Abstract. The current study clarified the accuracy of a circulating tumor cell (CTC) detection system to diagnose colorectal cancer using blood samples. The system uses the ‘polymeric CTC-chip,’ which is a microfluidic device that is used for CTC isolation. CTCs are considered sensitive diagnostic biomarkers. However, their concentration in the peripheral blood is low and requires highly sensitive and specific capturing techniques. The capture efficiency of the polymeric CTC-chip was first assessed using cell suspensions of the colorectal cancer cell line HCT-116, which was reported as 90.9% in a phosphate-buffered saline suspension and 65.0% in the blood. The CTC-chip was then used to detect CTCs in blood samples obtained from 13 patients with stage II-IV colorectal cancer. On average, the CTCs/ml was lower in patients with stages II and III colorectal cancer (3.3±2.3) than in those with stage IV (7.0±6.2). In patients with stages II-IV, 92% had ≥1 CTC per ml, which was significantly higher than the positive rate (15%) detected using the carbohydrate antigen 19-9 test (CA19-9). Furthermore, CTCs were detected in all patients with stage II and III colorectal cancer, including a number of patients with negative results for the carcinoembryonic antigen (CEA) and CA19-9 tests. With the polymeric CTC-chip detection system, CTCs can be effective cancer markers, particularly for patients with stage II and III colorectal cancer who often exhibit negative conventional serum marker test results. The CTC-chip system may also facilitate the detection of cancer progression based on CTC concentration.

Introduction

Colorectal cancer is the second most common cancer in women (9.5%) and the third in men (10.2%) and ranked as the second most common cause of cancer death (9.2%) worldwide (1). Therefore, sensitive and noninvasive diagnoses are important to improve its treatment outcomes. However, conventionally used biomarkers in blood samples, such as carcinoembryonic antigen (CEA) and carbohydrate antigen 19-9 (CA19-9), do not always indicate the pathological aspects of malignancy (2).

The use of circulating tumor cells (CTCs) as the next-generation cancer marker has been an active research topic in the field of oncology for the past two decades (3-6). The peripheral blood CTC concentration is reported to be extremely low: <10 cells/ml in patients with metastatic cancer (3,7). However, CTCs are not present or scarcely detected in the blood of healthy individuals nor in those with nonmalignant diseases (8,9). Nucleic acid evaluation in CTCs and direct enumeration of CTCs are typical methods used to detect CTCs requiring highly sensitive techniques. To evaluate nucleic acid levels in blood CTCs, related gene expressions are examined using reverse transcription-quantitative PCR (RT-qPCR), or DNA arrays (9-13). Cell capturing methods based on size or surface antibodies (6,10), such as the CellSearch™ System (Veridex LLC) (CellSearch) and other microfluidic devices, are used for direct enumeration (14-16). CellSearch is a semi-automated system for CTC quantification as a diagnostic method for metastatic breast, colorectal, and prostate cancers approved by the US Food and Drug Administration (FDA) (17,18). Moreover, the efficacy of this system has been reported in metastatic breast, prostate, esophageal, and colorectal cancers (19-25). However, the use of CellSearch is limited due to its high cost (26-28) and low sensitivity to some cancer types, such as hepatocellular carcinoma (28,29). Approximately 18% of non-metastatic and 41% of metastatic patients with colon cancer are positive with CTCs in the CellSearch system (30).

Nagrath et al (15) developed a microfluidic device known as the ‘CTC-chip’ to overcome these limitations. The ‘CTC-chip’ facilitates efficient and selective separation of CTCs from whole blood samples, mediated by the interaction of target CTCs with antibody-coated microposts under precisely
controlled laminar flow conditions (15,31). Subsequently, a novel ‘polymeric CTC-chip’ was developed to isolate CTCs, with lower cost, high transparency that facilitates observation through the chip, and convertibility of antibodies to coat the surface to arrest cancer cells than that of the existing CTC-chips (32-36).

In the present study, the capture efficiency of the polymeric CTC-chip was measured using colorectal cancer cells spiked in phosphate-buffered saline (PBS) or healthy whole blood at first. Next, CTCs in clinical blood samples were detected in patients with colorectal cancer. The sensitivity of CTC detection in the blood samples of patients with colorectal cancer was compared with that of the CEA and CA19-9 tests.

