Defining the dimensions of circulating tumor cells in a large series of breast, prostate, colon, and bladder cancer patients

Pauline A. J. Mendelaar¹, Jaco Kraan¹, Mai Van¹, Leonie L. Zeune², Leon W. M. Terstappen², Esther Oomen-de Hoop¹, John W. M. Martens¹ and Stefan Sleijfer¹

¹ Department of Medical Oncology, Erasmus MC University Medical Center, Cancer Institute, Rotterdam, The Netherlands
² Department of Medical Cell BioPhysics, University of Twente, Enschede, The Netherlands

Keywords
- cell morphology
- cell size
- circulating tumor cells
- metastatic cancer
- single cell isolation methods

Correspondence
P. A. J. Mendelaar, Erasmus MC University Medical Center, Room no: BE-414a, Dr. Molewaterplein 40, Rotterdam 3015 GD, The Netherlands and Mailbox 2040, Rotterdam 3000 CA, The Netherlands
Fax: +31 10 70 44 377
Tel: +31 10 70 44 375
E-mail: p.mendelaar@erasmusmc.nl

(Received 10 July 2020, revised 24 August 2020, accepted 10 September 2020, available online 4 October 2020)
doi:10.1002/1878-0261.12802

[Correction added on 21 December 2020, after first online publication: Peer review history is not available for this article, so the peer review history statement has been removed.]

1. Introduction

Circulating tumor cells (CTCs) in the blood of cancer patients are of high clinical relevance. Since detection and isolation of CTCs often rely on cell dimensions, knowledge of their size is key. We analyzed the median CTC size in a large cohort of breast (BC), prostate (PC), colorectal (CRC), and bladder (BLC) cancer patients. Images of patient-derived CTCs acquired on cartridges of the FDA-cleared CellSearch® method were retrospectively collected and automatically re-analyzed using the ACCEPT software package. The median CTC diameter (µm) was computed per tumor type. The size differences between the different tumor types and references (tumor cell lines and leukocytes) were nonparametrically tested. A total of 1962 CellSearch® cartridges containing 71,612 CTCs were included. In BC, the median computed diameter (CD) of patient-derived CTCs was 12.4 µm vs 18.4 µm for cultured cell line cells. For PC, CDs were 10.3 µm for CTCs vs 20.7 µm for cultured cell line cells. For CRC and BLC, CDs were 7.5 µm and 8.6 µm, respectively. Finally, leukocytes were 9.4 µm. CTC size differed statistically significantly between the four tumor types and between CTCs and the reference data. CTC size differences between tumor types are striking and CTCs are smaller than cell line tumor cells, whose size is often used as reference when developing CTC analysis methods. Based on our data, we suggest that the size of CTCs matters and should be kept in mind when designing and optimizing size-based isolation methods.

Abbreviations
- ACCEPT, Automated CTC Classification, Enumeration, and PhenoTyping software
- BC, breast cancer
- BLC, bladder cancer
- CD, computed diameter
- CEL, cultured tumor cell (cell line)
- CK, cytokeratin
- CRC, colorectal cancer
- CTC-L, circulating tumor cells derived from cerebrospinal fluid (liquor)
- CTCs, circulating tumor cells
- DAPI, 4′,6-diamidino-2-phenylindole
- EMT, epithelial–mesenchymal transition
- EpCAM, epithelial cell adhesion molecule
- IQR, interquartile range
- KW test, Kruskal–Wallis test
- MWU test, Mann–Whitney U test
- NCR, nucleus/cytoplasm ratio
- P2A, perimeter to area
- PC, prostate cancer
- TIF, tagged Image Format files
- TXT, text file
- µm, micrometer
- µm², square micrometers

Molecular Oncology 15 (2021) 116–125 © 2020 The Authors. Published by FEBS Press and John Wiley & Sons Ltd.
This is an open access article under the terms of the Creative Commons Attribution License, which permits use, distribution and reproduction in any medium, provided the original work is properly cited.
disease is estimated as 1 CTC per billion white blood cells, capturing them is technically challenging. This emphasizes the need for sensitive and specific CTC detection and isolation methods.

