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Accessibility
An RNA-based signature enables high specificity detection of circulating tumor cells in hepatocellular carcinoma

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Contributed by Kurt J. Isselbacher, December 11, 2016 (sent for review October 14, 2016; reviewed by Anil K. Rustgi and Timothy Wang)

Circulating tumor cells (CTCs) are shed into the bloodstream by invasive cancers, but the difficulty inherent in identifying these rare cells by microscopy has precluded their routine use in monitoring or screening for cancer. We recently described a high-throughput microfluidic CTC-CHIP, which efficiently depletes hematopoietic cells from blood specimens and enriches for CTCs with well-preserved RNA. Application of RNA-based digital PCR to detect CTC-derived signatures may thus enable highly accurate tissue lineage-based cancer detection in blood specimens. As proof of principle, we examined hepatocellular carcinoma (HCC), a cancer that is derived from liver cells bearing a unique gene expression profile. After identifying a digital signature of 10 liver-specific transcripts, we used a cross-validated logistic regression model to identify the presence of HCC-derived CTCs in nine of 16 (56%) untreated patients with HCC versus one of 31 (3%) patients with a nonmalignant liver disease at risk for developing HCC (P < 0.0001). Positive CTC scores declined in treated patients: Nine of 32 (28%) patients receiving therapy and only one of 15 (7%) patients who had undergone curative-intent ablation, surgery, or liver transplantation were positive. RNA-based digital CTC scoring was not correlated with the standard HCC serum protein marker alpha fetoprotein (P = 0.57). Modeling the sequential use of these two orthogonal markers for liver cancer screening in patients with high-risk cirrhosis generates positive and negative predictive values of 80% and 86%, respectively. Thus, digital RNA quantitation constitutes a sensitive and specific CTC readout, enabling high-throughput clinical applications, such as noninvasive screening for HCC in populations where viral hepatitis and cirrhosis are prevalent.

The shedding by epithelial cancers of circulating tumor cells (CTCs) into the bloodstream underlies the blood-borne dissemination of cancer, although only a small fraction of CTCs gives rise to metastases (1). Enumeration and analysis of CTCs may thus enable noninvasive monitoring of advanced cancers, as well as early detection of invasive but localized tumors before they give rise to viable metastases. Recent advances in CTC isolation provide sensitive and high-throughput platforms to enrich for these rare tumor cells within blood specimens, but antibody staining and microscopic imaging of captured cancer cells remain a critical bottleneck limiting broad application of the technology (2). Classical CTC staining criteria include the presence of cell surface epithelial cell adhesion molecule (EpCAM) and cytoplasmic epithelial cytokeratins and the absence of the hematopoietic CD45 marker (3), but epithelial marker expression is highly variable and extensive imaging criteria must be applied to score immunofluorescent signals reliably from rare cancer cells surrounded by contaminating leukocytes (4). Emerging microfluidic CTC isolation technologies that effectively deplete leukocytes without manipulating tumor cells (5) preserve cell viability and ensure high-quality RNA content, as demonstrated by single-cell RNA sequencing studies (6–8). These CTC isolation platforms now enable the application of powerful RNA-based digital PCR (dPCR) technologies to score molecular signatures of cancer cells, thus providing a potentially robust and high-throughput readout for the presence of CTCs within blood specimens. To test the feasibility of RNA-derived digital scoring of CTC-enriched cell populations, we applied this strategy to hepatocellular carcinoma (HCC), a cancer that lacks defining gene mutations but originates in liver cells with unique tissue-specific expression profiles.

Liver cancer is the second highest cause of cancer mortality worldwide, leading to 765,000 deaths in 2015 (9). In the developing world, the high prevalence of hepatitis B virus (HBV) infection drives the incidence of HCC; worldwide, it is estimated that 248 million individuals are infected with HBV (10). The risk of developing HCC is calculated as 0.5–1% per individual per year in HBV carriers without cirrhosis and as high as 8% per individual per year in patients with cirrhosis (11). Developed countries are also witnessing a rise in HCC incidence, linked with cirrhosis due to chronic hepatitis C virus (HCV) infection, alcohol abuse, obesity-associated nonalcoholic fatty liver disease

![Image](https://example.com/image.png)
(NAFLD), and nonalcoholic steatohepatitis (NASH) (9). Early-stage HCC is potentially curable by thermal ablation, surgical resection, or liver transplantation, with a 5-y survival of 50–80% following these therapies (9). Once the tumor disseminates within or outside the liver, however, therapeutic options are limited and 5-y survival declines to below 15% (11).