**Materials and methods**

**Preparation of cancer cells.** HCT116 (ATCC® CCL-247™) colorectal cancer cells were cultured and exhibited a high expression of epithelial cell-adhesion molecule (EpCAM), in McCoy's 5A medium (cat. no. 16600082; Invitrogen) with 10% fetal bovine serum and 1% penicillin-streptomycin at 37°C in a humidified 5% CO₂ atmosphere. Then, the EpCAM expression in HCT116 cells was evaluated with a flow cytometer (FACSVerso; BD Biosciences) using a PE/Cy7-conjugated anti-human CD326 (EpCAM) antibody (cat. no. 324221; BioLegend) and FlowJo software (ver.9; FlowJo LCC). To determine the EpCAM localization in the cells, Alexa Fluor® 594-conjugated anti-human CD326 (EpCAM) antibodies (cat. no. 324228; BioLegend) at 5 µg/ml was added to the HCT116 cell suspension; the mixture was allowed to sit for 2 h at room temperature and examined using a fluorescence microscope system (BZ-X710; Keyence) in a 24-well plastic dish (a cell culture plate with a lid; Sigma-Aldrich).

**Preparation of cancer cell suspensions.** To measure the capture efficiency, HCT116 cells were fluorescently labeled using the Cell Explorer™ Live Cell Tracking kit (cat. no. 22621; AAT Bioquest). The cells were spiked in PBS containing 5% bovine serum albumin (BSA; PBS suspension) or the whole blood obtained from a healthy donor and stored in a vacuum blood collection tube containing ethylenediaminetetraacetic acid (EDTA; VP-DK052K; Terumo; blood suspension) at 4°C. All cell suspensions were prepared at approximately 1,000 cells/ml concentration, and the precise concentration of each suspension was determined.

**Antibody coating on the chip surface.** An antibody coating of the polymeric CTC-chip surface was determined using the method described by Ohnaga et al (32), with the process outline illustrated in Fig. 1. The chip was washed with 70% ethyl alcohol once for hydrophilization and then exposed to goat anti-mouse IgG antibodies overnight (cat. no. 1032-01; Southern Biotech) in PBS at a 25 µg/ml concentration at 4°C. Then, the chip surface was washed with PBS once to remove any non-bonded anti-IgG antibodies and kept wet. Next, the chip surface was coated with mouse anti-human EpCAM antibodies (cat. no. sc-59906; Santa Cruz Biotechnology) in PBS at a 25 µg/ml concentration and stored at room temperature for 1 h. The chip was washed with PBS again after the antibody treatment.

**CTC capturing system and evaluation of the cell-capture efficiency.** The sample flow and CTC capturing were performed using the method described by Ohnaga et al (32). The workflow of CTC detection with the polymeric CTC-chip is outlined in Fig. 2. Briefly, the polymeric CTC-chip coated with antibodies was set in a holder and fixed on an inverted fluorescence microscope stage (CKX41; Olympus). The size of the polymeric CTC-chip was 75x25 mm, and surface micro-structures comprised two types of micropost arrays in a wide channel (32). The two ports of the holder were then connected to a syringe or sample tube with tubing and fittings, which allowed the liquid sample to flow through the channel (32). Then, the sample tube was shaken to prevent cell precipitation and adhesion. Each sample was sent to the chip using a syringe pump at a constant flow rate of 1 ml/h. After allowing the samples to flow through the polymeric CTC-chip, the tube was washed with PBS containing 5% BSA once at 1 ml/h for 15 min to remove the suspended cells. The chip surface was then examined using an inverted fluorescent microscope (CKX41) equipped with a digital video camera (HDR-CX555; Sony) during the flow test. In this study, three types of polymeric CTC-chips were used to assess the capture efficiency with PBS and blood suspensions: i) No antibody treatment (non-treated chip); ii) a goat anti-mouse IgG antibody solitary coating (IgG-chip); and iii) a primary coating of goat anti-mouse IgG antibody and a secondary coating of mouse anti-human EpCAM antibody (EpCAM-chip). Notably, the flow test was repeated three to four times for the two suspensions using each of the three types of chips.

Thus, the number of cells remaining in the chip after the flow test (Nr) was determined. The number of cells that passed through the chip inlet (Np) had been evaluated before the test based on the cell concentration in ea43ch suspension. The cell-capture efficiency, defined as Nr/Np, was evaluated (32) for each test.

