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Article

Blood Endocan as a Biomarker for Breast Cancer Recurrence

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Simple summary: Cancer recurrence is a life-threatening event for breast cancer patients as metastatic recurrence is, in particular, almost impossible to surgically be removed. Detecting or predicting recurrence early helps in the management of breast cancer. The purpose of this study is to examine if endocan, a protein found in the blood of some breast cancer patients, is capable of becoming a marker to detect breast cancer recurrence more sensitive than existing markers. We found that blood endocan reflected tumor burden in preclinical animal models and preliminary clinical observations with breast cancer patients. Despite a limited number of patients, endocan detected in the blood of recurrent breast cancer patients is more sensitive to breast cancer recurrence than existing markers. Measuring blood endocan could pick up the patients who are overlooked by the existing markers, leading to prolongation of the life of such patients.

Abstract: Blood-based biomarkers for early detection of recurrence have a significant clinical impact on the improved management of breast cancer patients, as cancer recurrence results in poor prognosis. This study aimed to investigate if the measurement of blood endocan in breast cancer patients with high *ESM1* expression could be an effective tool to detect postoperative recurrence compared with existing tumor markers. Blood was collected before and after the tumor resection from the mouse models of breast cancer, and endocan levels were measured while visualizing metastatic recurrence with noninvasive luminescence imaging. In clinical settings, blood was withdrawn from 16 breast cancer patients before and after the tumor resection, and the effect of lumpectomy on blood endocan level was evaluated. Additionally, the blood endocan from 20 patients diagnosed with postoperative recurrence was measured, and their positivity rate for endocan was compared with that for serum carcinoembryonic antigen (CEA) or cancer antigen 15-3 (CA15-3). Our preclinical and clinical experiments revealed that blood endocan levels reflected tumor burden. Furthermore, over 60% of patients suffering from postoperative recurrence who tested negative for CEA or CA15-3 were positive for endocan. Our results support the clinical significance of endocan in breast cancer patients for detecting breast cancer recurrence.

Keywords: endocan; recurrence; biomarker; animal models; clinical study

1. Introduction

Breast cancer is now the leading cause of cancer incidence in women worldwide, accounting for one in four cancer cases and one in six cancer deaths [1]. Although the overall prognosis for breast cancer patients is relatively good when detected early, relative to other cancers, it has a high recurrence rate after surgery [2]. Cancer recurrence affects the patient's prognosis and quality of life, posing a critical clinical problem in later years as they often resist chemotherapy. The potential for recurrence and metastasis of breast cancer varies by subtypes, proliferative potential, and dormancy [3,4]. Even in the same subtype, recurrence-free survival varies from patient to patient; some patients experience recurrence within one year, while others do not see any recurrence over five years. Therefore, it is beneficial to predict whether each breast cancer patient is prone to relapse individually so that frequency of follow-up tests would be personalized. For those who experience breast cancer recurrence, it is also helpful if recurrence can be diagnosed as early as possible by sensitive tests in a patient-friendly manner. Follow-up management after the initial treatment of breast cancer, such as adjuvant or neoadjuvant therapy, has generally involved physical examination and diagnostic imaging [5].

Simple and minimally invasive measurement of blood-based tumor markers is used for early detection of cancer and monitoring treatment response. Carcinoembryonic antigen (CEA) and cancer antigen 15-3 (CA15-3) are, in particular, often used as tumor markers for several cancers in clinical settings [6,7]. CEA is a glycoprotein involved in cell adhesion and known to be elevated in many cancers [8]. CA15-3 is known to be a glycan-containing protein antigen of the transmembrane glycoprotein MUC-1 and has been reported to be associated with breast cancer stage and survival [9]. These two markers have been validated in a number of studies to measure changes in patients' blood to evaluate their application for early detection of cancer recurrence and monitoring treatment response[10,11]; however, there are little data available about the utility of these markers in asymptomatic or early-stage cancers. Although the frequent measurements of tumor markers in the blood have not been strongly recommended by several guidelines [12,13], developing effective biomarkers to detect breast cancer recurrence is critical, since liquid biopsy is generally more sensitive and cost-effective than diagnostic imaging.

