Toward a real liquid biopsy in metastatic breast and prostate cancer: Diagnostic LeukApheresis increases CTC yields in a European prospective multicenter study (CTCTrap)

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Frequently, the number of circulating tumor cells (CTC) isolated in 7.5 mL of blood is too small to reliably determine tumor heterogeneity and to be representative as a “liquid biopsy”. In the EU FP7 program CTCTrap, we aimed to validate and optimize the recently introduced Diagnostic LeukApheresis (DLA) to screen liters of blood. Here we present the results obtained from 34 metastatic cancer patients subjected to DLA in the participating institutions. About 7.5 mL blood processed with CellSearch® was used as “gold standard” reference. DLAs were obtained from 22 metastatic prostate and 12 metastatic breast cancer patients at four different institutions without any noticeable side effects. DLA samples were prepared and processed with different analysis techniques. Processing DLA using CellSearch resulted in a 0–32 fold increase in CTC yield compared to processing 7.5 mL blood. Filtration of DLA through 5 μm pores microsieves was accompanied by large CTC losses. Leukocyte depletion of 18 mL followed by CellSearch yielded an increase of the number of CTC but a relative decrease in yield (37%) versus CellSearch DLA. In four out of seven patients with 0 CTC detected in 7.5 mL of blood, CTC were detected in DLA (range 1–4 CTC). The CTC obtained through DLA enables molecular characterization of the tumor. CTC enrichment niches however still need to be improved to isolate all the CTC present in the DLA.

Key words: circulating tumor cells, liquid biopsy, CellSearch, diagnostic leukapheresis, filtration

Abbreviations: CTC: circulating tumor cells; DLA: diagnostic leukapheresis; MNC: mononuclear cells; RT: room temperature; SOP: standard operating procedure; WBC: white blood cells

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INTRODUCTION
The presence of circulating tumor cells (CTC) in blood is associated with poor prognosis in patients with metastatic and non-metastatic disease.\textsuperscript{1–7} The numbers of CTC isolated in 7.5 mL of blood are frequently too low to obtain a “liquid biopsy” representative for the tumor, determine tumor heterogeneity and assess the optimal treatment strategy. Extrapolation of the CTC frequency distribution in 7.5 mL of blood from patients with metastatic breast, colon and prostate cancer showed that, probably, all these patients had tumor cells in circulation, but the sample volume was not sufficient to detect them in all patients.\textsuperscript{8} In recent years, numerous assays to detect CTC have been described. EpCAM-based technologies have offered requisites of robustness, reproducibility and cost effectiveness, providing the first in vitro diagnostic CTC assay. However, one of the major drawbacks in the use of CTC for the selection of personalized therapies in individual patients is that CTC are rare events. It was calculated that at a level of 1000 in vivo CTC, there is a probability of 95% that at least one CTC will be detected in 1 out of 5 samples of 7.5 mL of blood with the current technologies; below this tumor burden the accuracy of the detection at a time point is limited by the blood volume that can be obtained from a patient.\textsuperscript{9,10} One solution to overcome this problem is the use of leukapheresis to obtain the mononuclear cell (MNC) fraction believed to contain the majority of CTC from liters of blood. This procedure introduced by Fisher et al.\textsuperscript{11} was baptized diagnostic leukapheresis (DLA). The concept and feasibility of DLA was demonstrated by processing a small aliquot (~5%) of the DLA using the CellSearch® system, which represents the current gold standard for CTC detection. Our first aim was to validate the use of DLA for isolation of CTC in metastatic breast and prostate cancer within the EU FP7 program CTCTrap at multiple centers. Then, we evaluated different technologies to increase the percentage of the DLA that could be processed for the isolation of CTC.

PATIENTS AND METHODS

Patients
Twelve patients with metastatic breast cancer were enrolled in the study at the University Hospital of Duesseldorf and 22 patients with castration resistant prostate cancer; two at the University hospital of Padova, seven at the Institute Gustave Roussy and 13 at the Royal Marsden hospital. All patients provided written informed consent and the study was approved by the institutional review boards at each participating center.

