Detection and Molecular Characterization of Circulating Tumor Cells (CTCs) in Patient with Metastatic Melanoma: A Potential Application of Liquid Biopsy

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Abstract

Melanoma specific anti NG2 antibody conjugated with iron oxide nanoparticles that have been developed to isolate, detect, and culture CTCs from melanoma xenograft models and blood from patients with metastatic melanoma. The enrichment process involved, lysis of RBC from blood samples using RBC lysis buffer enhances the cancer cell recovery by immunomagnetic labeling and separation. Efficient cell capture was validated using fluorescently labeled cancer cells spiked into healthy human blood and clinical utility was demonstrated in specimens from patient with metastatic melanoma. The spiking experiment results with greater than 70% recovery in four sets of experimental. Spontaneous metastasis melanoma model, found that the number of CTCs increased during the tumor progression and correlated to lymphatic as well as lung metastasis development in vivo. CTCs were detected in 6 out of 7 patients’ blood samples that contained metastatic melanoma. The CTCs were positive for melanoma specific molecular markers such as S100, HMB45, MelA, MITF and the CTCs were positive for anti NG2-Q-dot and pERK2-Q-dot staining. The BRAFV600E gene expression pattern resulted that the CTCs were expressed BRAFV600E similar to the control melanoma cells. The iron oxide antibody nanoparticle isolation method is a highly reliable and cost effective detection method of CTCs from patients with metastatic melanoma and CTCs can be cultured for additional analytical studies or potential drug sensitivity testing.

Keywords: Melanoma; CTCs; Iron oxide; Antibody; Metastasis

Abbreviations: CTCs: Circulating Tumor Cells; IO: Iron Oxide; MITF: Microphthalmia-associated transcription factor; Q-dot: Quantum-dot

Introduction

Circulating tumor cells (CTCs) are cancer cells shed from primary and secondary metastatic tumor sites which are then transported through the blood stream to distant vital organs. These cells are then mixed with the other components of the blood making their isolation and characterization technologically difficult due to their rarity when compared to the frequency of other components of the blood. Detection and enumeration methods of CTCs from peripheral blood have been examined in many types of metastatic cancer such as breast [1-3], prostate [4-6] and colorectal [7-9] cancers in addition to melanoma [10-13]. The amount of CTCs present in the blood stream of cancer patients is extremely small and it has been estimated for every CTC in [10-13]. The enrichment process involved, lysis of RBC from blood samples using RBC lysis buffer enhances the cancer cell recovery by immunomagnetic labeling and separation. Efficient cell capture was validated using fluorescently labeled cancer cells spiked into healthy human blood and clinical utility was demonstrated in specimens from patient with metastatic melanoma. The spiking experiment results with greater than 70% recovery in four sets of experimental. Spontaneous metastasis melanoma model, found that the number of CTCs increased during the tumor progression and correlated to lymphatic as well as lung metastasis development in vivo. CTCs were detected in 6 out of 7 patients’ blood samples that contained metastatic melanoma. The CTCs were positive for melanoma specific molecular markers such as S100, HMB45, MelA, MITF and the CTCs were positive for anti NG2-Q-dot and pERK2-Q-dot staining. The BRAFV600E gene expression pattern resulted that the CTCs were expressed BRAFV600E similar to the control melanoma cells. The iron oxide antibody nanoparticle isolation method is a highly reliable and cost effective detection method of CTCs from patients with metastatic melanoma and CTCs can be cultured for additional analytical studies or potential drug sensitivity testing.

Numerous studies have shown that CTCs can be detected by reverse transcriptase polymerase chain reaction (RT-PCR) targeting specific mRNA gene expression in cancer cell types [19,20]. Melanoma markers such as tyrosinase, MUC18, MITF glycoprotein gp100/pmel17, and MART1/mela.A were all used as molecular markers in the detection of melanoma CTCs. The sensitivity of this molecular technique has been well demonstrated. However, the quality control for diagnostic use remains problematic because of the false positive and false negative results [21]. Advanced molecular techniques such as quantitative real time polymerase chain reaction (qRT-PCR) and microarray gene expression showed increased detection sensitivity as well as improved quantitative analysis of CTCs isolated from melanoma patients [22]. Several qRT-PCR based multi marker assays such as tyrosinase, p97, MUC16, melanA/MART1, MAGE3 and/or gp100/pMel-17 have been reported for CTC detection in melanoma patient blood samples [22-24].