Recently, endocan has received increasing attention as one of the blood-based biomarkers to detect various cancers. Endocan, encoded by the *ESM1* gene, is a human endothelial cell-specific molecule soluble dermatan sulfate proteoglycan known to circulate in the bloodstream [14,15]. Endocan expression is reportedly associated with clinicopathological parameters and poor prognosis in several cancers, including gastric cancer [16], hepatocellular carcinoma [17], prostate cancer [18], bladder cancer [19], and breast cancer [20]. Our previous study demonstrated that an elevated expression of *ESM1* in MDA-MB-231 cells, a human triple-negative breast cancer (TNBC) cell line, increased cell proliferation and tumor formation, and that TNBC patients with high *ESM1* expression in primary breast cancer had significantly shorter relapse-free survival [21]. In addition, blood endocan levels in luminal-type breast cancer patients were associated with cancer staging [20]. Therefore, since *ESM1* negatively impacts breast cancer patients and endocan, the gene product of *ESM1*, can be detected from the peripheral blood samples, the association between the status of breast cancer and blood endocan levels is clinically worth pursuing.

To bridge the gap between endocan's preclinical and clinical value as a blood-based biomarker, we sought to measure and assess the blood endocan levels in murine models of breast cancer and breast cancer patients. To the best of our knowledge, this is the first study to investigate the extent to which blood endocan levels in mice and patients with breast cancer are affected by the surgical removal of primary tumors and postoperative recurrence.

2. Materials and Methods

2.1. Cell Culture

In this study, MDA-MB-231BR cell line, a metastatic variant of human-derived triple-negative breast cancer cell line MDA-MB-231 [22], was used. Our previous study reported that MDA-MB-

231BR cells overexpressed *ESM1* and secreted measurable amounts of endocan, which was detected from the supernatant of the conditioned medium as well as the blood of mouse models bearing MDA-MB-231BR tumor, which was not the case with MDA-MB-231 [21]. To our knowledge, MDA-MB-231BR cell line is the only human breast cancer cell line overexpressing *ESM1* thus far, which is the reason why we used MDA-MB-231BR in this study.

MDA-MB-231BR was a kind gift from Dr. Patricia Steeg (National Cancer Institute, Bethesda, MD, U.S.A.). MDA-MB-231BR was cultured in RPMI-1640 (FujiFilm Wako Pure Chemical Corp., Osaka, Japan) with 10% fetal bovine serum (Thermo Fisher Scientific Inc.) and maintained under aseptic conditions of 5% CO₂ at 37°C. Contamination with *Mycoplasma* or fungi was routinely checked and only uncontaminated cells were used. Cell line authentication was outsourced to BEX Co, Ltd. (Tokyo, Japan), and MDA-MB-231BR/*mVenus-Akaluc*, detailed in the next section, was confirmed to be the identical cell strain to MDA-MB-231.

2.2. Generation of MDA-MB-231BR/*mVenus-Akaluc*

Primer was designed from pcDNA3 *Venus-Akaluc* vector (RDB15781; Riken BioResource Research Center, Ibaraki, Japan), and Q5® High-Fidelity 2X Master Mix (New England Biolabs, MA, USA) was used to amplify the DNA. The *mVenus-Akaluc* primers used for PCR amplification were as follows: FW, 5'-TAG AGC TAG CGA ATT ATG GTG AGC AAG GGC GAG-3' (33 bp); and RV, 5'-ATT TAA ATT CGA ATT CCA TAG AGC CCA CCG CAT-3' (33 bp). PiggyBac™ Transposon Vector System (System Biosciences, CA, U.S.A.) was used to transfect *mVenus-Akaluc* DNA into MDA-MB-231BR cells stably. The PB-CMV-MCS-EF1-Puro cDNA Cloning and Expression Vector (System Biosciences) was linearized by *Eco*RI-HF restriction enzymes, and a purified PCR fragment of *mVenus-Akaluc* was incorporated into the PB-linearized vector using In-Fusion® HD Cloning Kit (Takara Bio Ltd., Shiga, Japan). The resultant PB-*mVenus-Akaluc* plasmid was cloned with Stellar™ Competent Cells (Takara Bio Ltd.), followed by the purification with GenElute™ Plasmid Miniprep Kit (Sigma-Aldrich), and the sequence was verified by a DNA sequencing service (Eurofins Genomics Inc., Tokyo, Japan). MDA-MB-231BR was transfected with the plasmid DNA using Lipofectamine® LTX Reagent (Thermo Fisher Scientific Inc.), and stable lines, designated as MDA-MB-231BR/*mVenus-Akaluc* hereafter, were selected by 1 µg/mL of puromycin (Thermo Fisher Scientific Inc.). The basic function of MDA-MB-231BR/*mVenus-Akaluc*, such as cell proliferation and the visibility by fluorescence and luminescence imaging, was confirmed by Cell Counting Kit-8 assay, fluorescence microscopy, and IVIS Lumina Series III (PerkinElmer, Waltham, MA, U.S.A.).