**Characteristics of patients and volunteers.** We enrolled 13 patients (age range, 59-77 years; men, 9; women, 4) with stages II-IV (UICC classification) colorectal cancer (37) and 2 healthy volunteers (control group), without detectable cancer or serious diseases, in the Department of Coloproctological Surgery, Juntendo University Hospital (Tokyo), from August 2015 to March 2016 (Table I). Histological features and differentiation grades of cancer tissue samples were evaluated in the Department of Diagnostic Pathology of the Juntendo University Hospital, according to JSCCR classification (38): Each patient was classified into the following categories: well-differentiated tubular adenocarcinoma (tub1), moderately differentiated tubular adenocarcinoma (tub2), or poorly differentiated adenocarcinoma (por). In addition, CEA and CA19-9 levels in blood samples drawn from the participants before receiving any cancer treatment were evaluated. In this study, a value was considered ‘positive’ when higher than the set conventional cut-off values (5.0 ng/ml for CEA and 37.0 U/ml for CA19-9) (39-41). This study was conducted in accordance with the Declaration of Helsinki and was approved by the Ethics Committee of Juntendo University (cat. no. 2015036). Furthermore, written informed consent was obtained from all participants after adequate counseling using written documents that described the research aim and any possible risks involved.
Detection of CTCs from blood samples. Using a vacuum blood collection tube containing EDTA, 2 ml of blood samples were drawn from all patients one day preoperatively. In patients with stages II and III cancer, blood samples were obtained again 6 days postoperatively. The blood samples were then placed in the polymeric CTC-chip coated with both goat anti-mouse IgG antibodies and mouse anti-human EpCAM antibodies at a flow rate of 1 ml/h for 1 h, as previously described. After setting the blood samples, the chip was washed with PBS containing 5% BSA at a flow rate of 2 ml/h for 30 min to remove the blood cell constituents that were nonspecifically combined with the chip. After using a 15% formaldehyde to anchor the cells, the chip was washed with 0.1% Triton X-100. Then, the aqueous antibody solution was successively introduced into the chip.

The FDA-approved technique to detect CTCs relies on the use of antibodies that target EpCAM, followed by cytokeratin (CK) and CD45 staining, to confirm the epithelial phenotype (42). In the present study, fluorescein isothiocyanate (FITC)-conjugated anti-mouse CK 8 + 18 (207) antibodies (ab190366; Abcam Plc, Cambridge, UK; 5 µl: 50 µg/ml) and Alexa Fluor® 594-conjugated anti-mouse CD45 antibodies (cat. no. 103144; BioLegend; 5 µl: 50 µg/ml) were used to differentiate epithelial adenocarcinoma cells and leukocytes. Diamidino-2-phenylindole dihydrochloride (DAPI; 20 µl: 1.5 µg/ml), a nuclear staining reagent, was also used to evaluate the presence/absence of cell nuclei. A cell was defined as to contain CTC when it was positive for both DAPI and CK but negative for CD45. Furthermore, CTCs remaining on the chip were observed and counted with images obtained using the inverted fluorescence microscope (CKX41).

Statistical analyses. The effects of antibody treatments (none, IgG, and IgG + EpCAM) of the polymeric CTC-chips on the cell-capture efficiencies were examined using an analysis of variance (ANOVA) after the angular transformation on the cell-capture efficiency defined as Nr/Np (the captured and numbers of cells that passed through the chip), as previously described in both PBS and blood suspensions. Using Welch’s two-sample t-test, effects of the polymeric CTC-chip...
treatment on capture efficiency were assessed based on the difference between the average cell-capture efficiency values and the angular transformation between two different treatments: No treatment, IgG-chip; no treatment, EpCAM-chip; and IgG-chip, EpCAM-chip (in both PBS and blood suspensions). Welch’s t-test was also used to evaluate differences in the number of CTCs detected in blood samples obtained from patients with stages II-III and IV cancer. The paired t-test was used to determine the difference in numbers of CTCs/ml between blood samples pre- and postoperatively in patients with stages II-III cancer. Moreover, the Fisher exact test was used to compare the rates of patients detected with CTCs and those positive for CEA and CA19-9 tests. All statistical analyses were performed using the R software (v.3.2.5) (43).

Results

EpCAM expression of HCT116. The EpCAM expression in HCT116 cells was evaluated with flow cytometry using anti-EpCAM antibodies, showing that 97.3% of HCT116 cells were positive for EpCAM (Fig. 3). The immunofluorescence staining suggested that EpCAM was localized on the cell surface (Fig. 4).