The CellSearch® system is the only FDA-cleared CTC enumeration platform [2]. This method uses antibodies recognizing the epithelial cell adhesion molecule (EpCAM) coupled to ferrofluids to enrich peripheral blood for CTCs. The immunomagnetically enriched cells are stained with the nucleic acid dye 4′,6-diamidino-2-phenylindole (DAPI) and labeled with fluorescent antibodies against cytokeratins 8, 18+, and/or 19+ (CK) and CD45. The processed blood fraction is transferred into a cartridge of which images are taken. An expert user of the CellSearch® method assesses all presented events and identifies CTCs manually. CTCs are defined as CK+/DAPI+/CD45− events, whereas leukocytes are CK−/DAPI+/CD45+ events. CellSearch-based CTC enumeration has prognostic value in multiple tumor types [3–5]. However, since the method relies on EpCAM positivity, it will only detect CTCs which express EpCAM. Epithelial-mesenchymal transition (EMT) is thought to play a key role in dissemination of cancer [6,7]. To a certain degree, epithelial markers are down regulated in CTCs which have undergone EMT. This can lead to a detection failure using the CellSearch® method if EpCAM expression is too low or absent. Hence, EpCAM-based detection methods may introduce a selection bias in CTCs which can be studied.

Therefore, antibody-independent CTC detection and isolation methods are being developed and many of them are based on physical characteristics [8–13]. Size-based methods frequently use cultured cell line cells to characterize the performance of their system. Although evidence is lacking, CTCs are considered to be larger and less deformable than blood cells. Additionally, it is suggested that morphological differences of CTCs exist between tumor types [14,15]. This implies the need for improved knowledge on CTC size from different tumor types. Therefore, we determined the median CTC size, computed the approximate median diameter (CD) of the cytoplasm and nucleus per cell, and determined the nucleus/cytoplasm ratio (NCR) of CTCs from breast (BC), prostate (PC), colorectal (CRC), and bladder (BLC) cancer patients. For this purpose, a large series of images of CellSearch® cartridges were re-analyzed through the ‘Automated CTC Classification, Enumeration and PhenoTyping software package’ (ACCEPT) (https://github.com/LeonieZ/ACCEPT) [16]. In addition, we compared the CTC results to data derived from lymphocytes and well-known cancer cell lines.

2. Materials and methods

2.1. Data collection/CellSearch®

CellSearch® CTC enumeration is part of ongoing clinical research of the medical oncology department at the Erasmus Medical Center, Rotterdam, the Netherlands. Here, we retrospectively studied the images of CellSearch® cartridges of BC, PC, CRC, and BLC patients. All patients provided written informed consent and were included in clinical trials designed in accordance with the Helsinki Declaration and approved by the local ethics board of the Erasmus MC University Medical Center (Table S1). Subject numbers, cartridge numbers, and CellSearch® appointed CTC counts were collected for all patients. To acquire images of cultured tumor cells, cartridges of experiments in which three BC cell lines (MCF-7, SKBR3, and TD47D) and one PC cell line (LNCAP) were spiked into tubes of blood of healthy donors were studied. Leukocyte analysis was enabled through the selection CD45+/DAPI+/CK− events on one melanoma patient-derived cartridge. CellSearch® generated data consisting of a maximum of 175 Tagged Image Format files (.TIF) per sample, an Extensible Markup Language (.xml) containing the coordinates of the manually marked CTCs, were acquired per cartridge. The images were annotated by tumor type, patient, and material of origin (blood vs liquor). Data collection took place between January 2017 and January 2019.