Diagnostic leukapheresis procedure
Leukapheresis were performed at the clinics in Padua, Villejuif, Duesseldorf and Sutton using the Spectra Optia\textsuperscript{®} (Terumo BCT Inc., Lakewood, CO) according to manufacturer’s instructions. For DLA, the program of the apheresis device was set to the MNC collection procedure and set at a collection flow rate of 1.0 mL/min. Concurrent plasma collection was set to a volume of 0 mL. The objective was to collect a minimum volume of 40 mL DLA that was reached in approximately 90 min.

Post DLA sample handling
Samples were divided into aliquots immediately after the DLA procedure, under sterile conditions. White blood cell (WBC) counts and MNC counts were determined using an automated flow-cytometric based hematology analyzer. For CellSearch\textsuperscript{®} analysis an aliquot of the DLA product containing 2 × 10\textsuperscript{6} WBC was diluted to a final volume of 8 mL with CellSearch Circulating Tumor Cell Kit Dilution Buffer (Menarini Silicon Biosystems, Huntington Valley, PA) stored at room temperature (RT) and transferred into a CellSave\textsuperscript{®} tube containing CellSave preservative reagent (Menarini). For direct filtration 50 × 10\textsuperscript{8} WBCs were diluted in 7.5 mL dilution buffer and then transferred into a CellSave tube. For RosetteSep\textsuperscript{™} (Stemcell Technologies, Vancouver, Canada) 18 mL of DLA product was transferred to a 50 mL tube and CellSave preservative reagent from two CellSave tubes was added to the tube. All tubes were kept at RT, at least overnight, until analysis.

CellSearch sample processing
CellSave whole blood was run with CellSearch using the CTC kit (Menarini) according to manufacturer’s instructions. For the DLA product, containing 2 × 10\textsuperscript{8} WBC, the sample was processed using the CellTracks Autoprep system using the CTC kit. The cartridge from the DLA product was scanned, using the CellTracks analyzer II.

Filtration
For filtration, 50 × 10\textsuperscript{6} WBC diluted in CellSearch dilution buffer was filtered using a pump and filtration unit including a microsieve with 5 μm pores (VyCAP B.V., Deventer, The Netherlands). The sample was loaded onto the microsieve and a −100 mbar pressure was applied. Filtration was continued until the entire sample passed the microsieve, or for a duration of maximum 10 min. The successfully filtered sample volume was recorded and used to calculate recoveries. After filtration, the
microsieve was removed from the filtration unit and was washed once with a PBS/1% BSA/0.15% Saponin solution. A permeabilization buffer, containing PBS/1% BSA/0.15% Saponin, was added onto the microsieve and was incubated for 15 min at room temperature. Subsequently, a staining solution was added containing anti-CD45-PerCP (Life Technologies, MHC4531, clone HI30) at a final concentration of 4 μg/mL and anti-CKpan-NanoParticles 575 (AcZon, clone C11 and AE1/AE3) at a final concentration of 3.5 μg/mL in PBS/1% BSA/0.05% Saponin. Staining was performed for 15 min at 37 °C. After removal of the staining cocktail, the microsieve was washed twice using PBS/BSA 1%. The sample was then fixed using PBS/formaldehyde 1% (Sigma, St. Louis, MO) for 10 min at room temperature. Removal of the fluid during each of the staining and washing steps was done using a staining holder including a disposable sponge (VyCAP B.V.). Finally, the sieve was mounted using ProLong® Diamond Antifade Mountant with DAPI (ThermoFisher, Carlsbad, CA, P36971) and a custom cut coverslip (2 × 0.85 cm² custom cut, thickness #1 0.13–0.16 mm) was added to the filter (Menzel-Gläser, Braunschweig, Germany).