2.3. Animals

Female athymic *nu/nu* mice (Balb/c background, four weeks old, 17-20 g) and NOD.Cg-*Prkdc*^{scid} *Il2rg*^{tm1Wjl}/*SzJ* (NSG) mice (4 weeks old, 16-21 g) were purchased from CLEA Japan, Inc. (Tokyo, Japan) and The Jackson Laboratory Japan, Inc. (Kanagawa, Japan), respectively. The institutional animal care committee at Hoshi University approved the study protocol before the animal experiments.

2.4. Blood endocan Measurements and Luminescence Imaging of Mice Bearing Orthotopic Breast Cancer

One million MDA-MB-231BR/*mVenus-Akaluc* cells were orthotopically inoculated into the mammary fat pad of mice. The tumor volume was measured and calculated in a blinded manner using a caliper:

$$\text{Tumor volume (mm}^3\text{)} = (\text{length}) \times (\text{width})^2 \times \frac{\pi}{6}$$

Considering the susceptibility of mouse strain to metastatic recurrence, the primary tumor was resected from each mouse once the tumor volume exceeded 500 mm³ in nude mice and 200 mm³ in NSG mice.

Local or metastatic recurrence was visually detected by luminescence imaging with IVIS Lumina Series III after intraperitoneal injection of akaLumine *n*-hydrochloride (FujiFilm Wako Pure Chemical Corp.) at a dose of 1.8 nmol.

In order to quantitate endocan in the peripheral blood of nude and NSG mice, blood was collected from the tail vein at predetermined time points. Plasma was isolated by centrifugation at 4°C, 1,200 ×g for 10 min, and stored at -80°C until use.

2.5. Blood Collection from Breast Cancer Patients

Clinical studies on the measurement of endocan levels in the blood of breast cancer patients were approved by the Institutional Review Board from Showa University Hospital and Hoshi University, which preceded the participation of the patients in this study. Participation in the study was voluntary, and written informed consent was received from all subjects. Patients with diabetes, an infectious disease, or other types of cancer were excluded from this study because all of those conditions are known to affect blood endocan levels [23–29]. The patient's blood samples were sent to a certified clinical lab for the routine measurements of tumor markers, CEA and CA15-3. CEA concentration of over 5 ng/mL and CA15-3 concentration of over 31.5 U/mL indicated positive for serum CEA and CA15-3, respectively.

Blood was collected from breast cancer patients at the Breast Center of Showa University Hospital (Tokyo, Japan) between 2020 and 2021, and the patient population is summarized in Supplementary Tables S1 and S2. Plasma was isolated by centrifugation at 4°C, 1,200 ×g for 10 min, and stored at -80°C until use.

2.6. Enzyme-Linked Immunosorbent Assay (ELISA) for the Quantification of Endocan

Plasma samples were diluted twice with sample diluent, and endocan was quantitated using commercial ELISA kits (Lunginnov (Lille, France) for plasma from nude mice and CUSABIO (Wuhan, China) for plasma from NSG mice and humans) according to the manufacturer's instructions. The detection limit was set at 0.156 ng/mL in accordance with the protocol of the ELISA kit. Based on our previous study, an endocan level of 1.68 ng/mL or higher was considered positive for endocan [20]. Endocan expression of breast cancer cells was confirmed negative by immunohistochemistry in the patients who showed blood endocan levels below the cutoff value (Supplementary Fig. 1) and, thus, was excluded from postsurgical endocan measurements.

2.7. Statistical Analysis

The normality of the data distribution was estimated using StatPlus:mac software (AnalystSoft Inc., Alexandria, VA, USA). Student's *t*-test or the Wilcoxon matched-pairs signed-rank test was used unless otherwise noted. The null hypothesis was rejected when the *P*-value was smaller than or equal to the significance level ($\alpha=0.05$).