2.2. Data processing/ACCEPT and SPSS STATISTICS 25

ACCEPT is an open source program developed at the University of Twente, which enables the re-analysis of images of CTCs acquired through the CellSearch® method. [16]. Here, collected CellSearch® images were re-analyzed using the ‘marker characterization mode’ of ACCEPT. This software feature enables the analysis of only those events which were originally defined as CTCs by an expert user of the CellSearch® platform. ACCEPT used the coordinates of marked events present in the CellSearch® generated .xml file to trace back CTCs and selected spiked tumor cells and leukocytes. The program automatically detects the immunofluorescent signals present in the DAPI, CK, and CD45 channel and marks the borders of these events. Per individual event, ACCEPT reports multiple parameters of roundness, signal intensity, and size for the DAPI, CK, and CD45 channel per cartridge in an extended excel file (Microsoft Excel v.10, 365 Office; Microsoft,
analyses were performed using the SPSS file containing only those CTCs with a single nuclear signal. The measurement of the DAPI signal was assessed to select events, accepting enumeration results were compared to the known CellSearch® CT side CTCs. First, as ACCEPT did not detect additional criteria to analyze the cytoplasm/nucleus ratio signal.
The size analysis. To enable the analysis of leukocytes, only single, CK positive events were included in enable the selection of single CTCs for further analyses of CTCs, doublets, and small and large clusters to the CK channel was used to discriminate between single positive CTCs. Therefore secondly, a prior described method based on the ’perimeter to area’ (P2A) ratio of the CK channel was used to discriminate between single CTCs, doublets, and small and large clusters to confirm the size of CTCs per tumor type, it was important to analyze only those events containing confirmed single CTCs. Therefore, events which entered the size analysis had to meet two selection criteria to correct for any inaccuracies introduced by the ACCEPT processing. First, ACCEPT reported events had to have a positive CK signal. Importantly, the CellSearch® marked CTC coordinates in the .xml file could contain images of multiple CK positive CTCs. Therefore secondly, a prior described method based on the ‘perimeter to area’ (P2A) ratio of the CK channel was used to discriminate between single CTCs, doublets, and small and large clusters to enable the selection of single CTCs for further analyses [17]. Only single, CK positive events were included in the size analysis. To enable the analysis of leukocytes, the CK selection criteria were applied to the CD45 signal.

Single CK-positive events had to meet three additional criteria to analyze the cytoplasm/nucleus ratio (NCR) from each CTC. First, as ACCEPT did not detect a DAPI signal in all single CK positive CTCs, the DAPI signal had to be positive. This was defined as the DAPI size being > 0 µm². Secondly, the P2A measurement of the DAPI signal was assessed to select only those CTCs with a single nuclear signal. The third criterion to enter the analysis was that the nucleus had to be intact. This was defined by the fact that the DAPI signal (µm²) had to be smaller than the CK signal (µm²).

2.5. CK and DAPI size determination and cytoplasm/nucleus assessment

The size of the CK and DAPI signals in square micrometers (µm²) as reported by ACCEPT was used to compute an approximate diameter of the cytoplasm and nucleus per CTC. The median size of the predefined CK-positive single CTCs was calculated per tumor type. To assess whether a difference in CK and or DAPI size existed between the four tumor types, the Kruskal–Wallis test (KW test) was used. Subsequently, when a significant size difference existed, the Mann–Whitney U test (MWU test) was applied pairwise to assess the size differences between the tumor types. Within the BC CTCs, the MWU test was also used to assess the size difference between blood- and liquor-derived CTCs. Additionally, for BC and PC, the size difference between patient-derived CTCs and tumor cell line cells was assessed using the MWU test. Finally, this test was used to compare the size of leukocytes to the size of blood-derived CTCs of all tumor types. The computed DAPI and CK diameters were used to determine the NCR in CK positive single CTCs, which had an intact, single positive DAPI signal. The nuclear size and NCR differences between tumor types were assessed through the same nonparametrical testing as mentioned above.

