Fugitives on the run: circulating tumor cells (CTCs) in metastatic diseases

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Abstract
The presence of circulating tumor cells (CTCs) in the bloodstream signals the existence of a tumor and denotes risk of metastatic spread. CTCs can be isolated and analyzed to monitor cancer progression and therapeutic response. However, CTC isolation devices have shown considerable variation in detection rates, limiting their use as a routine diagnostic and monitoring tool. In this review, we discuss recent advances in CTC detection methodologies and associated clinical studies. We provide perspective on the future direction of CTC isolation and molecular characterization towards developing reliable biomarkers that monitor disease progression or therapeutic response.

Keywords Circulating tumor cells · Metastasis · Liquid biopsy · Biomarker

1 Introduction

The metastatic spread of cancer cells from a primary tumor to distant sites is the leading cause of mortality in cancer patients [1]. The underlying molecular mechanisms of metastasis are not well understood and remain a fundamental topic in the field of cancer research. The isolation and characterization of tumor cells that have broken off from the original tumor site and circulate within the peripheral blood stream can provide insight into the process of metastatic spread and can relay a real-time monitoring of disease progression and therapeutic response. This characterization of circulating tumor cells (CTCs) holds promise for guiding personalized therapies and developing novel anticancer drugs specifically targeting the metastatic process. CTCs exist in extremely low quantities in the blood stream, making this “liquid biopsy” difficult to capture and study. For this reason, there is continued interest in developing robust techniques for CTC isolation. Exploration into unique molecular signatures that can be used for effective enrichment is needed. To fortify the role of “liquid biopsies” and CTCs as a potential predictive biomarker in the clinical setting, focus should be placed on developing more precise and efficient CTC enrichment techniques.

2 Current use of CTCs

2.1 Trends in CTC enrichment techniques

In a liquid biopsy, there is roughly one CTC detectable out of ten million white blood cells per one milliliter of blood [2]. To add complexity, similar to a primary tumor, these circulating cells are known to exhibit heterogeneity with multiple sub-populations expressing different molecular markers [3]. The lack of a single ubiquitous marker or molecular signature present on all cancer cells creates a challenge in developing highly precise CTC isolation techniques. This has led to the invention of isolation devices that focus on exploiting epithelial marker proteins that are expressed on tumor cells and are deficient on the surrounding blood cells. In 2004, the U.S. Food and Drug Administration (FDA) approved an epithelial cell adhesion molecule (EpCAM)-dependent technique for use as a prognostic biomarker in breast cancer [4], which was then expanded for use in prostate and colorectal cancer patients [5]. EpCAM is an attractive target for CTC isolation because it is detectable on the majority of epithelial-derived cancer types and not detectable on leukocytes, making it possible to isolate CTCs effectively from other blood components.
Since its approval, the EpCAM-dependent device has become the “gold standard” for CTC isolation.

Approval of a CTC enrichment device facilitated investigations into the clinical use of CTCs for monitoring disease progression and therapeutic response, resulting in a steady increase in the number of clinical studies measuring CTCs (Fig. 1). Despite this progress, clinical use of the EpCAM-dependent device remains rather limited. Over half of all clinical studies on CTCs since 1999 were conducted in only three cancer types: prostate, breast, and lung cancer (Fig. 2). Although the role of CTCs in other cancer types has been studied, their clinical use is yet to be confirmed due to various limitations in enrichment [8–11].

2.2 Markers for CTC enrichment

Enrichment devices reliant on a specific cell surface marker for isolation can pose a bias by excluding sub-populations of CTCs that are deficient or express low levels of the marker. A key example is seen with EpCAM-dependent enrichment of CTCs; only CTCs that retain epithelial characteristics are isolated, excluding CTCs with mesenchymal traits. Cancer cells can biochemically alter their phenotype, undergoing epithelial to mesenchymal transition (EMT), gaining invasive mesenchymal and stem cell-like properties and losing epithelial characteristics, such as cell surface EpCAM expression. The CTC population can therefore be comprised of cells with a range of epithelial, epithelial-mesenchymal, and mesenchymal characteristics [12]. With EpCAM-dependent enrichment techniques, the marker-negative CTCs are undetectable, creating uncertainty in the accuracy of a patient’s CTC status. Analysis is then limited to cancers with CTCs that predominantly maintain epithelial characteristics.

