The Novel Association of Circulating Tumor Cells and Circulating Megakaryocytes with Prostate Cancer Prognosis

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Abstract

Purpose: To develop an approach for the investigation of different subtypes of circulating tumor cells (CTC) and other cells to evaluate their potential prognostic value of prostate cancer.

Experimental Design: Malignancy of CTCs undergoing epithelial-to-mesenchymal transition (EMT) was confirmed by repeated FISH. Subgroups of CTCs in 81 patients with prostate cancer (43 castration resistant and 38 untreated localized) were correlated to disease aggressiveness parameters. AUC analysis was applied to compare the performance for metastasis prediction between serum PSA level alone and a combined risk score using both PSA and EMTting CTC count. Circulating megakaryocytes and cancer patient survival association was performed using Cox model.

Results: The majority of vimentin (VIM)/CD45± cells were malignant, with genomic alterations in several genomic regions. The number of cytokeratin (CK)/VIM+/CD45± CTCs correlated with disease burden, tumor aggressiveness, and poorer survival. Meanwhile, CK+/VIM+/CD45± CTCs were associated with metastases better than other subtypes of CTCs in these limited samples. Combination of PSA level and the number of CK+/VIM+/CD45± CTCs enhanced the prediction of cancer metastases [AUC, 0.921; 95% confidence interval (CI), 0.858–0.985]. The number of circulating megakaryocytes was potentially associated with good patient survival in advanced prostate cancer (HR, 0.849; 95% CI, 0.628–1.146, per cell increase), and the difference between the number of mesenchymal CTCs and megakaryocytes strongly correlated to poor survival (HR, 10.17; 95% CI, 2.164–47.789, if score ≥2.0).

Conclusions: This CTC analysis approach and the potential association of megakaryocytes with cancer prognosis may greatly enhance our ability to investigate the cancer metastasis process and to predict/monitor cancer progression.

Clin Cancer Res; 23(17); 5112–22. ©2017 AACR.

Introduction

Cancer cells evolve during disease progression and in response to treatment (1), influencing their sensitivity to treatment. Circulating tumor cells (CTC) are considered to be the seeds of metastases (2) and, at the same time, a source of information on tumor tissue that can be acquired through a simple blood sample. Even before a metastatic tumor is clinically evident, it has been reported that a large number of CTCs already exist in the circulation (3, 4). This provides a more accessible “liquid biopsy” than tumor tissue biopsies to predict/monitor disease progression and therapy response at both cellular and molecular level. However, current CTC analysis mainly focused on cells expressing epithelial phenotype–specific markers (2). In fact, activation of epithelial–mesenchymal transition (EMT) is a key process in cancer metastasis (5) and an important factor in promoting invasiveness of cancer cells and their resistance to therapy (6). Both downregulation of epithelial markers, such as epithelial cell adhesion molecule (EpCAM), cytokeratin (CK), and E-cadherin, and upregulation of mesenchymal markers, such as vimentin (VIM) and N-cadherin, have been described (7, 8). Hence, CTCs undergoing EMT as part of the metastatic process may be missed when isolated on the basis of their epithelial characteristics only. To address this challenge, we have optimized a novel epitope–independent CTC isolation system, Parsortix, to capture CTCs based on the much larger size and less deformability nature of tumor cells compared with normal blood cells (9). The efficiency of Parsortix in capturing CTCs has been independently validated by other research groups in comparison with CellSearch in patients with small-cell lung cancer (10), and other types of cancers, including breast, colon, and lung (11). Here, we explored its clinical application in prostate cancer prognosis and disease state monitoring.

Prostate cancer is the most common cancer in Western men (12, 13). Current criteria for risk stratification of newly diagnosed
prostate cancer are mainly based on clinical features, including serum PSA, clinical stage, and biopsy/surgical specimen Gleason score (GS; ref. 14). However, these factors are not sufficient to discriminate between patients with indolent and aggressive disease. Furthermore, when metastatic disease progresses to castration-resistant prostate cancer (CRPC), the established PSA test is not satisfactory for disease prognosis and monitoring. These current clinical unmet needs urge us to identify novel biomarkers that can monitor disease status precisely, accessibly, and in real time. Therefore, we investigated the potential of analyzing CTCs with both epithelial and mesenchymal features for cancer prognosis using Parsortix system in prostate cancer as a disease model. We developed a novel technique to perform five rounds of fluorescence in situ hybridization (FISH) on the same slides after immunofluorescence staining. Using this technique, we simultaneously identified the epithelial and mesenchymal cell features and multiple genomic alterations, confirming the malignancy of circulating cells with mesenchymal phenotype. We also demonstrated the correlation of CTCs with advanced disease features. Unexpectedly, we discovered that an increase of circulating megakaryocytes had the trend to be correlated with good prognosis in patients with progressive disease and the combination of CTC and megakaryocyte count may effectively predict survival in advanced disease.