Leukocyte depletion
Eighteen milliliters of DLA was depleted from unwanted white blood cell populations using the RosetteSep CTC Enrichment Cocktail Containing Anti-CD36 (Stemcell Technologies, Catalog# 15167). First erythrocytes were isolated by centrifugation of two 9 mL EDTA blood tubes from each patient at 800xg for 10 min. The plasma and buffy coat were then removed and the erythrocytes from both tubes were pooled. Erythrocytes were then added to the DLA product to reach a final WBC to erythrocyte ratio of 1:40. Fifty microliters of the RosetteSep cocktail was then added for each 1 mL of sample and incubated for 20 min at RT. After incubation, the sample was diluted with an equal volume of PBS/2% FBS. The solution was then carefully layered on top of a Ficoll-Paque PLUS density gradient (GE Healthcare, Chalfont St. Giles, UK) and centrifuged at 1200xg for 20 min at RT without brake. The enriched cells where then collected and washed by adding two volumes of PBS/2% FBS and centrifuging for 8 min at 300xg. For filtration, cells were resuspended in 9 mL of 1× PBS and filtered through a 5 μm microsieve (VyCAP BV) at ~100 mbar. Staining of the microsieves was performed as described above. For CellSearch analysis after leukocyte depletion 9 mL of sample was transferred to a conical tube and CellSearch dilution buffer was added to a final volume of 14 mL. This sample was then processed as control within the CellTracks Autoprep system using the CTC kit.

Scanning
All CellSearch cartridges, with enriched CTC, were scanned using the CellTracks Analyzer II (Menarini). All microsieves were scanned using an automated fluorescence microscope available at each site. Each system should match the minimal requirements of having 10 times objective with a minimal numerical aperture of 0.45. In addition, the following filters for fluorescence detection were used: DAPI with excitation 377/50 nm, dichroic 409 nm LP, emission 409 nm LP (Spectra Physics Newport, Santa Clara, CA), PE with excitation 543/22 nm, dichroic 562 nm LP, emission 593/40 nm (Spectra Physics Newport, Santa Clara, CA) and PerCP with excitation 435/40 nm, dichroic 510 nm LP, emission 676/29 nm (Spectra Physics Newport, Santa Clara, CA).

Image analysis
CellSearch fluorescence images were analyzed according to manufacturer’s instructions. The fluorescent images from the microsieves were analyzed using the open-source software ICY. Operators were asked to annotate every DAPI+, CK+, CD45– event. In addition, raw images of both cartridges and microsieves were analyzed by the open source imaging program ACCEPT.13–15 The total number of nucleated events was determined to investigate the number of leukocytes present in the background during image analysis. To improve the detection of nucleated events even in images with crowded areas or images with background artifacts, we applied a high-pass Fourier filter to remove the background beforehand. This feature can be used in future versions of ACCEPT to improve image analysis results.

Statistical analysis
Statistical analysis was performed using OriginPro 9.1 (OriginLab Corporation, Northampton, MA) using the paired sample t-test.

RESULTS
CTC in 7.5 mL of blood versus 200 × 10⁶ cells (~5%) of DLA product
DLAs were obtained from 22 metastatic prostate cancer patients and 12 metastatic breast cancer patients at four different European academic medical institutions. Before starting the DLA procedure, 7.5 mL of whole blood was drawn and processed with the “gold standard” reference CellSearch® to obtain the CTC counts from whole blood. DLA samples were processed through the analysis techniques shown in Figure 1 and described in detail in the Standard Operating Procedures (SOP) developed for DLA in the CTCTrap consortium (https://www.utwente.nl/tnw/mcbp/protocolsandtools/).

In short, 200 × 10⁶ MNC of the DLA product (on average 3.7 mL) were analyzed using CellSearch, 50 × 10⁶ cells were analyzed using filtration and 18 mL of the DLA product was depleted of its white blood cells, allowing analysis of a larger part of the DLA product, followed by CTC enumeration by either filtration or CellSearch.