Although early detection of HCC in high-risk individuals offers a strategy for successful curative treatment, screening in patient populations with liver cirrhosis has been limited by the low test characteristics of the primary biomarker, serum levels of alpha fetoprotein (AFP) (11). CTCs have been reported in patients with HCC, but the characteristically low EpCAM cell surface expression in this tumor type has limited the utility of standard CTC measurements (12, 13). To establish a high-throughput, blood-based assay for HCC that would have broad applicability, we therefore adapted the microfluidic CTC-iChip readout combined with droplet detection of CTC-derived transcripts whose expression is retained in HCC.

We applied this molecular CTC assay as proof of principle in a pilot cohort of patients with HCC and high-risk patients suffering from liver disease.

Results

Establishment of the HCC-Specific RNA dPCR Assay. Fig. 1A outlines the RNA-based dPCR CTC-scoring assay. CTCs were isolated using the CTC-iChip microfluidic device, which depletes hematopoietic cells from blood by size-based exclusion of red blood cells, platelets, and plasma, followed by magnetic depletion of white blood cells (WBCs) tagged with magnetic bead-conjugated CD45, CD16, and CD66b antibodies (5, 14). The high efficiency depletion of WBCs (4- to 5-log purification) enriches CTCs, which are admixed with some contaminating WBCs (<500 WBCs per milliliter of processed whole blood) (5, 14). To improve CTC imaging with high-throughput detection of CTC-derived transcripts, we coupled whole-transcriptome amplification (WTA) of CTC-derived RNA with dPCR amplification, in which cDNA molecules are encapsulated within individual aqueous droplets and multiple transcripts of interest are quantified by in-droplet PCR amplification (Materials and Methods).

We curated liver-specific transcripts whose expression is preserved in HCC cells, but absent in blood components. Of the 20,000 genes measured from publicly available microarray datasets (15), the 100 most highly expressed genes in HCC were screened against expression profiles of hematopoietic cells and other normal tissues (16). These genes were then validated against a separate RNA-sequencing dataset, comparing 10 primary HCCs (17) with WBCs from eight healthy donor blood samples processed through the CTC-iChip (Fig. 1B). Quantitative RT-PCR (qRT-PCR) of cDNA from purified healthy donor WBCs (n = 3) was used to eliminate genes with low but detectable background signal (Fig. S1). Based on these results, 10 genes (AFP, alpha 2-HS glycoprotein (AHSG), albumin (ALB), apolipoprotein H (APOH), fatty acid binding protein 1 (FABP1), fibrinogen beta chain (FGB), fibrinogen gamma chain (FGG), glypican 3 (GPC3), retinol binding protein 4 (RBP4), and transferrin (TF)) were selected for developing the dPCR assay.

To technically validate this strategy, we determined the number of transcripts measured from the introduction of individually micromanipulated cells from the liver cancer cell line HepG2 known to express albumin into 4 mL of blood from healthy donors, followed by CTC-iChip processing and dPCR for the liver-specific transcript ALB (Fig. 1C). No ALB-positive droplets were observed in unspiked blood processed through the CTC-iChip, whereas a range from 2 to 100 spiked HepG2 cells generated from 240 to 28,800 ALB transcripts. Given the heterogeneity of HCC cells within clinical specimens, we applied low-template WTA to maximize starting material and optimized the 10-gene liver-specific panel to ensure that this dramatic signal amplification preserved the relative distribution among multiple markers. We ensured the absence of amplification bias in three independent experimental replicates (Fig. 1D and Fig. S2). Compared with unamplified cDNA derived from 1 ng of

