(f)].

In summary, molecular-targeted probes are emerging for performing minimally invasive in vivo microscopy with applications in oncology. An underexplored potential clinical use of these probes, in conjunction with fluorescence microendoscopy, is minimally invasive, repeated and quantitative monitor-
ing of cancer micrometastases in select sites is known to frequently harbor residual disease. CD45 (a leukocyte surface antigen) is a useful marker for identifying immune cells that uptake antibodies nonspecifically (e.g., macrophages) and removing these cells from analysis of micrometastases using cancer cell-targeted, activatable immunoconjugates. Multicolor approaches offer promise for treating and monitoring drug-resistant micrometastases presently invisible to clinicians with a sensitivity and specificity of ~98% for untreated tumors as small as a single cancer cell. Such an approach will also be of particular importance for monitoring micrometastasis destruc-
tion, to distinguish residual, tumor-associated inflammatory cells from residual cancer cells.

We anticipate that the overall approach of leukocyte imaging will be translated for use in humans and will be impactful for reducing false positives by identifying the immune cell uptake. In addition, the ROC analysis assay developed here has potential for translation to study cancer recognition using targeted molecular probes in human tissues. However, in the xenograft mouse model of epithelial ovarian cancer (EOC), we leveraged use of an antihuman CK8 probe that specifically binds human, but not mouse, protein to realize a high-fidelity gold standard marker for the human epithelial cancer cells. (In contrast, cetuximab binds both mouse and human EGFR.) Developing such a gold standard in human tissues will require some thought and further development. Furthermore, although EGFR is highly expressed by some tumors, it is unlikely that a single molecular marker can
be used to address inter- and intrapatient tumor heterogeneity and cancer cell clonal diversity; thus, new creative approaches will be needed to address multiple potential tumor biomarkers in human patients. Moreover, study of additional tissue types beyond the peritoneal tissues examined here is also warranted.

Of equal importance to probe and image analysis design is the development of clinical microendoscopy technology capable of more comprehensive samplings of micrometastatic burden. Elegant reports of comprehensive volumetric imaging of entire luminal organs have been accomplished, for instance, by rotary pull-back of tethered capsule probes through the gastrointestinal tract. However, the microscopic field-of-view does impose practical limits on sampling the entire micrometastatic burden in complex anatomical sites (e.g., the peritoneal cavity). Fiber-scanning microendoscope probes as well as multisolution, foveated lens systems can expand the field-of-view to enable efficient scanning of cavities and organs comparable to conventional endoscopy but with ultrathin probes and microscopic resolution. Furthermore, a number of advanced microscopy techniques, including multiphoton excitation and super-resolution imaging, have potential to or have already improved the imaging depth and resolution of microendoscopy.

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