3. Results

3.1. CellSearch® cartridges – Input data description

A total of 1962 CellSearch® cartridges from 1419 individuals were included. The images of 970 BC (655 individuals), 275 PC (185 individuals), 416 CRC (280 individuals), and 301 BLC cartridges (299 individuals) were re-analyzed (Table 1, Figs S1 and S2). Within the BC cohort, 43 of the 970 cartridges concerned liquor-derived images. According to the original CellSearch® data, 66.3%, 84.0%, 43.9%, and 25.2% of the BC, PC, CRC, and BLC cartridges contained CTCs, respectively. From the liquor-derived BC cartridges, 34.9% were CellSearch® CTC positive. Assessment of the original CellSearch® CTC counts showed that for BC, a total of 48 242 blood-derived (CTC-B) and 3285 CTCs derived from cerebrospinal fluid [CTC-L(liquor)]
CTCs were included. In PC, this was 20,442, whereas in CRC and BLC the input CTC count was significantly lower; 626 and 260, respectively (Table 1). Median CTC counts per cartridge were 2 [Interquartile range (IQR) 0–11], 11 (IQR 2–54), 0 (IQR 0–1), and 0 (IQR 0–1) for BC, PC, CRC, and BLC, respectively. The maximum CTC counts were 3254 for BC, 7333 for PC, 51 for CRC, and 32 for BLC samples.

All reference cell line cells were spiked in healthy donor blood and processed by the CellSearch® method. Regarding the BC cell line data, three cartridges containing SKBR3 cells, two cartridges with MCF-7 and one TD47D cells resulted in 530, 1616, and 1249 evaluable events, respectively. One PC cartridge resulted in 1054 LNCAP cells. Selection of leukocytes in one cartridge of a melanoma patient resulted in 130 evaluable events (Table S2).

3.2. ACCEPT output—data description and optimization cohort for CK and DAPI size analysis

ACCEPT detected a total of 72,840 events distributed over the four tumor types. In BC, the number of detected events in blood-derived cartridges was 45,695 and 5327 in liquor-derived cartridges. As summarized in Fig. S3 and Table S3, 99.5% of the events detected in blood and 99.9% of the events detected in liquor were CK positive, while selecting for both CK and DAPI positivity resulted in 86.2% and 99.1% of the events in blood and liquor, respectively. In PC, 20,624 events were detected of which 97.5% was CK positive and 89.3% was CK and DAPI positive. As seen in Fig. S4, the analysis of CRC cartridges resulted in the detection of 934 events of which 55.6% was CK positive and 47.32% was CK and DAPI positive. Finally, 260 events were detected in the BLC samples, of which 76.2% was CK positive and 72.7% was CK and DAPI positive.

Singularity determination of the CK-positive events resulted in 44,232 single blood-derived BC CTCs (97.3%), 1172 doublets, 68 small clusters, and 1 large cluster. In the re-analyzed cartridges containing liquor-derived BC samples, 4702 CTCs (88.4%) were single, while 513 doublets, 104 small clusters, and 3 large clusters occurred. In PC, 19,117 single CTCs (95.1%) as well as 916 doublets, 63 small, and 4 large clusters were found. Within CRC, this resulted in 503 single CTCs (96.9%), 12 doublets and, 3 small and 1 large cluster. Finally, in BLC, 182 of the CK positive events (91.9%) were single CTCs, while 15 doublets and 1 small cluster occurred (Table 2, Fig. 1).

Assessment of the 48,934 positive single CTCs in BC samples resulted in a total of 38,918 DAPI evaluable events (DAPI+ and DAPI signal < CK signal). Of these events, 38,211 showed a single DAPI signal, 682 showed a double signal and 25 showed a clustered signal. Within the 19,117 single CTCs in the PC samples, 13,356 were DAPI evaluable of which 13,272 were single, 224 double, and 5 were clustered DAPI signals, respectively. Out of the 503 single CRC CTCs, a total of 255 events were DAPI evaluable of which 252 had a single DAPI signal and 3 events had a double signal. Finally, from the 182 single BLC CTCs, the DAPI signal was evaluable in 99 events, resulting in 92 times a

Table 1. Description of CellSearch® input data.