3. Results

3.1. Blood Endocan Levels Dropped Below the Detection Limit After the Surgical Removal of Primary Breast Tumors from Nude Mice

MDA-MB-231BR/*mVenus-Akaluc* was generated to noninvasively visualize tumor recurrence and metastasis. Fluorescence and luminescence imaging confirmed the expressions of *mVenus* fluorescent protein (Figure 1b,c) and *Akaluc* enzyme (Figure 1d) in MDA-MB-231BR/*mVenus-Akaluc*. The doubling time of the generated cell line was 22.7 ± 1.9 h, while that of non-transfected original cells (MDA-MB-231BR) was 21.8 ± 2.1 h ($P=0.444$), confirming that the transfection of *mVenus-Akaluc* with PiggyBac™ Transposon Vector System did not affect cell proliferation.

At seven days after inoculation of the MDA-MB-231BR/*mVenus-Akaluc*, tumor volume reached 56–110 mm³, and blood endocan levels were 0.73 ng/mL (range; 0.38–1.54) (Figure 1f). When the tumor was resected, plasma endocan levels were below the detection limit of the ELISA kit three days after the surgery. Although weak luminescent signals were detected 131 and 145 days after surgery (Figure 1g), blood endocan levels remained below the detection limit (Figure 1f).

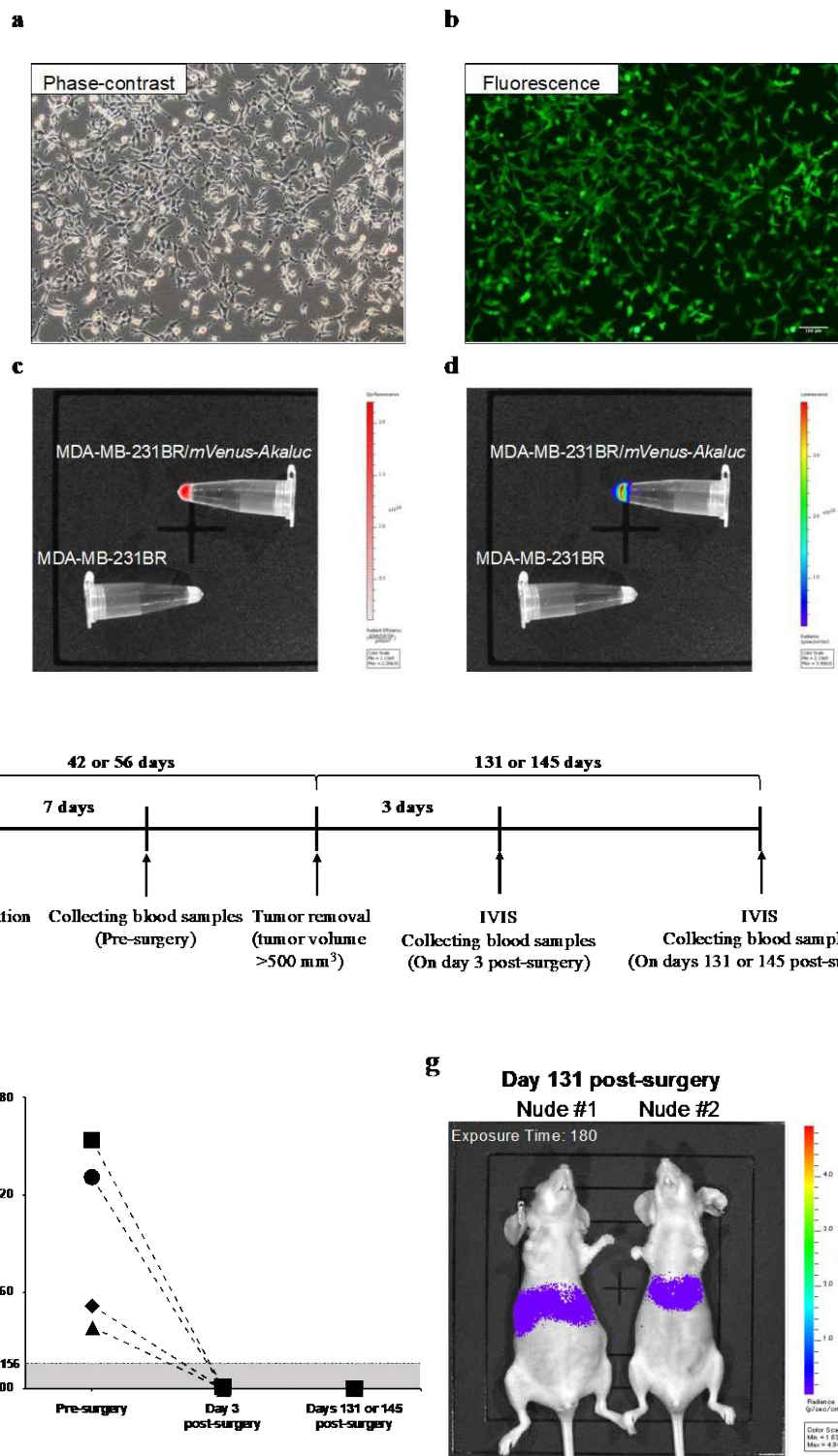


Figure 1. Effect of removing primary breast tumors on blood endocan levels of nude mice. (a,b), Fluorescence micrographs of MDA-MB-231BR/mVenus-Akaluc ((a), phase-contrast; (b), fluorescence). (c,d), Optical imaging ((c), fluorescence; (d), luminescence) of cell pellets of MDA-MB-231BR/mVenus-Akaluc (top) and non-transfected original cells (bottom). e, Schedule of in vivo experiments using nude mice ($N = 4$). (f), Changes in blood endocan levels of nude mice. The shaded area indicates endocan concentration below the detection limit (0.156 ng/mL). Each point represents the endocan concentration in an individual mouse. (g), Representative bioluminescence images of nude mice on day 131 after the surgery. Exposure time = 180 s.

3.2. Blood Endocan Levels Showed a Similar Pattern in NSG mice and Breast Cancer Patients After the Surgical Removal of the Primary Breast Tumor

Since elevation of blood endocan levels in the status of recurrence or metastasis was not observed in nude mice, more susceptible NSG mice were used in a similar experiment (Figure 2a). At seven days after inoculation of MDA-MB-231BR/*mVenus-Akaluc*, tumor size reached 77-147 mm³ and endocan was detected in all eight mice (range; 0.60-2.54). When the tumor was resected, blood endocan levels dropped in all but one mouse (Figure 2b). Luminescence imaging displayed weak luminescence signals in distant organs in some mice (Figure 2c).

To investigate how blood endocan levels respond to tumor resection in the clinical setting, we measured the blood endocan levels of 16 breast cancer patients pre- and post-surgery. Eight patients tested positive for endocan; the endocan level for pre-surgery ranged from 2.24 to 19.64 ng/mL. Blood endocan levels decreased after the surgery except for two patients (Figure 2d).