**Materials and Methods**

**Patients and blood samples**

A total of 81 patients with written consent were recruited in December 2014 from St Bartholomew’s Hospital, Barts Health NHS (London, United Kingdom), comprising 38 with untreated localized prostate cancer, and 43 with progressive CRPC (40 with metastasis) ready to commence an alternative treatment. The clinical information is summarized in Table 1, and details of sample collection and clinical information are shown in Supplementary Data and Supplementary Table S1. Blood specimens from 24 healthy male donors were collected with signed ethics committee–approved consent forms. Whole blood (7.5 mL) was donated from each participant for CTC enumeration. Use of blood samples from patients and healthy donors in this study was approved by National Research Ethics Service committee London City and East with a Research Ethics Committee reference of 09/H0704/4.5.

**CTC isolation and enumeration**

Isolation of CTCs from whole blood using a size- and deformability-based system Parsortix was performed as described previously (9). Sample harvest and the process of immunostaining are detailed in Supplementary Materials and Methods. Different DAPI+ populations of cells were recorded, including CK+/VIM−/CD45−, CK+/VIM+/CD45+, and CK−/VIM+/CD45−.

**FISH after immunofluorescence analysis**

Immunofluorescence signals were washed in a stripping buffer (containing 2% SDS, 0.0625 mol/L Tris-HCl pH 6.8, and 0.8% β-mercaptoethanol) at 50°C for 20 minutes, followed by washing three times in PBS for 5 minutes. Details of FISH probe and multiple FISH procedure are indicated in Supplementary Materials and Methods. Ten FISH probes for nine commonly altered genes/genomic regions in prostate cancer, including 6q16, NKX3.1, C-MYC, PTEN, CCND1, RB1, 16q22.1, ERG, and AR were used for CTC analysis. The copy number of each FISH probe was counted in CK+/VIM−/CD45−, CK+/VIM+/CD45+, and CK−/VIM+/CD45− cells. The copy number of lymphocytes from the prostate cancer patients and healthy donors was used to calculate the baseline for each genetic change. FISH signals from >100 lymphocytes were counted in each sample.

**Statistical analysis**

Unless specifically noted, Wilcoxon rank sum test was applied to assess the equality of CTCs between subgroups based on CTC score as well as different clinical features, such as metastasis, primary GS, and risk classification in localized disease. Data shown were as median [interquartile range (IQR)]. Spearman rank correlation was used to assess the association between CTC counts and concurrent PSA level. Bivariante logistic regression was performed with PSA and EMTing CTC counts as predictors for imaging-detected metastasis (yes, no). A combined risk score (CRS) was computed as the linear predictor of the ROC curve analysis was used to test the ability of different subtypes of CTCs as well as CRS to distinguish patients with metastasis. Optimal cut-off point was calculated with an optimal-corrected classified value to provide best available sensitivity and specificity. Roogold function was used to independently test the equality of the ROC area of each method against the PSA as a standard curve. The combined mesenchymal CTC and megakaryocyte score (CMS) was calculated as:

\[
\text{Mesenchymal CTC count} - \text{megakaryocyte count} \\
\text{Megakaryocyte count}
\]

Because of the possible absence of megakaryocytes detected in patients, megakaryocyte count = 0 was recorded as 1 for this score calculation. To determine the factors that predict survival time, variables considered as potential predictors were selected for univariate analyses using Cox model. The hazard ratio (HR) associated

**Translational Relevance**

Here, we firstly developed an optimized approach to robustly wash off immunofluorescence signals and perform multiple rounds of FISH analysis on the same cells. This facilitates the genomic analysis and confirmation of circulating tumor cells (CTC), valuable for cancer diagnosis, prognosis, and progression monitoring. Prostate cancer patients with advanced clinical features showed an increasing number of epithelial/EMTing/mesenchymal CTCs. The presence of metastasis was efficiently predicted by a combined risk score based on both serum PSA level and EMTing CTC count. These analyses greatly enhance our ability to investigate the metastasis process and to predict/monitor cancer progression. The most important novel finding was the association of circulating megakaryocytes with good prognosis. In combination with mesenchymal CTC count, they showed great cancer prognosis potential. The identification of circulating megakaryocytes and their association with cancer prognosis potentially opens a new revenue to investigate cancer progression and to develop novel prognostic biomarkers.