Fig. 1. dPCR quantitation of HCC cells after microfluidic enrichment from blood. (A) Schematic representation of the integrated platform for digital RNA-dPCR scoring of CTCs. Hematopoietic components are depleted from whole blood through CTC-iChip processing as previously described (5). RNA from the CTC-enriched product is subjected to WTA, encapsulation of cDNA molecules within lipid droplets, and PCR amplification for transcription of interest. (B) Heat maps derived from microarray (Left) and RNA-sequencing (RNAseq) (Right) datasets comparing expression of 10 liver-specific transcripts in HCC versus other tissues. The microarray dataset compares fetal liver, adult liver, and 10 cases of HCC (15 samples shown of 79 tissues tested) and blood components (15, 16). RNAseq compares 10 cases of HCC (17), with WBCs collected from eight healthy independent donor (HD) blood samples processed through the CTC-iChip. (C) dPCR quantitation of ALB transcripts from micromanipulated HepG2 spiked into whole blood and processed through the CTC-iChip. Each data point represents one-sixth of the CTC-iChip product. (D) Pie charts representing the distribution of transcripts for each of the 10 selected liver-specific genes, following dPCR analysis of 1 ng of HepG2-cell RNA. Samples were nonamplified or subjected to WTA (three independent reactions). (E) Total number of transcripts of interest after spiking increasing numbers of HepG2 cells into blood (n = 3), CTC-iChip processing, and dPCR. (F) Pie charts depicting the relative fraction of droplets for each of the 10 target transcripts after spiking increasing numbers of HepG2 cells into blood and CTC-iChip processing, as noted in E (n = 3).
HepG2 RNA, the increased overall signal resulting from WTA (25,000- to 100,000-fold amplification per gene) preserved the relative proportion of each transcript among the WTA replicates (Fig. S2 A and B).

To test the sensitivity of the 10-gene digital assay in rare tumor cells admixed with blood cells, we again micromanipulated either 1, 3, 5, 10, or 50 single HepG2 cells into 4 mL of healthy donor blood, which was then processed through the CTC-iChip, followed by WTA and dPCR. In two of three replicates, a single spiked HepG2 cell was detected, with an average 5,000-fold increase in signal over unpurified blood controls. All liver-specific genes were detected with progressively increasing signal and preservation of marker distribution as the numbers of input cells increased from 1 to 50 cells ($R^2 = 0.79$; Fig. 1 E and F). The high sensitivity of dPCR raised the possibility that CTC enrichment might not be required to detect tumor-derived transcripts in nucleated blood cell fractions. Past reports have suggested that standard RT-PCR amplification might identify PCR products comigrating with the expected ALB transcript from unpurified blood cells of patients with HCC (18); however, we were unable to reproduce this finding using the more sensitive and specific dPCR technology ($n=9$ HCC samples; Fig. S3). We therefore applied the combination of microfluidic CTC enrichment followed by dPCR detection of HCC-derived transcripts to clinical specimens from HCC patients.

**Digital CTC Detection in Patients with HCC.** We evaluated the performance of the optimized digital CTC-scoring assay in blood samples (5–15 mL) from six patient cohorts, per Institutional Review Board (IRB)-approved protocols at Massachusetts General Hospital. The cohorts studied included (i) healthy blood donors ($n=26$, median age = 55 y); (ii) patients with high-risk nonmalignant chronic liver disease (CLD), including hepatitis virus-associated cirrhosis, who were being routinely monitored for HCC development ($n=31$, median age = 58 y); (iii) newly diagnosed, untreated patients with HCC ($n=16$, median age = 66 y); (iv) patients with HCC actively receiving therapy with radiographically evident disease ($n=32$, median age = 67 y); (v) patients with HCC who had undergone curative-intent interventions, such as ablation, resection, or liver transplantation, and clinically had no evidence of disease (NED) any longer ($n=15$, median age = 66 y); and (vi) patients with primary malignancies other than HCC, with or without liver metastases ($n=44$, median age = 62 y). Patients in categories i through vi were categorized by a clinician blinded to the dPCR data. Clinical characteristics of these cohorts are provided in Tables S1–S6.

dPCR analysis of CTC-iChip–processed blood specimens from patients with HCC generated high signal for individual transcripts, compared with healthy donors, patients with CLD, or patients with other malignancies (Fig. 2A). As expected, intra-cohort variability was observed, with some HCC patient samples exhibiting high signal from multiple liver-specific transcripts and others containing few transcripts of interest. HCC cases in which no signal was detected may reflect the absence of CTCs within the single 5- to 15-mL blood sample or expression of transcripts that are not captured by the 10-gene panel.

To integrate the results of multiple genes into a statistically robust scoring model, we screened each transcript to determine if it served as a statistically significant single-gene predictor of HCC status (Fig. S4). Nine of the 10 genes met this selection criterion (all excluding GPC3). These genes were then combined into a single metric CTC score, using a leave-one-out cross-validated (LOOCV) multivariate logistic regression model (Fig. S5). The LOOCV allowed us to build and test the model using a single HCC patient cohort, although the high stringency associated with this approach may underestimate the predictive value of the model; Fig. 2C demonstrates the change in model performance with and
without cross-validation. We tested the multigene model with 48 patients with HCC versus 57 patients without cancer (Fig. 2C).