DLAs from metastatic cancer patients were performed for ~90 min without any noticeable side effects. DLA products had an average volume of 53 mL (range 21–98 mL, SD 16 mL) containing an average of 3.3 × 10⁹ (range 1.5 × 10⁷–9.0 × 10⁷; SD 2.0 × 10⁷) MNC representing ~1.6 L (range 0.03–3.5 L, SD 0.7 L) of blood. Figure 2a illustrates the concentration of MNC per mL of whole blood and DLA product.
The average MNC count in whole blood was $2.0 \times 10^6$/mL (range $0.3 \times 10^6$/mL–$4.3 \times 10^6$/mL, SD = $0.9 \times 10^6$/mL) and in DLA product $64.0 \times 10^6$/mL (range $0.4 \times 10^6$/mL–$164.3 \times 10^6$/mL, SD = $36.8 \times 10^6$/mL). In Figure 2b, the ratio of the concentration of MNC in DLA to MNC in blood is shown.

The number of CTC in 7.5 mL of blood ranged from 0 to 324 (mean 67, median 18) and CTC in DLA ranged from 0 to 2913 (mean 362, median 160) resulting in a significant increase in CTC yield ($p = 0.003$). The increase in CTC yield ranged from $0 \times$ to $32 \times$ (mean 6, median 5). The analyzed DLA volume represented 7–212 mL of blood (mean 98, median 97). In Figure 3, the absolute number of CTC in 7.5 mL of peripheral blood and in $200 \times 10^6$ MNC of the DLA product measured by CellSearch is illustrated for each patient. Extrapolation of the number of CTC obtained when the complete DLA volume could have been processed with...
CellSearch for each patient is also shown in the figure. The complete DLA volume represented 7–212 mL of blood (mean 98, median 97). The entire DLA product would comprise 0–9037 CTC (mean 3304, median 2873) or a 0× to 417× increase (mean 104, median 77) compared to 7.5 mL of blood.

**CTC in 2–18 mL (~5–45%) of DLA product**

The identification of CTC within enriched cell suspensions becomes increasingly more difficult when the number of leukocytes are so large that they are in close proximity to each other. In Figure 4a, representative microscopic images are shown after processing blood by CellSearch (Fig. 4a1), after processing DLA with CellSearch (Fig. 4a2), after filtration of DLA (Fig. 4a3) and after depletion of leukocytes in DLA product followed by either filtration (Fig. 4a4) or CellSearch (Fig. 4a5). From the images, it is clear that identification of CTC in DLA product directly processed by CellSearch or filtration is more difficult because of the larger background of leukocytes. To quantify the number of nucleated cells in the images, we used the open source imaging program ACCEPT. Gates were set to find all nucleated events by looking at the mean intensity of the DAPI signal and gates for the perimeter and eccentricity were set to identify cell like

![Figure 3](image-url) Absolute CTC counts in 7.5 mL of blood, 200 x 10⁶ cells of DLA product processed by CellSearch and the CTC count in the total DLA product by extrapolation of the DLA CTC counts, lines connect measurements from the same patient.

![Figure 4](image-url) (a) Typical microscopic images obtained after CTC enrichment and detection with the different techniques. Top left, the average number of nucleated cells in the enriched CTC samples assessed by ACCEPT. (b) CellSearch images from CTC detected in both blood and DLA from four patients. Cells show similar morphological characteristics (nucleus = purple, cytokeratin = green) [Color figure can be viewed at wileyonlinelibrary.com]
The gate used to define nucleated cells was a DNA mean intensity >5, a DNA perimeter >16 and a DNA eccentricity ≤0.95. The average number and standard deviation of nucleated cells present after isolation are shown in the top left part of the image. Blood volumes of 7.5 mL processed with CellSearch had on average 27,513 nucleated events in the background (SD = 6,716; n = 7), DLA products processed with CellSearch had 50,778 nucleated events (SD = 40,486; n = 8), directly filtered DLA exhibited 88,142 nucleated events (SD = 13,338; n = 8), DLAs depleted followed by filtration had 11,929 nucleated events (SD = 3,337; n = 5) and DLAs depleted followed by CellSearch yielded 3702 nucleated events (SD = 1,729; n = 5). The number of CTCs in samples with more than 100,000 WBC in the background is most likely underestimated as they may be obscured by leukocytes. To illustrate the similarities between the morphological characteristics of CTC in blood and DLA, a gallery of images from four patients is shown in Figure 4b.