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with each biomarker was derived from the Cox model as an increase in the hazard per unit increase in biomarker. Because of the small sample size, P values for the HR were further adjusted by false discovery rate (FDR) using Benjamini and Hochberg adjustment method (15). Survival curves were generated using the Kaplan–Meier method and compared using the log-rank test. All statistical tests were two sided and P values less than 0.05 (including multiple test adjustment where relevant) were considered as statistically significant. Bonferroni correction test was performed to modify P values for multiple tests through dividing the critical P value by the number of comparisons being made. Statistical analyses were performed using Stata 13.0.

### Results

#### Detection of circulating cells with epithelial and mesenchymal features

Using our optimized CTC isolation and detection method (9), we analyzed blood samples from 81 prostate cancer patients (Table 1; Supplementary Table S1) to identify CK+/VIM-/CD45- and CK+/VIM+/CD45- circulating cells (Fig. 1A). Three types of cells have been detected at high frequencies both in the blood of CRPC and localized cancer patients, and they are higher in CRPC patients than those in untreated patients (Table 1). Of 81 patients studied, 15 patients (19%) had no detectable 

#### Genetic evidence that CK+/VIM+/CD45- circulating cells are malignant cells with genomic alterations

To confirm the malignant feature of CK+/VIM+/CD45- circulating cells by detecting multiple genomic alterations, we developed a technology for repeated multiple rounds of FISH analysis after immunofluorescence staining and applied it on the CTC samples. This postimmunofluorescence multiple FISH analysis technique was developed using slides with lymphocytes spiked with PC3 cells, in which we removed the immunofluorescence signals completely by the stripping buffer, but not by 2× SSC buffer, fix solution, or proteinase K digestion. The length of polysilane slide coating time was optimized to 45 minutes to best preserve cells for downstream repeated FISH analysis after signal stripping. Less coating time frequently resulted in damaged or lost cells after signal stripping. Using these optimized conditions, we detected clear nuclear morphology and FISH signals in up to the fifth round of FISH on cells after immunofluorescence (Supplementary Fig. S1). In the leucocytes from the healthy donors, the average false-positive rates for the probes range from 0.7% to 7.1% (Supplementary Table S2).

The multiple FISH technique was then successfully applied to 11 prostate cancer CTC cases for five rounds of FISH (Fig. 1B) and one case for two rounds due to strong fluorescence background, to investigate the genomic alterations of nine genomic regions, including chromosomal copy number changes and/or rearrangements. In the leucocytes from 3 randomly selected patients, the average false-positive rates for the probes range from 1.5% to 7.8% (Supplementary Table S2). In CTCs from an individual, each of the genomic changes was only detected in a proportion of cells, indicating genomic heterogeneity. In the limited number of cases and limited number of CTCs in each case, no obvious genomic change patterns specific to a subgroup of epithelial or mesenchymal CTCs were observed. Changes of more than 30% of the genomic regions were detected in 68% of CK+/VIM+/CD45- cells, 57% of CK+/VIM+/CD45-, and 54% of CK+/VIM+/CD45- cells.