Capture efficiency of the polymeric CTC-chip covered with antibodies. Capturing efficiencies of colorectal cancer cells spiked in PBS and blood suspensions were tested using the
polymeric CTC-chip by conducting three different treatments: i) Primarily coated with goat anti-mouse IgG antibodies and secondarily coated with mouse anti-human EpCAM antibodies, ‘EpCAM-chip’; ii) coated only with goat anti-mouse IgG antibodies, ‘IgG-chip’; or iii) no antibody treatment, ‘non-treated chip.’ Each suspension sample was set on the chip using a syringe pump at 1 ml/h for 1 h. We determined that 1,109.4±392.8 (average number ± SD) cells in the PBS suspension and 1,160.8±119.9 cells in the blood suspension passed through the chip inlet during the suspension sending period. In the PBS suspension, capture efficiencies were 0.909±0.053 (average rate ± SD) for the EpCAM-chip (n=4), 0.060±0.010 for the IgG-chip (n=3), and 0.066±0.007 for the non-treated chip (n=3; Fig. 5), whereas in the blood suspension, the efficiencies were 0.650±0.064 for the EpCAM-chip (n=4), 0.066±0.016 for the IgG-chip (n=3), and 0.077±0.002 for the non-treated chip (n=3; Fig. 5).

Significant differences in the cell-capture efficiency were determined after the angular transformation among the three chip treatments in the PBS (P=7.610x10^-8) and blood (P=3.336x10^-7) suspensions using ANOVA. In the PBS suspension, the capture efficiency values after an angular transformation assessed using Welch’s t-test were significantly higher in the EpCAM-chip than those in the non-treated chip (n=3; Fig. 5).

**Table I. Patient characteristics with detected number of CTCs and CEA and CA19-9 values.**

| Patent no. | Age | Sex | Site | Histologic features | TNM classification | Stage | Before surgery | After surgery | CEA (ng/ml) | CA 19-9 (U/ml) |
|------------|-----|-----|------|---------------------|--------------------|-------|----------------|--------------|-------------|---------------|
| 1          | 74  | M   | A    | tub2                | T3 N0 M0           | II A  | 1             | 2            | 7.9         | 37            |
| 2          | 77  | F   | D    | tub2                | T4a N0 M0          | II B  | 6             | 2            | 6.8         | 7             |
| 3          | 76  | M   | Rb   | tub2                | T3 N0 M0           | II A  | 2             | 0            | 2.9         | 23            |
| 4          | 77  | M   | S    | tub1                | T4b N0 M0          | II C  | 4             | 1            | 4.9         | 34            |
| 5          | 66  | F   | C    | tub2                | T3 N1b M0          | III B | 1             | 0            | 23.6        | 24            |
| 6          | 59  | M   | Rb   | tub2                | T3 N2b M0          | III C | 6             | 3            | 3.5         | 8             |
| 7          | 70  | M   | RS   | tub2                | T3 N1b M1a (H2)    | IVA   | 12            |              | 387.7       | 94            |
| 8          | 72  | M   | A    | tub2                | T3 N0 M1a (H1)     | IVA   | 7             |              | 14.3        | 21            |
| 9          | 74  | M   | RS   | tub1                | T3 N2b M1a (LYM)   | IVA   | 18            |              | 12.8        | 27            |
| 10         | 67  | M   | S    | tub1                | T4b N2a M1a (LYM)  | IVA   | 5             |              | 34.8        | 6             |
| 11         | 77  | F   | A    | tub2                | T4a N2a M1a (H2)   | IVA   | 2             |              | 15.8        | 20            |
| 12         | 61  | F   | A    | tub2                | T3 N2b M1b (H1 PUL1) | IVA | 5             |              | 7.6         | 6             |
| 13         | 71  | M   | Ra   | por                 | T3 N1a M1a (H1)    | IVA   | 0             |              | 1.4         | 6             |

‘Postoperative samples were not examined. M, male; F, female; Site: C, cecum; A, ascending colon; D, descending colon; S, sigmoid colon; RS, rectosigmoid; upper R, upper rectum; lower R, lower rectum; tub, tubular adenocarcinoma; tub1, well differentiated; tub2, intermediate differentiated; por, poorly differentiated adenocarcinoma; TNM, tumor node metastasis; CTCs, circulating tumor cells; CEA, carcinoembryonic antigen; CA19-9, carbohydrate antigen 19-9.'
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(P=1.077x10⁻⁴, df=3.220) and the IgG-chip (P=6.994x10⁻⁵, df=3.426). In addition, values in the non-treated chip did not differ from those in the IgG-chip (P=0.4384, df=3.617).