Nine of 16 (56%) untreated patients with HCC were classified as positive by CTC score, compared with only one of 31 (3%) patients with at-risk nonmalignant CLD and two of 26 (7.6%) age-matched healthy blood donors (P < 0.0001, χ²; Fig. 2B). Patients with HCC undergoing therapy but with radiographically detectable disease had a lower fraction of cases with positive CTC scores [nine of 32 (28%)], but this fraction was still significantly higher than the control population (P = 0.004, χ²). Patients with NED after curative-intent treatment were only positive in one of 15 (7%) cases, an incidence comparable to the incidence of the control population (P = 0.56, not significant). Together, these results demonstrate that the CTC score can identify patients with active disease while maintaining a high degree of specificity. Among all patients with HCC, the CTC score was not associated with specific underlying risk factors (P = 0.73), but it was highly correlated with Barcelona Clinic Liver Cancer staging (P = 0.011) and trended toward significance when stratifying by vascular invasion (P = 0.06; Fig. S6).

We next determined the feasibility of differentiating between patients with HCC and patients with malignancies other than HCC using a separate logistic regression (Fig. S7). In this comparison, six of the 10 genes were statistically significant predictors of HCC vs. non-HCC status (AFP, AHSG, APOH, FABP, FGB, and FGG), yielding a multipredictor model with different features than the previous model (Fig. S7A and B). Among patients with pancreatic, prostate, breast, and non-small-cell lung cancers; cholangiocarcinoma; and melanoma, 39 of 44 (88%) cancers/carcinomas were correctly distinguished from HCC at a sensitivity of 50% for patients with HCC. Optimal differentiation of tissue of origin among CTCs will likely benefit from the inclusion of additional markers that are specific for other tumors, in addition to exclusion of HCC-associated transcripts.

Longitudinal Monitoring of Patients with HCC. The higher incidence of positive CTC scores in newly diagnosed, untreated patients with HCC (56%) compared with those patients with HCC who were undergoing active treatment (28%), and those patients with HCC who completed curative-intent therapy (7%) suggests a potential role for CTCs in longitudinal monitoring of tumor response. Furthermore, increasing tumor burden, as defined by the Barcelona Clinic Liver Cancer staging criteria, is associated with increased CTC score values (Fig. 2A and Fig. S6). A subset of patients with HCC in our study was monitored longitudinally for tumor response. Supporting the robustness of the assay, the CTC score remained high in two patients (HCC-041 and HCC-075), with no therapeutic intervention or change in clinical status between blood draws (Fig. 3A and Fig. S7E). Two other patients (HCC-058 and HCC-060) underwent surgical tumor resection and demonstrated a decrease in CTC score postoperatively (Fig. 3B). The CTC score for patient HCC-042 decreased impressively following an immune checkpoint inhibitor (nivolumab) treatment, and then showed a further reduction 3 wk after subsequent radioembolization of the tumor (Fig. 3C). Coincident computed tomography scans demonstrated a significant tumor response to radioembolization. Of note, for three of these five patients (HCC-042, HCC-058, and HCC-075), serum protein AFP measurements were below clinically informative values (<20 ng/mL) at all draw points. Although serum AFP protein measurements are often used for monitoring tumor response in patients with HCC, they are below detection in a significant fraction of cases. In such cases, CTC score monitoring may serve as a complementary marker to assess disease status.

Early Detection of HCC in High-Risk Populations. Although early detection of localized HCC in individuals with liver cirrhosis provides the only hope for curative therapy, serum AFP alone does not provide sufficient accuracy to enable screening of at-risk populations. Using a cutoff of 20 ng/mL, AFP has an estimated sensitivity of 53% with a specificity of 87%, leading to a positive predictive value (PPV) of 6% in populations where the expected prevalence of HCC is 1% (19). Raising the AFP threshold to 100 ng/mL improves specificity to 99% but reduces the test sensitivity to 31% (20). Given these poor test characteristics, the American Association for the Study of Liver Diseases does not recommend using AFP as a screening tool for HCC (11).