### Table 1. Summary of clinical characteristics and CTC count for all patients

<table>
<thead>
<tr>
<th></th>
<th>All patients, n = 81</th>
<th>Patient with untreated localized disease, n = 38</th>
<th>Patient with CRPC, n = 43</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, y</td>
<td>69 (76–62)</td>
<td>66.5 (72–57)</td>
<td>73 (81–67)</td>
</tr>
<tr>
<td>PSA, ng/mL</td>
<td>15 (71–7.65)</td>
<td>8.91 (12.25–5.40)</td>
<td>61 (367–23)</td>
</tr>
<tr>
<td>Primary GS, n (%)</td>
<td>6</td>
<td>11 (29)</td>
<td>2 (5)</td>
</tr>
<tr>
<td></td>
<td>3–4</td>
<td>14 (37)</td>
<td>3 (7)</td>
</tr>
<tr>
<td></td>
<td>4–8</td>
<td>9 (24)</td>
<td>3 (22)</td>
</tr>
<tr>
<td></td>
<td>B–10</td>
<td>4 (30)</td>
<td>20 (46)</td>
</tr>
<tr>
<td></td>
<td>n/a</td>
<td>0 (0)</td>
<td>13 (30)</td>
</tr>
<tr>
<td>Prior therapy, n (%)</td>
<td>No treatment</td>
<td>38 (47)</td>
<td>0 (0)</td>
</tr>
<tr>
<td></td>
<td>Systemic therapy</td>
<td>43 (53)</td>
<td>0 (0)</td>
</tr>
<tr>
<td></td>
<td>Metastasis, n (%)</td>
<td>40 (51)</td>
<td>38 (100)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>40 (49)</td>
<td>3 (7)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>40 (49)</td>
<td>40 (93)</td>
</tr>
<tr>
<td></td>
<td>CK+/VIM-/CD45- cell</td>
<td>49 (60)</td>
<td>18 (47)</td>
</tr>
<tr>
<td></td>
<td>Detectable patient number (%)</td>
<td>18 (47)</td>
<td>31 (72)</td>
</tr>
<tr>
<td></td>
<td>Cell number, median (IQR)</td>
<td>0 (2–0)</td>
<td>2 (2–0)</td>
</tr>
<tr>
<td></td>
<td>CK+/VIM+/CD45- cell</td>
<td>37 (46)</td>
<td>7 (38)</td>
</tr>
<tr>
<td></td>
<td>Detectable patient number (%)</td>
<td>0 (0–0)</td>
<td>30 (70)</td>
</tr>
<tr>
<td></td>
<td>Cell number, median (IQR)</td>
<td>2 (0–0)</td>
<td>2 (4–0)</td>
</tr>
<tr>
<td></td>
<td>CK+/VIM+/CD45- cell</td>
<td>59 (73)</td>
<td>21 (55)</td>
</tr>
<tr>
<td></td>
<td>Detectable patient number (%)</td>
<td>21 (55)</td>
<td>38 (88)</td>
</tr>
<tr>
<td></td>
<td>Cell number, median (IQR)</td>
<td>2 (0–0)</td>
<td>4 (8–2)</td>
</tr>
<tr>
<td></td>
<td>Total CTC</td>
<td>73 (90)</td>
<td>30 (79)</td>
</tr>
<tr>
<td></td>
<td>Detectable patient number (%)</td>
<td>30 (79)</td>
<td>43 (100)</td>
</tr>
<tr>
<td></td>
<td>Cell number, median (IQR)</td>
<td>6 (14–5)</td>
<td>11 (9–7)</td>
</tr>
</tbody>
</table>

*NOTE:* IQR, Q75%–Q25%.

Abbreviation: n/a, not available.
CD45− cells, but only detected in 3.7% of CK+/VIM−/CD45+ leucocytes and 7.8% of CK−/VIM+−/CD45− cells (Supplementary Table S3). The similar rate of genetic changes in the CK+/VIM+/CD45+, CK+/VIM+/CD45−, and CK−/VIM−/CD45− circulating cells indicates that the majority of CK+/VIM+/CD45+ cells were CTCs. Although it is possible that small proportions of the above three circulating cell categories are of nonmalignant origin, we considered all CK+/VIM+/CD45+, CK+/VIM+/CD45−, and CK−/VIM−/CD45− circulating cells found in noncancer healthy control cases were considered.

Association of CTC positivity with advanced clinical features in localized and metastatic prostate cancer

Analyzing blood samples from 24 healthy male donors, we detected one, two, and three CK+/VIM+/CD45− cells in three, two, and two samples, respectively, and none in the remaining 17 samples (median 0, range 0–3 cells/7.5 mL), and complete absence of CK−/VIM−/CD45− or CK−/VIM−/CD45+ circulating cells. Consequently, we defined positive CTC cases as those showing any CK+/VIM+/CD45+, any CK−/VIM−/CD45−, and/or >3 CK−/VIM−/CD45− cells to prevent potential false positive, although a better definition may be worked out from future large cohort studies. On the basis of these criteria, 24 of 38 patients with untreated localized disease (63%) and 41 of 43 CRPC patients (95%) scored positive for CTCs. Excluding mesenchymal CTCs reduced CTC positivity to 20 (53%) and 38 (88%) in these two groups of patients, respectively. In all 81 patients, high serum PSA level, high GS, and metastatic status were significantly correlated with CTC score–positive patients. In the 38 untreated localized diseases, CTC score–positive cases have a trend to be associated with high-risk classification based on National Comprehensive Cancer Network (NCCN) guideline version 1, 2016 (16), which should be further validated in a large sample cohort (Supplementary Table S4).