| Material | Number of cartridges (CTC+) | Number of patients (CTC+) | Number of CTCs |
|----------|----------------------------|---------------------------|----------------|
| Breast   |                            |                           |                |
| Total    | 970 (643)                  | 665 (497)                 | 51,527         |
| Blood    | 927 (628)                  | 655 (497)                 | 48,242         |
| Liquor   | 43 (15)                    | 35 (11)*                  | 3285           |
| Prostate |                            |                           |                |
| Blood    | 275 (231)                  | 185 (168)                 | 29,738         |
| Colorectal |                        |                           |                |
| Blood    | 416 (183)                  | 280 (136)                 | 626            |
| Bladder  | Blood 301 (76)             | 299 (76)                  | 260            |

* Patients also represented in the ‘breast-blood sample cohort’, therefore not unique.

Table 2. CTC counts per tumor type and type of event. Singularity determination of CK+ ACCEPT events through the use of the CK P2A results.

| Material | Single CTC | Doublet | Small cluster | Large cluster | Total |
|----------|------------|---------|---------------|---------------|-------|
| Breast   | Blood      | 44,232  | 1172          | 68            | 1     | 45,472|
| Prostate | Blood      | 19,117  | 916           | 63            | 4     | 20,100|
| Colorectal |          | 503     | 12            | 3             | 1     | 519   |
| Bladder  | Blood      | 182     | 15            | 1             | 0     | 198   |
| Total    |            | 68,736  | 2628          | 239           | 9     | 71,612|

Molecular Oncology 15 (2021) 116–125 © 2020 The Authors. Published by FEBS Press and John Wiley & Sons Ltd.
single signal, 4 times double, and 2 times a clustered signal (Table S4).

3.3. ACCEPT output—Analysis of size and nucleus/cytoplasm ratio

Figure 2 shows a median size of blood-derived BC CTCs of 120.4 μm² (IQR 104.9) [computed median diameter (CD): 12.4 μm]. Liquor-derived BC CTCs had a median size of 141.3 μm² (IQR 113.1, CD 13.4). The median size of cultured BC cells was 265.4 μm² (IQR 127.38, CD 18.4 μm). In increasing order of size, the median size of the three BC cell lines was 241.7 μm² (MCF-7, IQR 106.5, CD 17.54 μm), (T47D, IQR 95.25, CD 18.35 μm), and (SKBR3, IQR 169.58, CD 22.01 μm) (See Figs S5 and S6 for specific cell line details). PC CTCs had a median size of 83.6 μm² (IQR 63.5, CD 10.3 μm), while the median size of cultured PC cells was 337.9 (LNCAP IQR 142.5, CD 20.7 μm). The CTCs in CRC samples had a median
size of 44.6 \(\mu m^2\) (IQR 53.3, CD 7.5 \(\mu m\)), and the CTCs in the BLC samples had a median size of 57.8 \(\mu m^2\) (IQR 111.5, CD 8.6 \(\mu m\)). Finally, analysis of the lymphocyte data resulted in a median size of 69.6 \(\mu m^2\) (IQR 25.6, CD 9.4 \(\mu m\)). When nonparametrical testing was applied (Fig. 3), the independent
samples median test \((P < 0.001)\) and the KW test \((P < 0.001)\) showed an existing size difference within the patient-derived CTCs of the different tumor types. When testing the size differences between two tumor types at a time, the MWU test resulted in \(P < 0.001\) between every tumor type. MWU testing also showed that the median size difference of 20.9 \(\mu m\) between blood-derived and liquor-derived BC CTCs was significant \((P < 0.001)\). Additionally, the median size differences between cell line cells and patient-derived CTCs of 145 \(\mu m\) in BC and 254.3 \(\mu m\) in PC were both significant \((P < 0.001)\). Finally, the MWU test was applied to assess statistical significance between the median size differences between leukocytes and BC \((-50.8 \mu m, P < 0.001)\), PC \((-14.0 \mu m, P < 0.001)\), CRC \((+25 \mu m, P < 0.001)\), and BLC \((-11.8 \mu m, P = 0.31)\).