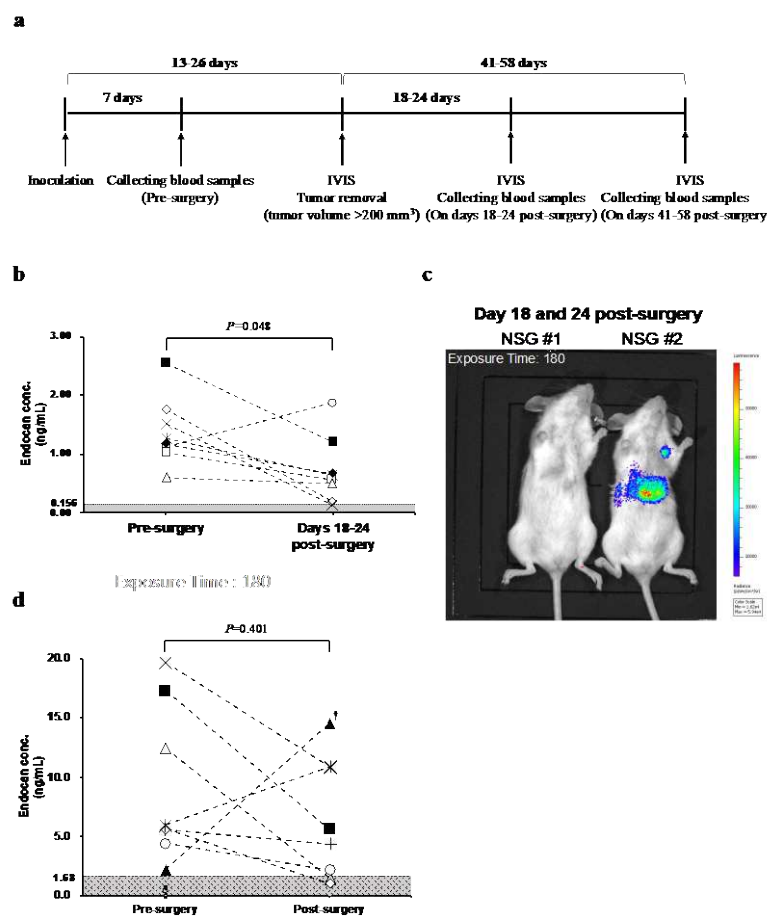


Figure 2. Changes in blood endocan levels before and after the surgical removal of primary breast tumors in NSG mice and breast cancer patients. **(a)**, Schedule of *in vivo* experiments using NSG mice ($N = 8$). **(b)**, Changes in blood endocan levels of NSG mice. The shaded area indicates endocan concentration below the detection limit (0.156 ng/mL). Each point represents the endocan concentration in an individual mouse. The statistical difference in blood endocan levels of NSG mice before and after the surgery was analyzed using a paired *t*-test. **(c)**, Representative bioluminescence images of NSG mice on days 18 and 24 after the surgery. Exposure time = 180 s. **(d)**, Changes in blood endocan levels of breast cancer patients ($N = 16$). The shaded area indicates endocan concentration below the cutoff value (1.68 ng/mL). Patients who showed blood endocan levels below the cutoff value were excluded from postsurgical endocan measurements ($N = 8$). †: The patient ran a marathon the day before the postsurgical blood collection. The statistical difference in blood endocan levels of breast cancer patients before and after the surgery was analyzed using a Wilcoxon matched-pairs signed-rank test.

3.3. Changes in Blood Endocan Level in Response to Breast Cancer Recurrence

In order to examine the changes in blood endocan levels with the growth of metastatic recurrence, the animals were monitored until recurrent tumor burdens were evident. On days 41-58, metastatic recurrence was observed with strong luminescent signals in all mice tested (Figure 3a). Relative to 18-24 days post-surgery, blood endocan levels increased on days 41-58 post-surgery in three out of six mice (Figure 3b). There was no correlation between the luminescence intensity of the metastases and blood endocan levels.

We next assessed the clinical benefit of blood endocan as a blood-based biomarker for breast cancer recurrence. Endocan in the blood of the patients diagnosed with postoperative recurrence ($N=20$) was measured, and the positivity rate for endocan was compared with that for CEA and CA15-3. As summarized in Table 1, 13 patients (65%) tested positive for endocan, which was much higher than CEA-positive patients (35%) and CA15-3-positive patients (25%). Moreover, among 13 patients who tested negative for CEA, eight patients (61.5%) were positive for endocan, and among 15 patients who tested negative for CA15-3, 10 patients (66.7%) were positive for endocan.

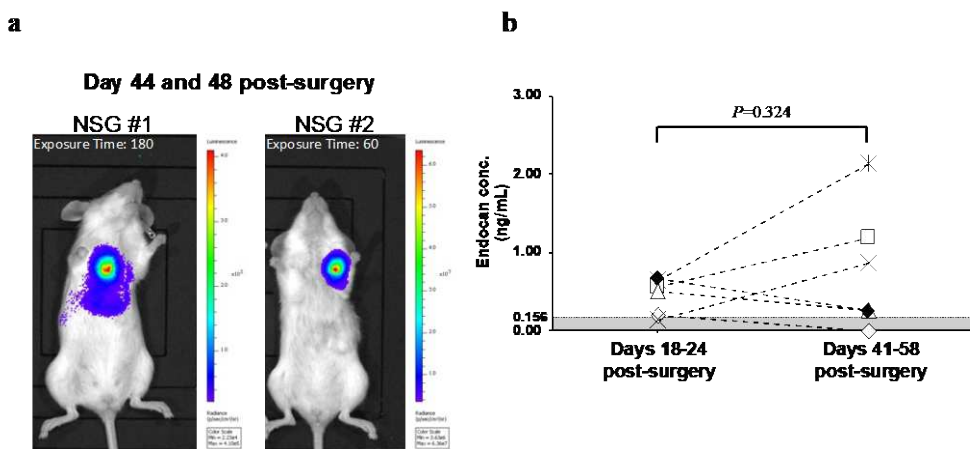


Figure 3. Effect of breast cancer recurrence on blood endocan levels of NSG mice ($N = 6$). (a), Representative bioluminescence images of NSG mice on days 44 and 48 after the surgery. Exposure time = 180 s. (b), Changes in blood endocan levels of NSG mice bearing recurrent breast tumors. The shaded area indicates endocan concentration below the detection limit (0.156 ng/mL). Each point represents the endocan concentration in an individual mouse. The statistical difference in blood endocan levels of breast cancer patients before and after the surgery was analyzed using a Wilcoxon matched-pairs signed-rank test.