Recent studies have focused on evaluating the epithelial and mesenchymal characteristics of CTCs to determine how these phenotypic differences can affect their clinical utility [13–16]. One such study assessing the clinical relevance of epithelial, mesenchymal and epithelial-mesenchymal CTCs in colorectal cancer found that only mesenchymal and epithelial-mesenchymal CTCs, not epithelial CTCs, correlated with clinical stage and metastasis [13]. Similarly, mesenchymal CTCs in esophageal squamous cell carcinoma have been shown to denote clinical stage and treatment efficacy [14]. These findings support the importance of isolating mesenchymal CTCs for inclusion in downstream analysis.

Despite the challenge in addressing the totality of CTCs, EpCAM-dependent enrichment is still the most commonly used marker in CTC isolation (Fig. 3). More than eight years of clinical studies have been conducted using this approach, with a steady increase in the number of studies over the past 20 years. These trials include enumeration, characterization, or a combination. Molecular characterization (blue), which encompasses genetic and proteomic characterization of CTCs, shows a steady increase and signals a trend in cancer therapeutic development.
EpCAM-dependent enrichment devices have been developed since 2004 for use in clinical trials (summarized in Table 1); with their use as prognostic tools pending confirmation. Alternative approaches are being developed to attempt collection of a phenotypically representative sample of the entire CTC population (Fig. 3), such as label-free devices that use size differentiation to capture CTCs or methods, such as negative depletion, which use immunomagnetic and microfluidic removal of blood cells leaving only the CTCs for downstream processing (Table 1). Size-based devices take advantage of large tumor cell diameters (> 8 μm) by using a porous filter with a size range to allow small diameter leukocytes and other blood elements to pass through [32]. Such a filtration approach is effective in isolating the diverse population of CTCs; yet, the inefficient depletion of white blood cells often leaves substantial background in the isolate, making downstream analysis difficult. Negative depletion typically involves red blood cell (RBC) lysis and removal of other blood elements by anti-CD45 immunomagnetic separation of leukocytes, allowing label-free CTC isolation. Following enrichment, immunocytochemistry is a common procedure to distinguish CTCs from cells in the peripheral blood stream not properly excluded during the isolation. This imaging technique often involves fixation and permeabilization, potentially interfering with further downstream molecular analysis of CTCs.

2.3 Clinical utility of CTCs

CTC enumeration is currently used as a prognostic marker of progression-free survival (PFS) and overall survival (OS) in breast, colorectal, and prostate cancer [33]. When using CTCs as a surrogate biomarker, a minimum cutoff value is designated to evaluate prognostic outcome. However, this cutoff value varies across isolation devices, the protein marker used, and the origin of the primary tumor. For example, ≥ 5 CTCs in 7.5 mL of blood was determined retrospectively to be a cutoff value to indicate poor prognosis based on OS and PFS in metastatic breast cancer patients when using an EpCAM-dependent device [34]. These findings established this cutoff value as “CTC-positivity,” which is now widely used for prognostication in breast cancer [35]. Similar retrospective determination of CTC-cutoff values were found in clinical trials for metastatic prostate cancer, which found use of ≥ 5 CTCs as a cutoff based on correlation with OS [36] and colorectal cancer (CRC), in which > 3 CTCs was associated with OS and PFS [37]. While correlations between enumeration and survival rates have been determined [38], the optimal
cutoff values vary across trials and is limited to epithelial-marker positive cancers [12, 32, 39–41]. Studies on other cancer types have difficulty defining a prognostic cutoff value [9] showing necessity for more effective CTC isolation and retrospective analyses.

In addition to use as a prognostic biomarker, CTC enumeration is being used as an exploratory indicator of efficacy and a measurement of treatment response [42–44]. A drop-off in CTC counts showed promise as an early efficacy endpoint in a study observing cabazitaxel response in metastatic castration resistant prostate cancer (mCRPC) patients with docetaxel resistance [42]. Enumeration was also tested as a patient stratification biomarker in the SWOG S0500 trial, which used changes in CTC counts in response to therapy as an early indicator of therapy resistance to determine a change in treatment regimen, though this prospective use was ineffective at prolonging OS in metastatic breast cancer patients [45]. To strengthen the use of CTCs as a biomarker, enumeration is being used as a secondary outcome measure to increase the accuracy of other prognostic markers used in clinical practice, such as tumor associated protein markers (i.e., PSA, CEA, CA125) and imaging tests. One study specifically found that using CTC enumeration in conjunction with positron-emission computerized tomography (PET-CT) had greater prognostic significance than either measure on its own [31].

To explore the predictive value of CTCs in the clinical setting, there must be uniformity in isolation and characterization methods. Without uniformity, there is inconsistency in the CTC population analyzed [46]—whether it be phenotypic variability or inconsistent capture efficiency in cancer types—limiting the scope of any significant findings. Clinical validation of markers identified in ongoing retrospective studies [47] will help to clarify how CTCs can be used to direct personalized medicine and drug development.