Figure 4 shows a median size of 56.1 \(\mu m^2\) (IQR 41.8) for the nucleus of the blood-derived BC CTCs resulting in a NCR of 0.47. The median size of BC cell line cells was 94.2 \(\mu m^2\) (IQR 50, NCR 0.37). Both nucleus size and NCR were statistically significant between patient-derived CTCs and BC cell line cells \((P < 0.001)\). In liquor-derived BC CTCs, the median size of the nucleus was 64.3 \(\mu m^2\) (IQR 36.9, NCR = 0.46). For PC CTCs, the median size of the nucleus was 52.0 \(\mu m^2\) (IQR 35.2, NCR = 0.62), while PC cell line cells had a median nucleus size of 161.0 \(\mu m^2\) (IQR 2.85, NCR = 0.48). Again, both the nucleus size difference as well as the NCR were statistically different between CTCs and cultured cells (MWU \(P < 0.001)\). The nucleus size of CRC CTCs was 26.83 \(\mu m^2\) (IQR 29.0, NCR = 0.60), and in BLC 43.4 \(\mu m^2\) (IQR 33.6, NCR = 0.75). Nonparametrical testing between the different tumor types was applied as stated above; the independent samples median test resulted in \(P < 0.001\) and the KW test in \(P < 0.001\). When testing the nuclear size differences between two tumor types at a time, the MWU test resulted in \(P < 0.001\) between every tumor type except for the PC vs BLC analysis \((P = 0.073)\).

4. Discussion

In this study, we assessed the median size of CTCs in a large cohort of BC, PC, CRC, and BLC patients using \(\text{ACCEPT}\) hypothesizing that there might be a difference in CTCs across tumor types and between patient-derived CTCs and cell line cells, implying the need for tumor-specific CTC isolation methods. We indeed found different median CTC sizes between the four tumor types with BC CTCs being the largest and CRC CTCs the smallest, respectively. Furthermore, we found that liquor-derived BC CTCs are larger than blood-derived CTCs in the same population of patients, suggesting a morphological difference between cells derived of the two types of origin. Finally, the median size of the nucleus also differed between the different tumor types although the variance was smaller than within the distribution of the CK signal.

Two prevailing dogmas regarding CTC size, which are highly important with respect to the future development of size-based isolation methods, are addressed.
by the analysis of our patient-derived CTC data in comparison with our reference data. The first widely accepted hypothesis states that CTCs are generally larger than white blood cells. Literature describes a median size of lymphocytes of 7.1–10.5 μm, and 8.7–9.9 μm for granulocytes [18], which implies that the CTC size as computed in this study is much more similar to the known size of white blood cells than postulated. Our data supports this hypothesis as the leukocytes we analyzed did not statistically differ from BLC CTCs and were even significantly larger than CRC CTCs. Secondly, the development of size-based isolation methods heavily relies on spike experiments using cell line tumor cells, based on the assumption that their median size is comparable to that of CTCs. Our data show that patient-derived CTCs are generally smaller than the cultured cell line cells of the respective tumor type as described in the literature. In BC, the CD of patient-derived CTCs was 12.4 μm, compared to the larger, cultured cells which were 15–17 μm (SKBR-3), and 16.5 μm (MCF-7). In prostate cancer, the median CD of 10.3 μm is also significantly smaller than the size of a cultured cells: 18–21 μm (PC3-9). This is also the case in colorectal cancer, where our computed median diameter is 7.5 μm vs 11 μm in cultured cells (SW480) [19]. Our analysis of three BC cell lines (median size of 18.4 μm) and one PC cell line (20.7 μm) were comparable to these data. With the results of this study, we show that the median size of cell line cells is indeed significantly larger than patient-derived CTCs in both BC and PC.