Association of subgroups of CTCs with serum PSA level, primary biopsy GS, and the risk of localized tumor

We further analyzed the relationship of different subpopulations of CTCs to PSA level, primary GS, and the risk of localized cancer. Although all three subpopulations of CTC numbers were significantly correlated to serum PSA levels (Spearman r = 0.28, 0.26, and 0.36, and P = 0.01, 0.019, and 0.0009, respectively, for CK+/VIM+−/CD45+, CK−/VIM−/CD45−, and CK+/VIM+/CD45+ CTCs), the mesenchymal type had the most significant association (Fig. 2A). As expected, the total number of CTCs had an even more significant correlation (Spearman r = 0.49, P < 0.0001) due to the greater number of events included for analysis. Because of the multiple tests for analysis, Bonferroni correction test was performed to modify the significant P values. After adjustment, mesenchymal CTC alone, but not epithelial and EMTing CTC alone, reached statistical significance. Traditional CTCs (combination of epithelial and EMTing CTC) had a significant correlation. Higher CTC count was also detected in patients with higher GS of primary tumor regardless of therapies received, and mesenchymal CTCs had the most significant association (P = 0.001). When total CTCs were included for analysis, a stronger association was observed (P = 0.0001; Fig. 2B; Supplementary Table S5). After multiple test correction, both epithelial and mesenchymal subtypes of CTCs alone were correlated with higher GS.

When the 38 untreated patients with localized disease were divided into low/intermediate and high-risk groups based on NCCN guideline, the number of each subtype of CTCs was higher.
Figure 2.
The correlation of CTC count with clinical features. A, Association of PSA with epithelial CTCs, EMTing CTCs, mesenchymal CTCs, total CK⁺ CTCs, total VIM⁺ CTCs, and total CTCs. The numbers of all types of CTCs were higher in patients with higher GS of primary tumor (B), patients with untreated high-risk localized disease (C), and patients with metastases (D). E, ROC analysis of the efficiencies of serum PSA level (AUC = 0.823) and CRS (AUC = 0.921) in discriminating metastatic prostate cancer patients from those without metastasis are shown in crosses and black dots, respectively. An AUC equal to 1 suggests perfect discrimination.
in high-risk group compared with low/intermediate group (Fig. 2C; Supplementary Table S5). After adjustment for multiple tests, EMTing CTCs and the traditional CTCs (epithelial and EMTing CTCs) still remained statistically significant.

EMTing CTC number was significantly associated with the presence of metastasis

When we correlated the numbers of different subpopulations of CTCs to cancer metastasis, the presence of metastases was significantly associated with higher number of any type of CTCs and the association with EMTing CTCs was most significant (P = 0.0001), which was similar in significance when considering total CTCs (Fig. 2D; Supplementary Table S5). Each subtype of CTCs alone achieved a statistical significance after P value adjustment for multiple tests. When a ROC curve was applied to explore the role of CTCs in metastases prediction in comparison with PSA, EMTing CTC count (with an optimal cut-off point at ≥2 cells) had the highest area under the ROC curve (AUC) score [0.755; 95% confidence interval (CI), 0.654–0.856] of all types of CTCs. Although the efficiency was lower than PSA (AUC = 0.823; 95% CI, 0.720–0.927, with an optimal cut-off point at ≥23 ng/mL; Supplementary Table S6), when EMTing CTC count and PSA were combined to create a CRS (as CRS = 0.012 · PSA + 0.115 · EMTing CTC count), the CRS (AUC = 0.921; 95% CI, 0.858–0.985) was a significantly (P = 0.03) better predictor of metastasis than PSA score alone (Fig. 2E). The increased performance of the CRS for metastasis prediction compared with PSA at different sensitivity and specificity level is presented in Supplementary Table S7. Box plots of CRS in patients with/without metastasis are shown in Supplementary Fig. S2A.

Mesenchymal CTC count is potentially associated with poor survival

Survival analysis was performed in the 40 metastatic CRPC (mCRPC) patients with a median follow-up time of 11.8 months (range, 0.7–19.7 months), of which 11 (27.5%) had died. Median follow-up time for patients still alive at the endpoint was 13.3 months (range, 4.1–19.7 months). Table 2 shows the association between potential risk factors and survival derived from Cox model with adjusted P value using Benjamini and Hochberg method (15). High PSA level (P = 0.015), high serum ALP level (P = 0.015), and high serum LDH level (P = 0.040) were found to be associated with increased risk of death in univariate analyses. None of the subtypes of CTCs had a significant association in this small cohort. However, when a cutoff was selected to optimally demonstrate the significance (Supplementary Fig. S3), patients with mesenchymal CTCs (≥5 cells) had an HR of 8.458 (95% CI, 1.815–39.411; P = 0.001). Patients with epithelial (≥3 cells), EMTing (≥2 cells), and total CK+/≥2 cells CTCs had an HR of 2.765 (95% CI, 0.732–10.438), 1.858 (95% CI, 0.491–7.033), and 1.537 (95% CI, 0.449–5.261), respectively; none reached statistical significance.