The presence of tumor cells in blood is predictive of short survival in several cancers and their isolation and characterization can guide towards the use of more effective treatment. The CTCs are highly heterogeneous in nature and number of clinical research studies has been undergoing the validation of CTCs observation and characterization. In this paper, we demonstrated the separation of circulating tumor cells (CTCs) from in vivo melanoma xenografts.

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models and metastatic melanoma patient blood samples using a melanoma specific anti NG2 (9.2.27) antibody conjugated with iron-oxide nanoparticles under a magnetic field. This cells surface antigen is highly expressed on melanoma cells. The isolated rare CTCs cells were able to proliferate under in vitro conditions.

**Materials and Methods**

**Cell lines**

The B16-F10 murine melanoma, C8161, 1205Lu, and 451Lu A375M melanoma cell lines were cultured in DMEM containing 10% fetal bovine serum. Primary human melanoma cells from melanoma patient’s tumor tissues by previously published protocol [25]. Briefly, the tissue samples were washed with PBS and the tissues were mechanically chopped into small pieces followed by enzymatic digestion using 20 µg/mL concentration of collagenase type 1 solution and incubation of the tissues at 37°C for one hour. The cells were sieved through a cell strainer (100 µm) size and the cells were washed with PBS and cultured in DMEM supplemented with 10% FBS and Pen/Strep solution. For GFP lent viral production, a GFP plenti viral vector was co-transfected into 293T cells with a packing vector (pCMVΔ8.2durp and pCMV, VSVG), (generous gift from Dr. Xiaowei Xu at university of Pennsylvania), and the viral supernatants were collected at 48 and 72 h post transfection [26].

**Patient and blood sample collection**

After the informed consent for participant in this institutional review board-approved study, each patient’s blood (5-10 mL) was collected in EDTA tubes at the time of surgery in the university hospital and processed within 2 hrs. The research operators were blinded to all clinical and histopathological correlative information during the sampling analysis process.

**Red blood cell lysis**

In the clinical blood samples from cancer patients and healthy donors, the red blood cells were removed by applying a RBC lysis buffer (154 mM NH₄Cl, 10 mM KHCO₃ and 0.1 mM EDTA). Briefly, 1 mL of a blood sample was added into 25 mL of pre-warmed RBC lysis buffer and incubated for 5 min in a shaking condition at room temperature. The sample was then centrifuged at 300 g for 5 min. The cell pellets were washed two times with pre-warmed PBS and the cells were suspended in 1 mL DMEM. The cell suspension was then ready for magnetic labeling.

**Immunomagnetic labeling and magnetic separation**

To perform immunomagnetic labeling, we have used a carboxyl magnetic iron oxide nanocrystal conjugation kit. The IO np-Ab conjugation was performed according to the manufacturer’s instructions (Ocean Nano Tech, Springdale, AR). Briefly, 200 µL of magnetic iron oxide nanoparticles and 200 µL of activation buffer were aliquoted in a sterile 96 well plate and 100 µL of EDAC/NHS was added then incubated for 10 min with continuous mixing at room temperature. 500 µL of anti NG2 (9.2.27) purified antibody and 500 µL of coupling buffer were added into the tube and mixed well then incubated for 2 hours with continuous mixing. Next, 10 µL of quenching solution was combined with the above mentioned reaction mixture and incubated for another 10 min with continuous shaking. The reaction mixture was then transferred into a plastic cuvette and 1 x wash/storage buffer was added. The reaction mixture in the cuvette was then placed into the Super Mag Separator™ magnetic separator (Ocean Nano Tech, Springdale, AR) separate the magnetic iron oxide nanoparticles at 4°C for 4 hrs. The separated iron oxide- anti NG2 [9.2.27] antibody complex was washed with 3 mL of 1 x wash buffer and stored in 1 mL of 1 x wash/storage buffer. To perform cell separation 50 µL of iron oxide- 9.2.27 antibody conjugated nanoparticles was incubated with 1 mL of RBC lysed blood sample for 2 hrs at room temperature and the immunolabeled cells were separated by using a Super Mag Separator™ magnetic separator for 2 hrs. The magnetically separated cells were washed with 1x PBS and the cells were then resuspended in DMEM supplemented with 10% FBS and Pen/strep solution then plated in a 24 well plate and incubated in CO₂ incubator at 37°C. The media was changed after 24 hrs with tumor fibroblast condition media with 10% DMEM (1:1 ratio).