Table 1. Comparison of endocan with CEA or CA15-3 in the blood samples of patients with postoperative breast cancer recurrence as a blood-based biomarker.

	CEA-Negative Patients ($N=13$; 65%)	CEA-Positive Patients ($N=7$; 35%)	CA15-3-Negative Patients ($N=15$; 75%)	CA15-3-Positive Patients ($N=5$; 25%)
Endocan-negative patients ($N=7$; 35%)	5/13 (38.5%)	2/7 (28.6%)	5/15 (33.3%)	2/5 (40%)
Endocan-positive patients ($N=13$; 65%)	8/13 (61.5%)	5/7 (71.4%)	10/15 (66.7%)	3/5 (60%)

4. Discussion

For successful clinical translation of endocan as a blood-based breast cancer biomarker, it is critical to be accurately reflected by the status of tumors. Surgical resection of primary breast tumors is expected to reduce blood endocan levels, while recurrence potentially increases blood endocan

levels. As expected, the removal of primary tumors decreased blood endocan levels of the majority of tumor-bearing mice and breast cancer patients. However, most of them did not reach an undetectable level, except for nude mice. Since MDA-MB-231BR/*mVenus-Akaluc* cells are the only source of human endocan in animal models, a large number of cells that are sufficient for detectable levels of endocan possibly remained in the body of NSG mice.

Although luminescence signals were detected in nude mice long after the tumor resection, luminescence signals that appeared in nude mice were much weaker than those detected in NSG mice. NSG mice are more susceptible to engraftment, growth, metastasis, and recurrence of xenogeneic cancer cells than conventional immunodeficient mice, including nude mice [30,31], which may explain why the number of MDA-MB-231BR/*mVenus-Akaluc* cells that remained in the body of nude mice was small and did not produce enough endocan detectable in the peripheral blood. In addition, there was no correlation between the luminescence intensities of recurrent tumors and blood endocan levels. *ESM1* expression levels were different in an individual cell obtained by single-cell cloning of MDA-MB-231BR/*mVenus-Akaluc* cells (Supplementary Fig. 2), implying that intratumoral heterogeneity of MDA-MB-231BR/*mVenus-Akaluc* cells could be responsible for the inconsistency between luminescence intensities and blood endocan levels in the NSG mouse model.

The case is more complicated for breast cancer patients, however, because various diseases, including inflammatory disease [27], diabetes [28], and sepsis [29], are known to increase blood endocan levels. Although such patients were excluded from this study, unnoticeable inflammation or undeclared events might have occurred during the study, which might be why the blood endocan levels were not entirely down to below the cutoff value.

Blood endocan levels of a few individuals were increased even after the surgery, both in NSG mice and breast cancer patients. Although the blood endocan level in one patient whose blood was collected on day 29 post-surgery was increased, that in other two patients whose blood was collected on day 32 post-surgery was decreased (Supplementary Table S3), implying that surgical inflammation was not considered to affect the postsurgical blood endocan levels. Therefore, other than one patient who ran a marathon, which might cause an increase in the blood endocan level, the day before the postsurgical blood collection, it is difficult to identify the reasons for the discrepancy with limited information and sample size.

CEA and CA15-3 have been used in clinical practice for years but are not well supported by clinical evidence [12,32]. Despite a limited number of patients participating in this study, endocan seems more responsive to breast cancer recurrence than CEA and CA15-3. Due to technical issues, CEA and CA15-3 in mouse models were not measured in this study, as it is necessary to obtain sufficient amounts of mouse plasma in order to quantitate all three markers by each ELISA kit. More importantly, the main objective of this study is not just to compare blood endocan with the existing biomarkers, but eventually to show the “clinical” benefit of blood endocan measurement. Since CEA and CA15-3 are routinely measured in clinical settings, we can compare blood endocan with the existing biomarkers of the same patient, which is more valuable information than preclinical comparisons for future clinical use. Still, one-fourth of the patients with breast cancer recurrence were negative for all three markers (Supplementary Table S4). Not all breast cancer patients have primary tumors expressing endocan, and the patients with postoperative recurrence participating in this study were not prescreened by *de novo* endocan expression. A longitudinal study tracking endocan-positive breast cancer patients is necessary to corroborate the clinical benefit of blood endocan measurements.