The presence of circulating megakaryocytes was associated with better survival

Unexpectedly, we found a rare population of cells in the harvest samples, with big nuclei (larger than both lymphocytes and most CTCs) of strong DAPI staining, but negative for CK, VIM, and CD45 (called BigNeg below; Fig. 3A). Because of the strong DAPI staining and larger size of nuclei, such cells were easily identified from other cells with DAPI staining alone (Fig. 3B). Through the absence of CK, VIM, and CD45 staining, it was possible to exclude CTCs. FISH analysis for these cells showed 100% polyplody and most (61.9%) of them had more than 10 copies of the genome in each cell (Fig. 3C), suggesting a special type of cells, distinct to lymphocytes or CTCs. The mean nuclear diameter of BigNeg cells was 17.5 ± 4.9 μm (range, 10–32 μm), whereas the mean nuclear diameter for lymphocytes was 6.5 ± 0.8 μm (range, 4–8 μm). No overlapping in nuclear size was observed between these two categories (Fig. 3D). Therefore, BigNeg cells were distinguishable from other cells by means of immunostaining and FISH measurement using the following criteria: cells with a strong DAPI-stained nucleus ≥10 μm in size and negative for CD45, CK, and VIM staining. Using these criteria, such cells were counted in all 81 blood samples. The numbers were variable and lower in mCRPC patients who died during follow-up (median 1 cell/7.5 mL; IQR, 2–1 cells/7.5 mL) than those mCRPC patients still alive (median 2 cells/7.5 mL, IQR: 4–1 cells/7.5 mL), and also than patients with untreated localized disease (median 2 cells/7.5 mL, IQR: 4.25–0 cells/7.5 mL), although none of the differences reached significance (P value of 0.18 and 0.33, respectively). In the 40 mCRPC patients, high BigNeg cell count had a trend to be associated with better survival (HR, 0.849; 95% CI, 0.628–1.146; Padj = 0.31; Table 2). When a cutoff was selected, patients with BigNeg (≥3 cells) had an HR of 0.144 (95% CI, 0.018–1.129; P = 0.02). The estimated survival rates were significantly higher in those with ≥3 BigNeg cells by Kaplan–Meier curve (P = 0.03, Fig. 4A). On the basis of previous results that high mesenchymal CTC counts generally represent more aggressive tumors and are associated with a higher risk for death, we hypothesized that the difference between the number of mesenchymal CTCs and BigNeg cells may further enhance the ability to predict survival. Using the formula [(mesenchymal CTC count – BigNeg cell count)/BigNeg cell count as detailed in the method and a univariate Cox model for survival analysis, the HR for per unit increase of the score is 1.282 (95% CI, 1.097–1.499; Padj = 0.015); Table 2]. When a cutoff was selected, patients with BigNeg (≥3 cells) had an HR of 0.144 (95% CI, 0.018–1.129; P = 0.02). The estimated survival rates were significantly higher in those with ≥3 BigNeg cells by Kaplan–Meier curve (P = 0.03, Fig. 4A).

In blood samples from the 24 healthy donors, the median number of BigNeg cells was 1 cell/7.5 mL (range, 0–15; IQR, 2.5–0; Supplementary Fig. S2B), which was slightly lower than that in the
38 patients with untreated localized disease (median, 2; range, 0–20; IQR, 4.25–0; Supplementary Fig. S2B; \( P = 0.19 \)) and marginally significantly lower than that in 43 CRPC patients (median, 2; range, 0–23; IQR, 3–1; Supplementary Fig. S2B; \( P = 0.078 \)).

Finally, we characterized the BigNeg cells. Their consistent hyperploidy revealed by FISH analysis highly suggested a potential megakaryocyte-like origin. We established the megakaryocytes immunoﬂuorescence analysis method by detecting CD34 and CD41 expression in megakaryocytes induced by phorbol myristate acetate from K562 cells (Fig. 4C). All BigNeg cells were CD34+ with 92% (101/110) CD41+, confirming them as a specific group of megakaryocytes (Fig. 4D). CD45+ lymphocytes, epithelial CTCs, EMTing CTCs, and mesenchymal CTCs were all negative for CD34 and CD41.