**Anti NG2 [9.2.27] antibody production**

The murine hybridoma cells secrete an IgG monoclonal antibody against a 240 kD human melanoma surface glycoprotein. The anti NG2 hybridoma cells were grown in a T75 flask containing filter sterilized DMEM with 4 mM L-glutamine and 1.5 g/L sodium bicarbonate supplemented with 10% fetal bovine serum. The culture was maintained at 37°C in 5% CO₂ incubator. To make anti NG2 antibody, the viable hybridoma cells were seeded into a flask containing a combination of BD cell™ M Ab medium animal component free media (BD biosciences CA and DMEM supplemented with 10% FBS in the ratio of 50:50. The monoclonal antibody containing media was collected according to the hybridoma culture method and the concentration of antibody was determined by standard protein estimation methods.

**In vivo studies**

All animal studies were performed under a protocol approved by the IACUC at the Pennsylvania State University. Green Fluorescent Protein (GFP) tagged murine melanoma cells B16-F10 (2 x 10⁵ cells per injection) and metastatic human melanoma cells 451Lu, 1205Lu and A375M (2.5 x 10⁵) were injected into the flank region of the intradermal layer of C57B/L6 mice for B16- F10 GFP cells and in athymic nude mice injected 451Lu, 1205Lu and A375M cells. The tumor challenging experiments were preformed observed for 21 to 24 days and measure tumor size, collect blood for CTC analysis, dissect out the lymphnodes and other internal organs at each time points.

**Q- Dot staining, immunocytochemistry, western blotting and qRT-PCR**

The melanoma and CTC cells were grown on eight well chamber slides and the monolayer cells were fixed with 4% paraformaldehyde and stained with primary antibodies specific for MITF (Neomarker, Carlsbad, CA) S100, HMB45, Mela-A, CD45 (Dako North America, INC Carpinteria, CA) and anti-NG2 (9.2.27) antibodies. The cells were incubated with primary antibodies overnight at 4°C. After washings, the cells were incubated with appropriate Alexa fluor secondary antibodies (Life Technologies, Grand Island, NY) and mounted on DAPI containing mounting media (Vector laboratories Inc., Burlingame, CA, USA). Q-dot staining was performed according to the manufactures protocol (Life Technologies, Grand Island, NY), for western blot analysis, a monolayer of melanoma cells and the CTC cells were washed with ice- cold PBS and lysed in RIPA buffer or lamella buffer to analyze total protein content. Fifty microgram of proteins were separated in Nu-PAGE4-15% Bis-Tris Gel (Life Technologies, Grand Island, NY) and α-Enolase (Santa Cruz, CA) in 5% milk TBST buffer (150 mmol/L Tris-HCl. (pH 8.0), 150 mmol/l NaCl and 0.01% Tween.
20 and 5% non-fat milk. The membranes were washed thrice with wash buffer for 5 min and incubated with horseradish peroxidase-conjugated secondary antibody and washed again before being processed with chemiluminescence reagents (ECL Western Blotting Detection System; Pierce, Rockford, IL, USA). Quantitative RT-PCR was performed according to previously published protocol [26].

**Statistical analyses**

The data represent mean ± S.E.M values. The differences between two experimental groups were determined using Student’s t-test. A two-tailed value of $P<0.05$ was considered statistically significant.