### 2.4 Molecular characterization

Interest in molecular characterization, specifically genetic and proteomic characterization, has been on the rise as the clinical utility of CTCs for therapy personalization has become more apparent (Fig. 1, Fig. 3). Genetic characterization explores the presence of a specific mutation or activation of an oncogene that can be targeted through therapy or correlated with progression or response. NCT03366116 is a clinical trial in progress exploring the efficacy of the nucleoside analog abbreviated Aza-TdC intended to upregulate tumor suppressor genes, which will be monitored in isolated CTCs [48]. Whole transcriptome sequencing of CTCs can identify specific genetic events that occur during the metastatic process and therapeutic response, holding promise for cancer drug development [49]. Technical challenges within CTC capture efficiency have made progress in this field difficult, limiting the number of genomic studies and significant findings to date [50]. Sequencing methods, like mRNA-Seq, require high capture purity and viable CTCs for downstream analysis, which is difficult to achieve with current isolation devices due to the intensive fixation and labeling methods used [51]. One study was able to identify a CTC gene signature that could predict survival in pancreatic cancer patients using a unique method of negative depletion followed by gene microarray analysis [52].

Detecting the presence of specific proteins on the surface of CTCs is more commonly used and can be done by incorporating additional protein labeling into the isolation-devices’ standard immunochemistry (including CD45, DAPI, cytokeratins, and EpCAM). The ongoing DETECT studies (III, IV, and V) incorporate human epidermal growth factor
| Device Feature for isolation | Technology | Comments | Clinical studies |
|------------------------------|------------|----------|------------------|
| **Positive enrichment** | CellSearch (Veridex) EpCAM | FDA approved, ferromagnetic beads labeled with EpCAM-antibodies, stained for CK⁺, CD45⁻, DAPI⁺ | No detection of EMT, single marker, expensive, no further downstream analysis | [17, 18] |
| GILUPI CellCollector EpCAM | In vivo metal probe with anti-EpCAM antibodies on surface, high sensitivity | Invasive 30-min insertion, no detection of EMT, single marker, no further downstream analysis | NCT03071900, NCT02507778 |
| IsoFlux (Fluxion Biosciences) EpCAM and optional additional markers | Antibody based magnetic separation by flow cytometry, additional markers available | Based on cell surface antigen presence, automated identification, incorporates Next Generation Sequencing | [19] |
| Herringbone CTC-Chip EpCAM | Antibody micropost capture, microfluidic device | Increased interactions with antibody surface, based on cell surface antigen presence, single marker, no detection of EMT | [20] |
| MAC MicroBead Technology (Miltenyi) EpCAM | Superparamagnetic nano-sized beads | No differentiation of low and high expression levels [21], single marker, no detection of EMT | [22] |
| BD FACScanto II EpCAM and CKs | Flow-cytometry antibody fluorescent separation | No detection of EMT, based on cell surface antigen presence | [23, 24] |
| GEDI EpCAM | Microfluidic device with EpCAM-antibody labeled obstacles, stained for CD45⁻, DAPI⁺, Pdx-1⁺ | No detection of EMT, single marker, no further downstream analysis | [25] |
| EPISPOT Many cell surface proteins | CD45 depletion followed by immuno-fluorescence using fluorochrome labeled secondary antibodies | Utilize CTC secreted proteins for enrichment, protein characterization possible | [26] |
| **Negative enrichment** | RosetteSep CTC Enrichment Cocktail/ CD45 Depletion Cocktail (STEMCELL) Density | RBC lysis with or without CD45 depletion followed by density gradient to remove lymphocytes and other unwanted cells | Requires downstream analysis to confirm CTC status | NCT02349867 |
| ApoStream Dielectric property | Dielectrophoresis field-flow that repels blood cells into eluent | Takes advantage of CTC phenotypic and dielectric properties | NCT02349867 |
| MAC MicroBead Technology (Miltenyi) CD45 | Superparamagnetic nano-sized beads deplete CD45⁺ leukocytes | No differentiation of low and high expression levels [21] | NCT02349867 |
| Size-based CellSieve (Creatv Microtech) Size | Biocompatible polymer filter with 7-μm diameter pores | Loss of small diameter CTCs, high leukocyte retention | [27, 28] |
| ISET Size | Polycarbonate membrane with 8-μm diameter pores | Loss of small diameter CTCs, high leukocyte retention | NCT01776385, NCT01943500 |
| Viator Therapeutic Oncophoresis System Size and deformability | Ex vivo extracorporeal fluid circuit, cross-flow filtration, returns other blood elements back to the body | Time-consuming, patient compliance with procedure, loss of small diameter CTCs | NCT02490800, NCT03427450, NCT02785731 |
| Parosotix System (ANGEL) Size and deformability | Microfluidic device, separates cells when flowing through steps of the cassette | | NCT02490800, NCT03427450, NCT02785731 |
| CanPatrol (SurExam) Size | RBC lysis followed by filtration in an 8-μm diameter porous membrane | Loss of small diameter CTCs, high leukocyte retention | [14, 29] |
| ScreenCell CytoFiltration Size | 8-μm diameter porous membrane | | NCT02951897 |
| Density-based Ficoll-Paque Density | Reagent separates peripheral blood mononuclear cells for downstream analysis of CTCs | Requires downstream analysis to confirm CTC status | [31] |