When the results of our analysis were compared to the expected input data retrieved from the CellSearch® data files, the efficiency of ACCEPT to analyze previously CellSearch® marked events was high. However, the increased number of ACCEPT detected events compared to the CellSearch® marked cell count in CRC was striking. A possible explanation is that cartridges of the specific clinical trial in which this occurred contained the images of the Buffy coat of 30 mL blood instead of the usual 7.5 mL whole blood. Within this technique, 30 mL of blood is pooled into 2 x 15 mL tubes, after which plasma separation and the creation of a Buffy coat is achieved through a centrifuging step. After pipetting off the blood plasma, the two Buffy coats are pooled into one CellSave tube. After these preparation steps, the CellSearch technique is applied in the same manner as on 7.5 mL blood. It is possible that the high background of leukocytes in these samples led to a more difficult distinction for ACCEPT to recognize CTCs as a single event.

The main limitations of this study concern the introduction of a selection bias in the analyzed data. First, its retrospective nature led to the inclusion of a cohort of patients with a wide range of disease characteristics. The cartridge data of all patients included in one of the historical clinical trials were re-analyzed, irrespective of the disease stage, metastatic status, and the treatment line or choice. This could lead to an under- or overestimation of the CTC counts within the current cohort. Second, as we made use of images acquired through the CellSearch® method, the analysis is limited to EpCAM positive CTCs. Although this type of CTCs is of prognostic value in all studied tumor types, it is increasingly suggested that cells which have undergone epithelial-mesenchymal transition (EMT) have a higher metastatic potential and are therefore of interest regarding their characterization. These EMT cells could not be studied through the current method. The third limitation regards the marker characterization mode of ACCEPT. We chose to analyze only those events which were previously marked as CTCs and therefore had to meet the CellSearch® criteria of CK positivity. CK signal ≥ 4 μm², DAPI signal which overlaps the CK signal for ≥ 50%, and a morphological appearance of an intact cell. It is possible that the studied CTC cohort reflects a certain population of cells due to this selection.

5. Conclusion

Although the re-analyzed data in this study were subject to a certain degree of selection bias, to the best of our knowledge it does reflect the largest cohort of morphological studied CellSearch® depicted CTCs which are directly compared to reference data. In addition, one could suggest that the size difference as found in this cohort, irrespective of a possible selection bias, reflects an even more pronounced difference when corrected for this limitation. The shown differences in CTC size between tumor types, and between CTCs and our reference data of cultured tumor cells and patient-derived leukocytes is striking. In conclusion, we therefore suggest that the size of CTCs does matter and should be kept in mind when designing and optimizing size-based isolation methods.

Conflict of interest

The authors declare no conflict of interest.

Author contributions

PAJM, JWMM, LWMMT, and SS designed the project. PAJM, JK, and MV acquired the data. LLZ and LWMMT advised in the use of ACCEPT. EOH
References

1. Allard WJ, Matera J, Miller MC, Repollet M, Connelly MC, Rao C, Tibbe AG, Uhr JW & Terstappen LW (2004) Tumor cells circulate in the peripheral blood of all major carcinomas but not in healthy subjects or patients with nonmalignant diseases. Clin Cancer Res 10, 6897–6904.

2. Riethdorf S, Fritsche H, Muller V, Rau T, Schindlbeck C, Rack B, Janni W, Coith C, Beck K, Pantel K et al. (2007) Detection of circulating tumor cells in peripheral blood of patients with metastatic breast cancer: a validation study of the cell search system. Clin Cancer Res 13, 920–928.

3. Cristofanilli M, Budd TG, Ellis MJ, Stopeck A, Matera J, Miller CM, Reuben JM, Doyle GV, Allard JW, Terstappen L et al. (2004) Circulating tumor cells, disease progression, and survival in metastatic breast cancer. N Engl J Med 351, 781–791.

4. Cohen SJ, Punt CJA, Iannotti N, Saidman BH, Sabbath KD, Gabrail NY, Picus J, Morse M, Mitchell E, Meropol NJ et al. (2008) Relationship of circulating tumor cells to tumor response, progression-free survival, and overall survival in patients with metastatic colorectal cancer. J Clin Oncol 26, 3213–3221.