A similar tendency in the blood suspension was observed, showing that capture efficiency values after an angular transformation two-tailed Welch's t-test were significantly higher in the EpCAM-chip than those in the non-treated chip (P=2.647x10⁻⁴, df=3.028) and IgG-chip (P=2.180x10⁻⁵, df=4.561). Values in the non-treated chip did not differ from those in the IgG-chip (P=0.3794, df=2.057). Furthermore, the capture efficiency of the EpCAM-chip was significantly higher in the PBS suspension than that in the blood suspension (P=0.001071, df=5.643).

Detection of CTCs in blood samples of patients with colorectal cancer and healthy volunteers. CTCs in the preoperative peripheral blood samples of 13 patients with stage II-IV colorectal cancer were enumerated on the chip (Figs. 6 and 7, Table I). The mean number of CTCs/ml in patients with stages II and III cancers (3.3±2.3, n=6) tended to be lower than those with stage IV, with near-marginal significance (7.0±6.2, n=7; t=1.4563, P=0.0919, one-tailed Welch's t-test). No CTCs were detected in one patient with stage IV cancer, which was different from all other cases of well or moderately differentiated tubular adenocarcinoma (tub1 or tub2).

After excluding this patient, the mean number of CTCs/ml associated with stage IV cancer (8.2±5.8, n=6) tended to be higher than those with stages II and III cancer (3.3±2.3, n=6), with difference that is close to being statistically significant (t=1.8806, P=0.0524, one-tailed Welch's t-test).

The number of CTC/ml in blood samples postoperatively (mean ± SD: 1.3±1.2) was significantly lower than that preoperatively (3.3±2.3) in patients with stages II and III cancers (n=6; t=2.7386, P=0.0204, one-tailed paired t-test). In addition, no CTCs were detected in two healthy volunteer blood samples on polymeric CTC-chips.

The sensitivity of CTC detection using the polymeric CTC-chip compared with conventional biomarkers. The detection rate of CTCs was compared with that of positive rates detected using cancer markers. Nine patients had CEA or CA19-9 levels of higher than the cut-off values (CEA, 5.0 ng/ml; CA19-9, 37.0 U/ml) (Table I; Fig. 7). At least one CTC (per ml) was detected in 12 of 13 patients, whereas the CEA and CA19-9 levels were positive, i.e., above the cut-off values, in nine and two patients, respectively. For all cancer stages (II-IV), a significant difference was observed between the positivity rate from CTC detection and that from two biomarkers (P=0.0002251 according to the 3x2 two-sided Fisher's exact test). In addition, the detection rate of CTCs in all patients was higher than that of CA19-9 (P=0.0002127, Fisher's exact test) but not significantly higher than that of CEA (P=0.3217). These two conventional markers exhibited negative levels in the patient (no. 13 in Fig. 7) in whom no CTCs were detected. In all six patients with stage IV cancer (except for no. 13), both CTC and CEA values were positive, whereas the CA19-9 value was positive in only one patient. Although CTCs were
detected in all six patients with stages II and III cancer, CEA and CA19-9 values were positive in only two and one, respectively, of the four patients with stage II cancer, and in one and no patient, respectively, in two patients with stage III cancer.

**Discussion**

Capturing device development is expected to make CTCs a sensitive clinical biomarker to diagnose and predict the prognosis and treatment effects in cancers (6,10,16). Using one of the new microfluidic devices to capture CTCs, the polymeric CTC-chip, CTCs were detected in the blood samples of most patients with colorectal cancer (12 of 13) who participated in this study. The CTC detection was confirmed as more sensitive than the two most common conventional marker tests for the diagnosis of colorectal cancer.

Newly developed devices for CTC detection are typically examined based on their capture efficiency of the target cancer cells (15,16,44,45). In this study, the efficiency of the polymeric CTC-chip was first assessed on the colorectal cancer cell line, HCT116. The polymeric CTC-chip secondarily treated with the anti-EpCAM antibody exhibited a considerably high capture rate for HCT116 cells: 90% in the PBS suspension and 65% in the whole blood suspension, which were comparable to or exceeded those reported in previous studies (15,16,44,45). The efficiency seemed to exceed with those in existing devices. Nagrath et al (15) reported that the capture efficiency of the silicon microchip was 65-80% in the PBS suspension for several cell lines, which fit the prediction that HCT116, an EpCAM-positive cell line, could be captured with the polymeric CTC-chip treated with the anti-EpCAM antibody. However, 5-7% of CTCs were captured using the chip without the anti-EpCAM antibody treatment, demonstrating that the chip trapped some cells with nonspecific bonds.