**Results**

**Characterization of Iron Oxide (IO) nanoparticles**

Carboxyl magnetic iron oxide (IO) nanocrystal conjugation kit was purchased from Ocean Nano Tech (Springdale, AR). The IO nanocrystals were conjugated with melanoma specific antibody anti NG2 (9.2.27) according to the manufacture’s protocol. The size and charge of the anti NG2- Iron Oxide nanoparticles were measured by zeta analyzer (Malvern Zetasizer- nanoseries, Malvern Instruments limited, Worcestershire, UK). The particle size observed for the antibody conjugated IO particle was increased (Figure 1A and 1B) along with an increase in the zeta potential of the IO-Ab nanoparticles from -36.8 mV to -35.6 mV. The less negative zeta potential can be attributed to the surface charge shielding of the carboxyl groups on the surface of the unconjugated iron oxide [IO] nanoparticles after conjugated to the amine groups of the antibodies. The increase in size may be attributed to the conjugation of the antibodies plus the water of hydration on the conjugated surface. The migration of the conjugated IO-antiNG2 nanocrystals observed during gel electrophoresis in a 1% agarose gel in a 1% agarose gel towards the positive pole was much slower than the unconjugated IO nanocrystals. This can be attributed to the increased molecular weight of the IO-antiNG2 complex and less negative zeta potential (Figure 1C).

Since the capture and conjugation of antibody was performed directly from the reaction media, it is important to check the binding affinity of antibody with the iron oxide nanobeads, which was done by resolving the conjugated antibody on a SDS-PAGE gel (data not shown) and the results indicated that both the heavy (55 KDa) and the light chain (25 KDa) seen in iron oxide conjugated antiNG2 antibody.

**Anti NG2 protein expression in melanoma cells**

Anti NG2 (9.2.27) hybridoma cells were cultured in a hybridoma specific culture medium. After obtaining a continuous culture, the monoclonal antibody was isolated from the culture medium and purified by affinity column chromatography. The purified anti NG2 antibodies were the subjected to western blot analysis as well as immunocytochemistry (Figure 2) melanoma cells (1205Lu, 451Lu, A375M, WM793 and UACC903), and breast cancer cells (MCF7) used as a negative control. The western blot results showed that melanoma cells were expressed high level of anti NG2 protein (Figure 2A). Furthermore, we have detected this cell surface marker protein in melanoma cells by immunocytochemical staining methods. Three independent observations were performed for the evaluation of cell surface protein. Melanoma Cell C8161 (Figure 2B), and tumor derived melanoma cell lines (data not shown) showed a high intensity of anti NG2 staining.

**Spiking melanoma cells into human blood and culture in vitro**

There are certain limitations of CTC isolation methods used to isolate CTCs from the blood of tumor-bearing animals that only permit in situ analysis of collected cells within the separation device. In this method, we demonstrated that melanoma cells spiked into the blood can be isolated and then cultured in vitro. It is important to note that this experiment does not mimic actual model of melanoma metastasis in which cells would have to detach from a primary tumor site and survive circulation through the entire vasculature before being isolated. For the effective capture and detection of CTCs, we used melanoma-specific antibody NG2 conjugated with iron- oxide nanocrystals. We spiked different cell numbers (1000, 500) of metastatic melanoma cells (1205Lu-GFP, 451Lu-GFP) and cells derived from metastatic melanoma tissue samples tagged with Green Flourescent Protein (GFP) into fresh healthy human whole blood. The result shown in Figure 2C, the 1205Lu-GFP, 451Lu-GFP, and melanoma tumor cells (data not shown) were successfully captured directly from whole blood and RBC lysed samples. The same amount of cells spiked into the whole blood and RBC lysis samples were seeded in a separate chamber as a positive control. In addition, used breast cancer cell line (MCF7) and human fibroblast (FF10) was used as negative control for testing the binding affinity of Iron oxide conjugated Anti NG2 antibody. The percentage of recovery was calculated according to previously published methods [27]. The percentage of recovery was higher The samples that underwent RBC lysis with percent recoveries of 70% and 85% which were substantially higher than the percent recovery values from the whole blood samples (less than 30%). The captured cells were cultured in 10% DMEM and successful colony formation and proliferation was observed without any deterioration of cell morphology and viability (Figure 2D). We obtained blood and tumor tissue samples from patients diagnosed with a BRAFV600E-mutant and BRAF WT cutaneous melanoma. The BRAF mutation appears to have critical role in tumor initiation and melanoma progression. We investigated the gene expression pattern of BRAFVT [Sbc3, WM3211] and BRAFV600E [WM793, 1205Lu] melanoma cell lines and CTCs isolated from melanoma patient’s blood samples. Our result demonstrated that, the expression of BRAFV600E expression was homologous in melanoma primary cell lines and in CTCs of patients with metastatic melanoma (Figure 2E). However, the melanocyte and BRAFWT cell lines, the BRAFV600E mRNA expression remain same or less expressed.