**Discussion**

We have previously shown that CD45− cells with both epithelial and mesenchymal features can be detected in prostate cancer patients using Parsortix (9). Here, we developed a repeated FISH analysis approach following immunoﬂuorescence analysis on the same slide to investigate multiple genomic regions for alterations and, consequently, establish the malignancy of VIM+/CD45− cell populations, which were associated with disease burden and poor prognosis and have the potential to serve as an additional marker to traditional epithelial CTCs for clinical application. As EMT is a critical step for cancer metastasis, analyzing EMT CTCs has great potential for cancer prognosis and progression monitoring (7, 17). Several non–EpCAM-based CTC isolation platforms have been developed to analyze CTCs with EMT, such as ISET (18, 19) and Vitatex CAM platform (20). CTCs with ongoing EMT or completely changed into mesenchymal features have been identified, and their biological and genetic differences from CellSearch captured CTCs have been revealed. To our knowledge, the current study is the first genetic investigation of EMT CTCs using repeated FISH and the first investigation of potential prognostic value of EMT CTCs in prostate cancer.

Repeated FISH analysis of CTCs generates genomic alteration information for multiple genomic regions in single cells, enabling the conﬁrmation of the malignant genomic feature of suspicious CTCs, investigation of the heterogeneity of cancer cells by analyzing the differences in genomic alterations between individual cells, and correlation of genomic alterations with cellular features and different types of CTCs to understand mechanisms of metastases. FISH analysis performed on cells analyzed by immuno-staining (21–24) has been previously performed by using traditional fixation with ethanol or methanol mixed with acetic acid (24) and treatment with pepsin (21) to remove immuno-
signals. However, based on our experience, it was difficult to achieve complete removal of previous immunofluorescence signals using these pretreatments and the leftover fluorescence signals often interfered with FISH signal interpretation. Our stripping buffer method, robustly removing immunofluorescence completely without damaging cell morphology, facilitates the analysis of multiple genomic alterations on the same cells after immunofluorescence, which increases the chance to detect genomic alterations in most CTCs to confirm their malignancy nature. A panel of genes is better than a single gene or genetic change for cancer prognosis (25). Multiple genomic region analysis also makes it possible to combine a number of genomic changes for the development of CTC-based genetic prognostic biomarkers. However, in this study, FISH was primarily used to confirm the malignancy of EMT CTCs. We observed a high frequency of alterations in CTCs at the stage of cancer metastasis. This is consistent with a previous report that metastatic prostate cancer has more genomic alterations than primary tumor (26). Our results also showed that genomic heterogeneity exists in prostate cancer CTCs.

In this study, we demonstrated the association of CTC number with GS and the aggressiveness of localized disease, which have not been reported previously. Association of detection of CTCs with poor prognosis in patients with advanced prostate cancer was initially reported around a decade ago (27–29) and has been recently further validated in several large-scaled clinical trials (30–33). In our small patient cohort, we observed the associations of ≥3 epithelial CTCs (CK⁺/VIM⁻/CD45⁻), ≥2 EMTing CTCs (CK⁺/VIM⁺/CD45⁻), and ≥6 traditional CTCs (combination of epithelial and EMTing type) with advanced clinical features,
CRPC development, and poorer survival, which is consistent with these previous reports analyzing epithelial CTCs. By analyzing mesenchymal CTCs, we detected an increase of CTC-positive cases in both CRPC and localized diseases. Of interest, high baseline mesenchymal CTC count (≥5 cells/7.5 mL) before an alternative treatment in CRPC patients was better associated with poorer survival than the traditional CK+ CTCs in our cohort, which warrants further investigation. Moreover, mesenchymal CTCs associated with PSA level and GS better than epithelial CTCs and EMTing CTCs were most significantly correlated to cancer metastasis. All these data indicate a potential better prognostic value of mesenchymal than epithelial CTCs in prostate cancer. As the number of patients in this study is still small, these associations should be validated in large cohorts of samples. EMT has been increasingly recognized for its role in tumor cell invasion, metastatic dissemination, and acquisition of therapeutic resistance (6). Loss of epithelial markers, such as E-cadherin, and gain of mesenchymal markers, such as VIM and N-cadherin, have been proven to be associated with more invasive phenotype or higher GS in prostate cancer cells (34, 35). EMTing CTCs were most significantly associated with both high-risk localized disease and metastasis, suggesting that these cells under active EMT process are potential indicators for cancer-aggressive invasion and metastasis. Taken together, compared with traditional CK+ CTCs, CTCs undergoing or undergone EMT provided more information about disease burden and intrinsic tumor biology.

Radionuclide bone-scan and CT, the current gold standard procedures to detect metastatic sites, are costly, time-consuming, and expose patients to radiation. Patients who are unlikely to have metastases are better off avoiding these costly and potentially harmful procedures. PSA level has been used to predict bone metastases and select patients for bone scans (16). Although our data also showed a strong association between PSA and metastasis and none of the CTC analyses taken alone outperformed PSA for metastatic correlation, a CRS based on both PSA level and EMTing CTC count significantly improved the metastasis prediction accuracy compared with PSA alone. This suggests that although PSA level correlates with CTC count, they are independent factors contributing to cancer metastasis, potentially by representing different aspects of tumor biology. Large-scale trials are warranted to confirm this superior metastasis prediction model.