**Figure 5.** Expected number of CTC based on CellSearch CTC count in 200 × 10^6 cells of DLA product plotted against the actual measured CTC count for each of the analysis techniques. (a) CTC recovery after filtration (n = 16). (b) CTC recovery after depletion of DLA product followed by filtration (n = 22). (c) CTC recovery of depletion of DLA product followed by CellSearch analysis (n = 5).

**DISCUSSION**

In this European multicenter study we showed that in 34 metastatic cancer patients the DLA procedure was well tolerated and...
a DLA product could be obtained. The mononuclear blood fraction obtained represented 0.03–3.5 L of blood (mean = 1.6, SD = 0.7 L). In one patient the represented blood volume was only 30 mL, whereas in all other patients, volumes represented 400 mL or more peripheral blood. Due to the high MNC concentration resulting in high leukocyte-carry-over impeding CTC detection, only around 5% (200 × 10^6 cells) of the DLA product can be directly processed in a CellSearch run. Processing of 200 × 10^6 cells with CellSearch can already result in a carryover that is too high for accurate identification of CTC. A simple solution to this problem is to dilute the sample and divide it over multiple cartridges for scanning and image analysis. However, this approach would lead to a practical problem consisting in the availability of empty cartridges.

For a thorough molecular characterization of the tumor an assessment of heterogeneity is important, thereby implying the need for an assessment of the individual cells. A question that cannot be answered is how many molecular characterized tumor cells from how many tumor sites are needed. However, we do know that the tumor cells present in the blood can come from a variety of metastatic sites, enabling their direct analysis. Extensive heterogeneity has been observed in CTC that match the findings in the metastatic sites to various degrees. The molecular characterization of single CTC requires technology for their isolation, which can be performed by FACS,22,23 DEPArray,4,25 Punch26,27 or micromanipulation.28 However, employing these methods, cell loss is inevitable. Another problem, if only very few cells are available for analysis is that CTCs are frequently in poor condition disabling their further DNA and/or RNA analysis.29 Therefore, there is agreement that the more tumor cells are available the better the chance to accurately characterize them. With DLA, the increase in CTC yield after processing 5% of the DLA product with CellSearch ranged from 0 to 32 fold (mean 6, median 5) compared to analysis of 7.5 mL of matched PB. In 23% of the patients >100 CTC were detected in 7.5 mL of blood and this increased to 53% of patients in 2 mL of DLA and to 68% if all the DLA could have been processed. A lower number may be sufficient for molecular characterization and for example in 59% of the patients >10 CTC were detected in 7.5 mL of blood and this increased to 65% of patients in 2 mL of DLA and to 79% if all the DLA could have been processed. Molecular characterization and expansion of CTC through in vivo culture in mice is being explored in some of these samples.30,31

Clearly, some improvements have to be made before the use of DLA can become practice in a clinical setting. For example, methodologies for molecular characterization of CTCs isolated from DLA need to be standardized. Also, to proceed with molecular characterization, more effective methods need to be established to process the whole DLA product, e.g. processing of 20 CellSearch runs is accompanied by practical and economical limitations. We therefore evaluated other means of CTC enrichment in DLA products. The use of DLA filtration through microsieves was also limited by overwhelming number of leukocytes, which restricted the use of only ~1.25% (50 × 10^6 cells) of the DLA product. Depletion of leukocytes prior to filtration or CellSearch enrichment indeed resulted in a number of leukocytes that could be handled by the image recognition of CTC but the depletion procedure itself was accompanied by CTC losses. The overall number of CTC obtained however increased (Fig. 5) thereby raising the ability to characterize the different tumor cells. We conclude that the use of DLA increases the number of CTC that can be isolated. However, further improvements on CTC enrichment technologies are needed to truly gain advantages of DLA as a means to obtain sufficient CTC for the characterization of the tumor and ultimately to guide therapy.

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