**Assessment of spontaneous metastasis**

Here we report that syngenic model of spontaneous metastatisis of B16-F10 mouse melanoma cells in C57BL/6 and highly metastatic melanoma cell (451Lu, 1205Lu and A375M) in athymic nude mice with a very high frequency that mimics clinical metastatic melanoma. Two hundred thousand of B16-F10 mouse melanoma cells as well as metastatic human melanoma cells (2.5 × 10^6) were injected intradermally into the flank region of the skin. Metastatic melanoma is typically first detected by the presence of lymph node metastases and less often by systemic metastases. Lympathic and visceral metastases were observed in all animals tested. The GFP tagged melanoma cells were observed in all animals at day 7 and an increase in the amount of GFP cells during the tumor progression was apparent. The tumor weight was correlated with the tumor growth time and also the significant number of micro metastases in the lymph nodes. However the appearance of CTCs and lung metastasis was not seen during the early stages of the tumor development in both tumor progression models (Figure 3A and 3B). To evaluate the efficacy of anti NG2-iron oxide nanocrystals techniques for in vivo CTC capture, we separated circulating tumor cells (CTCs) from the blood of melanoma xenograft models. The blood samples (~1 ml) were collected from those animals 2 weeks after tumor cell implantation were mixed with anti NG2 coated magnetic beads and were subjected to magnetic separation. Isolated CTCs were then washed with PBS and plated in 24
Figure 1: Characterization of Iron Oxide (IO) nanoparticles. A and B) Charge and size of iron oxide conjugated with melanoma specific antibody anti-NG2. The biocompatible water soluble iron nanoparticle size around 40 nm ± 5 with a zeta potential of -36.6 mV and after antibody conjugation with size of iron oxide was around 61 ± 5 with a zetapotential of -35.6 mV. C) Agarose gel electrophoresis of biocompatible water soluble iron oxide [IO] (on left) and IO-anti NG2 antibody (on right), the electric pole is negative for top and positive for the bottom.
Figure 2: Melanoma cell separation from DMEM and in spiked human blood and RBC lysis buffer. A) Western blot analysis of 9.2.27 protein expression in metastatic melanoma cell lines. Erk2 antibody used as loading control. B) Immunostaining of anti-NG2 antibody in human melanoma 451Lu GFP cells. GFP over expressing melanoma cells 1205Lu. C) 451Lu cells separated using anti NG2 conjugated iron oxide (n=3). The percentage of recovery rate was significantly high in RBC lysis method. D) Image of melanoma cells recovered from blood spiking using anti NG2 iron oxide nanobeads (Scale 200X). E) Quantitative RT-PCR was performed and showed that BRAFV600E expression level in different melanoma cell lines and melanoma patient CTCs.
well plates containing DMEM supplemented with 10% FBS and pen/strep solution. After 24 hr, GFP tagged melanoma cells were counted under a fluorescent microscope. We observed that the CTCs appeared in the blood after the 14-16th day of post cancer cell injection in both xenograft models. The number of CTCs increased during the tumor progression. The CTCs adhered well and proliferated on the culture dishes used (Supplementary Figure 1A). The isolated CTCs remains amephenotypic and genotypic characteristic of the parental cells. These results clearly indicate that rare CTCs isolated from the blood collected from xenograft models retain their viability and proliferation capacity during the transition from in vivo to in vitro conditions.