To model the potential combination of AFP and CTC score in screening for HCC, we determined the correlation between these two biomarkers in all of the patients with HCC for whom corresponding assay results were available. CTC score and serum AFP levels were not significantly correlated (R² = 0.0007, P = 0.74), with concordance restricted to cases with high serum AFP levels (Fig. 4A). The discordance between AFP protein levels and the CTC score is consistent with the underlying basis for these assays: The multigene CTC score is a digital signature to quantify HCC cells that have invaded into blood, whereas serum AFP measures a single protein produced by HCC cells and released into blood from tumor deposits. As such, the two assays may be orthogonal and have added value as blood-based biomarkers for HCC.

Although our pilot study was not powered to test the accuracy of the CTC score directly in early detection of HCC, we modeled two strategies using either a high or low cutoff for AFP measurements. First, we tested the additive value of CTC score positivity and high-threshold AFP (100 ng/mL). Of the 15 patients with newly diagnosed HCC for whom both AFP and CTC scores were available, four (27%) were detected by CTC score alone, one (7%) by AFP alone, and five (33%) by both assays (Fig. 4B). Together, either AFP or CTC score was positive in 67% of patients, leaving only one-third undetected by either method. Importantly, among all 16 patients with newly diagnosed HCC, six (38%) patients met the Milan criteria for liver transplantation, a clinical indication that the disease is sufficiently localized to enable curative therapy. CTC scores were available for all six patients who met the Milan criteria, of whom two (33%) were positive (Fig. 4C). AFP levels were available for five of six patients, but none were above the 100-ng/mL threshold. Thus, a
of heterogeneous CTCs among an excess of contaminating WBCs. Moreover, the dramatic signal amplification and the molecular specificity derived from dPCR provide an effective way to detect the rare but highly biologically significant occurrence of intact tumor cells in the blood circulation. As an initial proof of concept, we applied this digital CTC measurement strategy to HCC, a highly lethal malignancy with worldwide impact, for which early detection strategies are currently inadequate (11).

The concept of liquid biopsies for noninvasive monitoring of cancer has emerged as one of the most promising approaches in cancer diagnostics, with applications ranging from early detection to treatment selection and monitoring response (23, 24). Three complementary technologies each interrogate different biological specimens and rely upon distinct technological assays. Circulating tumor DNA (ctDNA) is derived from small fragments of genomic DNA shed into the vasculature by dying tumor cells, amid the background of DNA released from normal tissues, and it provides powerful opportunities for targeted DNA-based genotyping (25). However, an inherent limitation of ctDNA-based genotyping is that it does not provide information as to the tissue of origin for mutations detected in the blood, in contrast to CTCs, which provide a source of intact RNA for lineage-based analysis. Genotype-based cancer detection is also of limited utility in tumors such as HCC, where highly prevalent mutations have not been identified. An alternative blood-based cancer detection technology takes advantage of exosomes, small membrane-bound cellular fragments encapsulating cytoplasmic RNA and other cellular components that are released by both tumor and normal cells. Although strategies for enrichment of tumor-derived exosomes are still being optimized, the analysis of pooled exosomes has allowed RNA-based detection of cancer-associated mutations (26). However, the fact that normal tissues also abundantly shed exosomes precludes the use of lineage and tissue markers to identify tumor-derived expression signatures in blood specimens. In contrast to exosomes, whole cells derived from normal tissues are extraordinarily rare in the bloodstream. Hence, the initial isolation of intact CTCs in the bloodstream, followed by their molecular quantitation, may provide a highly specific diagnostic assay that is amenable to large-scale clinical applications. HCC, which arises within well-defined at-risk populations, and in which early detection may be curative, is particularly appropriate to serve as a “proof of principle” for CTC-based screening. Of note, the sensitivity and specificity of this molecular CTC assay hinge on the successful identification of HCC-derived transcripts: Additional technologies that capture the full heterogeneity of heterogeneous CTCs, and analysis of larger blood volumes may also improve the likelihood of CTC detection, especially in patients with small lesions.