*CK, cytokeratin; DAPI, 4′,6-diamidino-2-phenylindole
receptor 2 (HER2) staining into an EpCAM-dependent protocol to identify HER2-positive CTCs in patients with HER2-negative primary tumors [53]. Patients with HER2-negative breast cancer that are found to have HER2-positive CTCs are allocated to the DETECT III study where efficacy of HER2-targeted therapy is observed as a means of controlling the metastatic process [54]. An alternative study using the ProOne Assay (Prometheus Laboratories) to identify HER2-positive CTCs in HER2-negative metastatic breast cancer patients found that HER2-targeted therapy was not effective in this population due to continued disease progression [55]. This discordance between primary tumor and CTC phenotypes may contribute to recurrence and disease progression. For this reason, biopsy and analysis of the primary tumor must still be used in conjunction with CTC molecular characterization for accurate determination of guided therapies. The intrinsic heterogeneity of protein expression across the CTC population and between the primary and circulating tumor populations makes the use of CTCs as a “liquid biopsy” for real-time monitoring of cancer lesions and targeted treatment response challenging [56].

With CTC heterogeneity posing a challenge to understanding and targeting the metastatic process, identification of a more commonly expressed molecular marker and development of techniques that utilize such a marker is imperative. For biomarker discovery, label-free devices and techniques that leave CTCs relatively “untouched” for downstream analysis, such as size filtration techniques, can be used. Recently, a novel cancer cell-specific surface marker was identified and was shown to increase the efficiency and specificity of CTC isolation from patient blood samples. This marker, a unique proteoglycan modification of chondroitin sulfate (CS) called oncofetal CS (ofCS), was found to be expressed on 95% of patient-derived human cancer cell lines of hematopoietic, epithelial, and mesenchymal origin [57]. CELLection beads and microfluidics designed to bind ofCS were stated to be capable of isolating CTCs with high efficiency [39]. ofCS is present on cancer cell types independent of origin or disease stage, meaning it exists on EpCAM-negative and EMT CTCs as well [39]. Identification of a CTC-specific surface marker, such as ofCS, may eliminate the need for subtype specific markers in CTC isolation, reducing current concerns with establishing consistency in CTC clinical use and significance.

3 Perspectives

Understanding the role CTCs play in the metastatic process can aid the development of anticancer therapies and improve disease evaluation [58]. The current state of CTC isolation strategies underlines the need for more reliable enrichment techniques that can isolate the entire CTC population and be combined with downstream molecular characterization. These techniques may include more ubiquitously expressed markers or molecular signatures on CTCs, like ofCS, to investigate their clinical utility and application in different cancer types. The identification of a more global signature for CTC identification and enumeration can eliminate many of the inconsistencies across methods, improving the isolation and targeting of this metastatic population. Implementation of clinical guidelines for CTC studies has been proposed as a method of improving standardization within trials with respect to experimental design and analysis [59]. Once a guideline for CTC clinical studies has become well established, clinical implementation of CTCs as markers of progression or treatment response can be more clearly defined.

Heterogeneity in CTC phenotype and discordance with primary tumor type also needs to be addressed. The process of EMT highlights how CTC phenotype evolves and can provide valuable insight into the metastatic process, disease progression, and chemotherapy resistance. Molecular characterization holds promise for personalized cancer therapy, not only providing a potential target for therapeutic intervention, but also a reliable and consistent means of isolating the entire CTC population. CTC characterization used prospectively in treatment decision and drug development could reshape cancer therapeutic practices.

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Compliance with ethical standards

Conflict of interest On behalf of all authors, the corresponding author states that there is no conflict of interest.

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