Capture of CTCs from patient with metastatic melanoma

Blood samples from six patients in various stages of treatment for metastatic melanoma were processed with anti NG2 conjugated iron-oxide nanoparticle. We also processed blood from a healthy individual as a control (n=3). The captured cells were cultured in regular culture conditions and CTC cells were attached and grown on the surface of the culture dish with the media being changed every two days. After a week, the growth of CTC cells was clearly visible in the dish and the CTCs had morphological features analogous to the pointed spindle shape of melanocytes (Supplementary Figure 1B) and could not detect any CTCs cells in control samples. The CTCs cells were trypsinized and grown in chamber slides for immunocytochemical examination with melanoma specific antibodies such as S100, MITF, HMB45, Mela A and a leukocyte marker CD45 (Figure 4A). Furthermore, we quantified the positively stained CTCs cells against DAPI staining to confirm percentage of melanoma marker staining (Figure 4B). The western blot analysis showed that four out of five CTCs from melanoma patients expressed MITF protein, (Figure 4C). The western blot analysis used
Figure 4: Immuno fluorescent analysis of magnetically isolated CTCs from metastatic melanoma patient blood samples. A) Immunostaining of CTC cells stained with melanoma specific biomarker protein S100, MITF, HMB45 and MelaA. B) Percentage of CTCs cells stained with melanoma against DAPI staining (Scale 200X). C) Western blot analysis of MITF protein expression in CTCs isolated from melanoma metastatic blood sample using anti NG2 conjugated iron oxide nanoparticle. D) Q-DOT staining of CTCs, anti NG2antibody conjugated to q-dot probe (Scale 400X). E) Wright-Gysma staining leukocyte staining of CTCs isolated from melanoma patient blood sample.
metastatic melanoma cells 1205Lu as a positive control, fibroblast cells as a negative control, and α-enolase as a loading control. The isolated CTCs conformation that had melanoma specific antigen, we performed a Qdot staining with anti NG2 antibody conjugated with a Qdot probe (diameter=585 nm ±), and a significant level of low abundance expression of melanoma specific antigen in these cells was observed (Figure 4D). The melanoma CTCs were expressing phosphoERK2 after staining with Q-dot probe conjugated with pERK antibody (diameter=605 nm ±) (Supplementary Figure 1C). To further validation, we performed a double stain with S100 and CD45. The result showed that the melanoma specific S100 were co-localized with CD-45 leukocyte markers (Supplementary Figure 1D). Leukocyte contamination is common problem associated with CTCs isolation methods. So we stained the isolated CTC’s cells with Wright-Gymsa dye to localize leukocytes contamination in the isolated CTC’s cells. It showed that some cells show positive with the staining but some cell are were not stained with Wright-Gymsa dye (Figure 4E).

Discussion

Tumor cell invasion and extravasation leads to the presence of circulating tumor cells (CTCs) in peripheral blood. However, the CTC counts in patients with cancer would be helpful for analyzing the metastatic mechanism in addition to clinical applications for diagnosis and therapy. In the present study, we have successfully captured and cultured circulating tumor cells (CTCs) from blood samples from melanoma xenograft models and metastatic melanoma patients. The process involved polymer coated 30 nm iron oxide nanocrystals that were modified with melanoma specific NG2 antibodies. In this study, we demonstrated the separation of CTCs using melanoma specific antibody conjugated with iron oxide nanoparticles (IO-Ab) in an applied magnetic field. The iron oxide nanoparticles were synthesized using a pyrolysis-based method in organic solvent allowing precise control of particle size and crystallinity [28]. The iron oxide nanoparticles which were soluble in organic solvent were modified with polymers to make them water soluble before further modification of the iron oxide nanoparticle with antibody attachment. The successful conjugation of the anti NG2 Ab to the iron oxide nanoparticles was tested using gel electrophoresis. After evaluation of stability, the antibody conjugated IO nanoparticles were used for cancer cell separation from culture media (DMEM) and separation of cancer cells spiked into fresh human blood with and without RBC lysis buffer treatment. The most common problem associated with nanomaterials in biological applications is their compatibility/stability at various biological parameters including, pH, salinity, viscosity and toxicity [29,30]. The nanomaterial releases toxic heavy metal ions that cause problems in biological sample separation especially in CTC capturing and the establishment of growth of CTCs in culture conditions. In our studies we successfully captured and grew CTCs from melanoma xenograft models and melanoma patient blood samples without any toxicity problem associated with the iron-oxide nanoparticles.