In order to achieve the clinical use of blood endocan measurement for breast cancer recurrence, several possible limitations need to be addressed. First, as described above, blood endocan rises in patients with inflammatory disease, diabetes, and sepsis. Therefore, it is important to include patients without breast cancer recurrence and compare with patients with breast cancer recurrence in order to determine if blood endocan levels are specifically elevated when breast cancer recurrence occurs. However, providing conclusive evidence of “non-recurrence” in patients is challenging as physicians typically do not perform diagnostic imaging if patients do not show any abnormalities during routine follow-up, leaving the recurrence status undetermined. Since performing additional diagnostic tests

to confirm “non-recurrence” may impose medical costs, potential radiation exposure from mammography, and unnecessary tissue diagnosis, this study primarily focused on the sensitivity of the blood endocan measurement and left the specificity for future studies. Generally speaking, sensitivity is more important when the consequence of missing a positive case is serious, such as in screening for a life-threatening disease, like cancer, which is the case for this study. In addition, some ELISA kits from other suppliers did not work well with measuring blood endocan in our study. Epitopes that anti-endocan antibodies recognize may vary depending on each ELISA kit supplied by manufacturers, which might be responsible for inconsistent results when ELISA kits from different suppliers were used. We have worked on developing a new alternative to ELISA for future clinical use of blood endocan measurement.

5. Conclusions

This study is the first report on the effect of the surgical resection of primary breast tumors and recurrence on blood endocan levels, and our findings are consistent with the earlier reports for low-grade glioma [33] and renal cell carcinoma [23]. More importantly, the current study did demonstrate that blood endocan measurement could detect breast cancer recurrence with a higher probability compared to conventional tumor markers, CEA and CA15-3, in a limited number of patients ($N = 20$). Although this study deals with only a limited number of patients, this positive finding is worth being validated by future large-scale studies, which would strengthen the robustness of the findings. According to clinicaltrials.gov search, many clinical trials regarding endocan levels in the peripheral blood of patients suffering from various diseases are ongoing or completed, which warrants the application of blood endocan measurements to breast cancer. Taken together, our findings indicate that blood endocan could be a more sensitive marker than CEA or CA15-3 in breast cancer patients with postoperative recurrence.

Supplementary Materials: The following supporting information can be downloaded at the website of this paper posted on Preprints.org. Figure S1. Pathological assessment of endocan expression in primary breast tumors removed by the surgery. Figure S2. Difference in *ESM1* expression of MDA-MB-231BR/mVenus-Akaluc obtained by limiting dilution cloning. Table S1. Clinical characteristics of breast cancer patients whose blood was collected before and after the surgery. Table S2. Clinical characteristics of breast cancer patients with postoperative recurrence. Table S3. Effect of the interval between the surgery and the blood collection on changes in blood endocan levels. Table S4. Endocan, CEA, and CA15-3 concentrations in the blood samples of patients with postoperative breast cancer recurrence.

Authors' contributions: Conceptualization, SN and YKato; Data curation, KD, YKanada, AN and KT; Formal analysis, KD, TY and HO; Funding acquisition, YKato; Investigation, KD, YKanada, AN, HO and YKato; Methodology, KD, YKanada, AN, KT, KI, TY, FS and YK; Project administration, SN and YKato; Resources, YKanada, AN and KT; Supervision, KI, FS, SN and YKato; Validation, KI; Writing – original draft, KD; Writing – review & editing, FS, SN and YKato. All authors read and approved the final manuscript.

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Institutional Review Board Statement: The institutional animal care committee of Hoshi University approved the animal experiments (Protocol #: 19-083). Clinical studies were approved by the Institutional Review Board from Showa University Hospital (#2880) and Hoshi University (#2019-12, #2021-04).

Informed Consent Statement: Written informed consent was taken from each volunteer with the assistance of Showa University Hospital.

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Conflicts of interest: The authors declare that they have no competing interests.

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