The number of CTCs/ml tended to be higher in patients with stage IV than those with stages II and III cancers. Thus, CTC concentration seemed to increase with cancer state progression. Moreover, the number of CTCs/ml in patients with stages II and III cancers postoperatively became lower than that preoperatively. These results suggested that the polymeric CTC-chip system could potentially be used to monitor disease progression and the treatment effects in patients with colorectal cancer.

CTCs are considered specific to cancer and are not detected in the peripheral blood of healthy individuals (8,9). No CTCs were detected on the polymeric CTC-chip in the blood samples of two healthy volunteers. In this study, each patient was classified as ‘CTC positive’ when more than one CTCs were detected in the sample. The CTC positive rate in all stages tended to be higher than the positive rate obtained by blood tests with two conventional markers. CTCs were detected in all patients with stages II and III cancers (six patients), although three of them tested negative for CEA and five tested negative for CA19-9, suggesting that CTCs could be an effective cancer marker because of they are potentially detected in earlier stages than existing tumor markers.

The anti-EpCAM antibody binds with the EpCAM, which is typically expressed on epithelial cells (3,5,11). However, the use of an anti-EpCAM antibody might not adequately capture EpCAM-negative cancer cells (14,29,42,45). In this study, CTCs were not detected in the blood sample of one patient with stage IV cancer. In this case, CEA and CA19-9 values did not exceed the cut-off values despite cancer progression and distant metastasis. Histologically, the cancer tissue sample of this case was classified as poorly differentiated adenocarcinoma (por), whereas those of all other cases were classified as differentiated adenocarcinoma (tub1, 2). The cause of this failure of CTC detection remains unknown, although it may be related to the characteristics of the cancer tissue.

Other microfluidic devices treated with anti-PSMA, HER2, or epidermal growth factor receptor (EGFR) antibodies have detected CTCs in the blood samples of patients with prostate, breast, and lung cancers (45-47). Antibodies used in the secondary treatment on the polymeric CTC-chip are easily changeable. To take advantage of this feature, EpCAM-negative mesothelioma cells were captured using this polymeric CTC-chip treated with anti-podoplanin antibodies (34,35).

Using anti-EGFR antibodies on the surface of the polymeric CTC-chip, Ohnaga et al (36) recently reported a high capturing efficiency against several cell lines expressing EGFR.

Using alive CTCs alone, the expression level of oncogenes that are assumed to be related to colorectal cancer of a single cell or its offspring cells was analyzed. In circulating lung cancer cells collected from a microfluidic device, gene mutations in EGFR were detected (47). Future advancements of the CTC isolation technique in the polymeric CTC-chip will facilitate the evaluation of molecular markers.

This study validated the usefulness of CTC detection in blood samples obtained from patients with colorectal cancer using the polymeric CTC-chip; nevertheless, the number of participants was limited. Further studies should be conducted with more patients, including those in earlier stages of the disease, to assess the sensitivity of the polymeric CTC-chip in diagnosing colorectal cancer.

| Stage | II | III | IV | % positive |
|-------|----|-----|----|-----------|
| Patient No. | 1  | 2  | 3  | 4  | 5  | 6  | 7  | 8  | 9  | 10 | 11 | 12 | 13 | II,III | IV | II,III,IV |
| CTCs   | 100| 86 | 92 |
| CEA    | 50 | 86 | 69 |
| CA19-9 | 17 | 14 | 15 |

Figure 7. CTC detection using the polymeric CTC-chip and based on the positivity of CEA and CA19-9 in blood samples of patients with colorectal cancer. Shaded boxes indicated a positive case with at least one or more CTCs detected or CEA and CA19-9 values higher than the respective cut-off values (5.0 ng/ml and 37.0 U/ml). CTC, circulating tumor cell; CEA, carcinoembryonic antigen; CA19-9, carbohydrate antigen 19-9 test.
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Availability ofdata and materials

The datasets used and/analyzed during the present study are available from the corresponding author on reasonable request.

Authors’ contributions

HK designed the study and provided the overall guidance with advice from KS and YT. TO prepared the polymeric CTC-chip. Bray F, Ferlay J, Soerjomataram I, Siegel RL, Torre LA and Paterlini-Brechot P and Benali NL: Circulating tumor cells (CTC) detection: Clinical impact and future directions. Cancer Lett 253: 180-204, 2007.

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Competing interests

The authors declare that they have no competing interests.

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