The most striking finding is the unexpected discovery of increased circulating megakaryocytes in cancer patients and its association with the survival of CRPC patients. As the nuclei of megakaryocytes are big and with strong DAPI staining, they can be easily identified during CTC analysis. A group of large hyperploid cells with no detectable biomarkers (including epithelial and hepatocellular cancer specific markers) have also been reported recently (36). As the authors did not include any mesenchymal markers, they considered all these cells as CTCs with EMT. On the basis of our study, these cells might be a mixture of mesenchymal CTCs and circulating megakaryocytes. Megakaryocytes have also been identified in a previous CTC study of prostate cancer, but the investigators excluded them for further analysis (23). Taking advantage of the size and/or deformability-based CTC isolation platform, for example, Parsortix here, such cells can be captured with high efficiency for cancer prognosis analysis. With the development of multiple treatment approaches for CRPC patients, real-time prognostic and therapeutic response–predictive biomarkers are critical for stratified patient management. However, currently, only CellSearch-based CTC analysis, which mainly detects cells with epithelial features, is approved by FDA for CRPC patient survival prediction (2). Here, we not only showed that mesenchymal CTCs has the potential to be better associated with patient survival than CTCs with epithelial features, but also found that the combination of mesenchymal CTC and megakaryocyte counts has a great power to predict CRPC patient survival, with 1.28-fold risk of death per unit increase of the CMS and 10-fold risk for patients with a score ≥2.0.

In addition to the bone marrow and the spleen, megakaryocytes have also been detected at very low numbers in circulation (37, 38). CD34 expresses in hematopoietic stem cells and megakaryocyte progenitor cells, which are all diploidy cells (39). Its expression can also be seen on mature-appearing megakaryocytes in reactive and disorders of bone marrows (40). CD41 is a megakaryocytic lineage-specific marker, expressing neither in endothelial cells nor monocytes (39, 41, 42). The circulating megakaryocytes detected in our study with high DNA content and consistent expression of CD34 represent a specific and potentially abnormal type of megakaryocytes, in which endonuclear DNA replication is active but the CD34 expression is not switched off, hence premature high DNA content megakaryocytes without apparent lobulated nuclei. This is consistent to the finding of CD34 expression in mature-appearing megakaryocytes in reactive and disorders of bone marrows (40), and increased ploidyization of megakaryocytes has been reported in patients with metastatic tumors (43). Platelets, which are released from megakaryocytes in bone marrow, have recently been recognized to play an important role in cancer metastasis (44, 45), and the gene expression profile of platelets has been shown to efficiently distinguish individuals with or without cancers (46). Platelet counts have also been investigated for cancer prognosis, and the data are still in conflict (47). In this study, there was no correlation of the number of platelets with either circulating megakaryocytes or cancer prognosis.

The association of high number of megakaryocytes with a good survival of CRPC patients suggests an antitumor effect of megakaryocytes. To support this, megakaryocytes have been reported to increase their number in response to cancer bone metastasis and inhibit prostate and breast cancer cell growth both in vitro and in vivo (48–50). Although the prognostic value and biological functions of megakaryocytes in cancer development and progression need to be further investigated, models that analyze the number difference between CTCs and megakaryocytes should be developed to make an efficient biomarker for survival prediction for patients with mCRPCs and potentially for other cancers as well. With further validation and cutoff optimization in large sample cohorts, this study has the potential to be translated into clinical use, for patients not only with prostate cancer but also a range of other human malignancies.

Disclosure of Potential Conflicts of Interest

G. Shaw reports receiving speakers bureau honoraria from and is a consultant/advisory board member for Merck Sharp & Dohme. Y. Lu reports receiving commercial research grants from ANGLE plc. No potential conflicts of interest were disclosed by the other authors.

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Acknowledgments
We would like to thank N. Lemoine, J. Fitzgibbon, and T.V. Sharp for critical review of the manuscript and thank all patients and healthy donors participating in this study.

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Grant Support
This work was supported by Orchid, Cancer Research UK [C16420/A18066] and Angle PLC. L. Xu and T. Guo also thank Chinese Scholarship Council for funding support for their PhD studentship (201306100035 and 201506330098).

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Received December 7, 2016; revised March 10, 2017; accepted June 7, 2017; published OnlineFirst June 14, 2017.


The Novel Association of Circulating Tumor Cells and Circulating Megakaryocytes with Prostate Cancer Prognosis

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