The poor performance of AFP alone in HCC screening stems from the fact that only a minority of patients with liver cancer have very high elevations in this marker (>100 ng/mL), whereas low levels (>20 ng/mL) are common in conditions that predispose to HCC, including viral hepatitis. However, the combination of AFP screening with a second more specific assay has been shown to be effective in reducing disease mortality. In a large randomized trial, a 37% reduction in HCC-related mortality was reported among high-risk patients who underwent surveillance with AFP and liver ultrasound (27). Ultrasound-based surveillance is now practiced in many centers, but image quality is operator-dependent and degraded in the setting of obesity or cirrhosis. Furthermore, access to high-quality ultrasound is limited in developing countries, which bear the greatest burden of HCC. The technology that we describe here, together with the analysis of pilot clinical cohorts and clinical modeling studies, raises the possibility that digital CTC scoring may provide an important new tool for HCC detection. The scalability of digital CTC monitoring may be particularly useful in underserved populations that lack access to MRI and ultrasound screening. Although large population-based studies are now required to test and validate this technology’s performance against existing screening standards, we favor the combination of AFP screening.
and CTC-scoring assays as orthogonal markers that, together, may provide both the sensitivity and specificity required for noninvasive and cost-effective HCC screening in high-risk populations that se-
rum AFP alone is unable to provide. Alternative CTC-scoring al-
gorithms may also enhance the differentiability of HCC from other hepatic lesions, an application of particular importance in the United States, where the use of imaging tests currently drives HCC
detection and monitoring. Thus, quantitative analysis of multiple
tissue-specific transcripts derived from CTCs may be optimized for
distinct applications in the diagnosis and treatment of patients with HCC.

Finally, we note that digital scoring of CTCs for cancer moni-
toring is broadly applicable to other cancer types. Indeed, many
cancers originate in tissues that express specialized gene tran-
scripts that are absent in normal blood cells, and the curation and
testing of these markers may enable high-sensitivity detection and
monitoring of rare cancer cells in the blood. Such blood-based
molecular monitoring for CTCs holds considerable promise for
the early detection of cancer.

Materials and Methods

Gene Target Identification and Validation. Publicly available RNA-sequencing
and microarray datasets were used to identify the top 100 genes highly expressed in HCC
in comparison to expression in other tissues and blood components (15–17). The low expression of candidate genes within WBCs persisting in the CTC-
Chip output was confirmed by RNA-sequencing of processed blood from healthy
donors, and qRT-PCR of WBCs purified from whole blood was used as an addi-
tional exclusion criterion. Ten genes were selected to establish a signature
of HCC-derived CTCs, enriched within a background of normal blood cells.

Cell Culture and RNA Processing. HepG2 cells were cultured following American
Type Culture Collection-recommended culturing conditions. RNA was isolated
using the RNeasy Plus Micro kit (Qiagen), and cDNA was generated using the
SuperScript III Reverse Transcriptase kit (Thermo Fisher). Buffy coat WBC prepa-
rations were generated using standard procedures, followed by TRIzol RNA
extraction (Ambion). For qRT-PCR assays, 5 ng of cDNA generated from HepG2
cells or from WBCs (buffy coats) of three independent blood donors was amplified
and compared with GAPDH using the Applied Biosystems 7500 RT-qPCR assay (40
cycles). Primer and probe combinations are provided upon request. For single-cell
molecular manipulation and spike-in studies, individual cells were micropipetted using an
Eppendorf TransferMan NK2 micromanipulator and introduced into whole blood
samples from healthy donors, before processing through the CTC-iChip.

Patient Cohorts and Blood Processing Through the Microfluidic CTC-Chip. Pa-
tient cohorts and clinical characteristics are provided in Tables S1–S6. Dana–
Farber/Harvard Cancer Center IRB-approved protocols were used to obtain
consent from all patients and healthy donors. Blood was processed using the
CTC-iChip as previously described (5). A detailed description of each cohort
and blood processing conditions can be found in SI Materials and Methods.

WTA and Droplet dPCR. WTA was performed on CTC-iChip-derived RNA using
the SMARTer Ultra Low-input RNA kit, version 3 (Clontech Laboratory). One-
third of the RNA extracted from the CTC-iChip product was loaded into each
reaction. The dPCR experiments were performed using an AutoDG automated
droplet generator, C1000 Touch Deep-well thermocycler, and QX200 plate
reader (BioRad) following manufacturer recommendations, with 1% of the WTA
product. Thermocycling conditions and primers are provided upon request.

Statistical Calculations and Logistic Regression. Genes that served as statisti-
cally significant predictors of disease status were used to build a multi-
predictor logistic regression whose performance parameters were determined
using LOOCV (Fig. 2C and Figs. S4, S5, and S7). For longitudinally collected
cells or from WBCs (buffy coats) of three independent blood donors was amplified
and compared with GAPDH using the Applied Biosystems 7500 RT-qPCR assay (40
cycles). Primer and probe combinations are provided upon request. For single-cell
molecular manipulation and spike-in studies, individual cells were micropipetted using an

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