5. de Bono JS, Scher HI, Montgomery RB, Parker C, Miller MC, Tissing H, Doyle GV, Terstappen L, Pienta KJ & Raghavan D (2008) Circulating tumor cells predict survival benefit from treatment in metastatic castration-resistant prostate cancer. Clin Cancer Res 14, 6302–6309.

6. Alix-Panabières C, Mader S & Pantel K (2017) Epithelial-mesenchymal plasticity in circulating tumor cells. J Mol Med 95, 133–142.

7. Nieto AM, Huang R, Jackson RA & Thiery J (2016) EMT. Cell 166, 21–45.

8. Pantel K & Alix-Panabières C (2019) Liquid biopsy and minimal residual disease – latest advances and implications for cure. Nat Rev Clin Oncol 16, 409–424.

9. Vona G, Sabile A, Louha M, Sitruk V, Romana S, Schütze K, Capron F, Franco D, Pazzagli M, Paterlini-Brechet P et al. (2000) Isolation by size of epithelial tumor cells: a new method for the immunomorphological and molecular characterization of circulating tumor cells. Am J Pathol 156, 57–63.

10. Harouaka RA, Nisic M & Zheng S-Y (2013) Circulating tumor cell enrichment based on physical properties. J Lab Autom 18, 455–468.

11. Gabriel MT, Calleja LR, Chalopin A, Ory B & Heymann D (2016) Circulating tumor cells: a review of non-EpCAM-based approaches for cell enrichment and isolation. Clin Chem 62, 571–581.

12. Valihrach L, Androvic P & Kubista M (2018) Platforms for single-cell collection and analysis. Int J Mol Sci 19, 807.

13. Yu M, Stott S, Toner M, Maheswaran S & Haber DA (2011) Circulating tumor cells: approaches to isolation and characterization. J Cell Biol 192, 373–382.

14. Hao S-JJ, Wan Y, Xia Y-QQ, Zou X & Zheng S-Y (2018) Size-based separation methods of circulating tumor cells. Adv Drug Deliv Rev 125, 3–20.

15. Park S, Ang RR, Duffy SP, Bazov J, Chi KN, Black PC & Ma H (2014) Morphological differences between circulating tumor cells from prostate cancer patients and cultured prostate cancer cells. PLoS One 9, e85264.

16. Zeune LL (2019) Automated CTC classification, enumeration and phenotyping: where math meets biology [MAppS thesis], 207 p. University of Twente, Enschede, The Netherlands.

17. de Wit S, Zeune LL, Hiltermann JTN, Groen HJM, van Dalum G & Terstappen L (2018) Classification of cells in CTC-enriched samples by advanced image analysis. Cancers 10, 377.

18. Shapiro HM, Schildkraut ER, Curbelo R, Laird CW, Turner B & Hirschfeld T (1976) Combined blood cell counting and classification with fluorochrome stains and flow instrumentation. J Histochem Cytochem 24, 396–401.

19. Coumans FAW, van Dalum G, Beck M & Terstappen L (2013) Filter characteristics influencing circulating tumor cell enrichment from whole blood. PLoS One 8, e61770.

Supporting information

Additional supporting information may be found online in the Supporting Information section at the end of the article.

Fig. S1. Number of CellSearch cartridges per tumor type.

Fig. S2. Number of patients per tumor type.

Fig. S3. ACCEPT efficiency vs. CellSearch and ACCEPT results in breast and prostate cancer.

Fig. S4. ACCEPT efficiency vs. CellSearch and ACCEPT results in colorectal and bladder cancer.

Fig. S5. Cell diameter (µm) and size (µm^2) per cell line.

Fig. S6. Nucleus diameter (µm) and size (µm^2) per cell line.

Table S1. Trial details of included cartridges.
Table S2. ACCEPT results reference data.
Table S3. Description of ACCEPT events in relation to CellSearch data.

Table S4. Nucleus singularity determination (in CK+, single CTC, DAPI+, DAPI<CK ACCEPT events).