Recently Clawson et al. reported that CTCs were isolated from metastatic melanoma patient blood samples by simple centrifugation method (OncoQuick column) and characterized by staining with panKRT a melanocytic marker which was aberrantly expressed in melanocyte differentiation [31]. Previous studies were reported that CTCs can be isolated from patient with different types of cancer such as breast cancer using anti HER2 antibody [28] and for prostate cancer using anti PSA antibody conjugated iron oxide nanoparticles [32]. However no studies have been reported with melanoma specific antibody antiNG2 antibody conjugated with iron oxide nanoparticles for CTC isolation and characterization from metastatic melanoma patient blood samples. The accurate quantitative measurement and stability of performance of standards remains challenging. There has been inaccurate assessment as to how many CTCs are released from the primary tumor into the blood daily. The isolation of CTCs is technically challenging due to the rare number (1 in 10^9 red blood cells). Multiple batch approaches have been employed to detect CTCs, including cell size based separation [33,34] and fast scanning cytometry [35]. In most of the cases the CTCs are rendered non-viable due to the pretreatment process. However the IO based CTC capturing technologies have superiority over other methods, because cell viability remain unchanged and they would proliferate to some extent. While conducting the melanoma cell spiking experiment, we used 6 different cell lines such as UACC903, 1205Lu, 451Lu, A375M, C8161 and B16 mouse melanoma cells which over express GFP protein. The recovery rate was consistent with all the cell lines either in blood or in RBC lysis methods. However the CTCs capturing performance was not same as from cancer patient blood samples. Gertler et al. proposed that CTCs can be distinguished from normal blood cells depends on its physiological properties such as size and difference in cell density, charge, migratory properties and melanocytic granules [36]. Several studies showed that CTCs have significant variation in size difference, and buoyant density compared to the leukocytes [34,37]. These results provide significance outcome in capturing cancer cell lines spiked into blood and further validation required with clinical samples in which heterogeneous nature of cancer cells in the blood. The molecular characterization of CTCs has been conducted in many previous studies. Both qRT-PCR and RT-PCR based methods can detect CTCs and that positive markers are associated with metastatic disease and poor prognosis in melanoma patients [38]. The specificity of these qRT-PCR assays remains controversial because of the indication of false positive results due to the heterogeneous nature of mRNA expression of melanoma associated biomarkers in the blood of melanoma patients [39]. The use of immunomagenetic nanobead capture of melanoma cells in our approach exploits the selective expression of melanoma surface antigen in melanoma cells. The CTC isolation techniques rely on antibody-based capture of CTCs which express melanoma specific cell surface markers that are absent from normal leukocytes. Conjugation of antiNG2 (9.2.27) to iron oxide nano beads followed by purification of captured cells through a magnetic field and subsequent staining with melanoma specific antibodies to visualize the expression of melanoma marker protein in captured CTCs. Staining for the leukocyte specific marker CD45 widely used as a negative control to exclude containing leukocyte [40]. Wechler et al. (ASCO meeting abstract) isolated CTCs from melanoma patient using Screen Cell method, the isolated CTCs were express S100, MART1/MelaA and HMB45 and their results were consistent with our techniques of iron oxide nano beads conjugated with antiNG2 antibody and cancer cell recovery rates were comparatively higher than Screen Cell system. Our findings revealed the potential clinical usefulness of an immunomagenetic nanobead assay for capturing and detecting CTCs in the blood of melanoma patients and melanoma xenograft models. Future studies involving blood analysis and long term follow-up of patients may be needed for a more detailed assessment of efficacy of drug studies and the clinical utility of this assay.

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References


