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Single-cell tracking reveals super-spreading brain cancer cells with high persistence

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A B S T R A C T

Cell migration is a fundamental characteristic of vital processes such as tissue morphogenesis, wound healing and immune cell homing to lymph nodes and inflamed or infected sites. Therefore, various brain defect diseases, chronic inflammatory diseases as well as tumor formation and metastasis are associated with aberrant or absent cell migration. We embedded multicellular brain cancer spheroids in Matrigel™ and utilized single-particle tracking to extract the paths of cells migrating away from the spheroids. We found that – in contrast to local invasion – single cell migration is independent of Matrigel™ concentration and is characterized by high directionality and persistence. Furthermore, we identified a subpopulation of super-spreading cells with >200-fold longer persistence times than the majority of cells. These results highlight yet another aspect of cell heterogeneity in tumors.

1. Introduction

Tumor cell migration is a hallmark of cancer and is typically divided into three major types: amoeboid-, mesenchymal-, or collective migration [1]. Amoeboid and mesenchymal migration is the movement of single cells. Although cancers tend to exhibit one specific kind of migration – such as amoeboid migration by lymphoma cells – migrating cancer cells are known for their high plasticity and ability to interconvert between different modes of migration in response to environmental cues [2,3]. In contrast, collective migration i.e. invasion involves the cooperative transport of whole groups of cells and describes the local expansion of tumor cells into the extracellular matrix (ECM).

A common system used to model tumors in 3D is that of tumor spheroids. Spheroids are dense multicellular structures that form due to most cells’ natural tendency to aggregate under certain environmental conditions [4]. The advantages of spheroids as tumor models include tighter cell–cell adhesion than cells cultured in 2D and that cells are not equally exposed to the local environment; instead there is an innermost and an outermost region emerging. This gives rise to nutrient, oxygen and cell proliferation gradients from the surface to the core [5,6]. Cells in spheroids therefore exhibit different phenotypes depending on their proximity to the surface much like cancer cells in solid tumors which are e.g. dormant or necrotic at the core [6,7]. For studies of metastasis, spheroids are embedded in a collagen, hydrogel or Matrigel™ matrix to mimic the natural cancer environment [8–11]. Thus, this model system allows true three-dimensional migration/invasion of the cancer cells into the surrounding matrix.

Migration of tumor cells is a multistep process initiated by the epithelial to mesenchymal transition (EMT) where cells lose their apical-basal polarity and change their expression of e.g. surface adhesion molecules to allow a more migratory phenotype [12]. This transition is in itself a multistep process with multiple semi-stable intermediate states [7,13,14]. In order to migrate, mesenchymal cells have to develop pseudopods which protrude from their body and form focal contacts with the ECM. Proteases expressed on the cell surface then up-concentrate at these contacts to degrade the ECM locally and carve tunnels for the cell as it moves. The actual motion is driven by actomyosin contraction within the cell followed by detachment of the trailing edge by disassembly of focal contacts [2]. Due in part to the plasticity of cancer cells, the exact integrins, proteases and signaling molecules involved in this cascade of events depend not only on the cancer type but also on the microenvironment and the homogeneity among individual cells [2,7].

A number of different random walk models have been used to describe cell migration in various species [15,16]. Specifically, persistent random walk (PRW) models derived from the Ornstein–Uhlenbeck process have been very useful for describing the motility of various species from Dictyostelium and mouse fibroblasts to human endothelial cells [17–21]. However, there are several examples of motile cell types that are not exhaustively modeled as simple persistent random walk models.
walkers [22–26]. In the specific case of glioblastoma invasion and migration, a sigmoidal Gompertzian model has been used to describe the growth of glioblastoma spheroids in vitro [27], while a cellular automaton model was used to highlight the importance of cell–cell adhesion/attraction during glioma cell migration [28]. Further, the PRW model has been used to describe both the growth and diffusion of cells away from glioblastoma spheroids [29]. While this work separated the proliferation and migration into two distinct populations, it has been shown that the combined growth and migration can be described by a density-dependency i.e. cell migration is inversely proportional to cell density, which is highest in the proliferating zone [30–32]. A number of studies have also modeled glioblastoma behavior regarding morphology, metabolism, vasculature, and medical treatment [33], for instance the targeting of chemotherapeutic peptides [34]. However, studies on the dynamics of glioblastoma cell migration are still limited and there seems to be no consensus on the subject.

We therefore set out to further characterize the invasion and migration of brain glioblastoma cells (U87-MG) from multicellular tumor spheroids, a cell type forming particularly tight spheroids [35]. We developed a brightfield live-cell imaging setup to follow the invasion of Matrigel™-embedded spheroids as well as subsequent migration of individual cells away from it. We found that the ability to invade the extracellular matrix is compromised by matrix stiffening. However, our analysis – based on machine learning algorithms [36] – also yielded insights into the migration behavior of individual cells and we found this to be highly ECM density independent. Surprisingly, we identified a subpopulation of fast migrating cancer cells. We found that these super-spreaders are not moving significantly faster along their trajectory, but with much more persistence.

2. Results

We cultured gravitation-assisted brain cancer spheroids by seeding ~ 650 U87-MG cells in 200 μL low-attachment U-formed wells. The spheroids were harvested after 4 days of incubation. At this point, they had an average diameter of 278 ± 11 μm (mean ± SD, N = 34), an average aspect ratio of 1.32 ± 0.09 (mean ± SD, N = 34), and a density of 3 · 10^4 cells/μm^3 [37]. To kick-start invasion, we interchanged growth medium with the ECM formulation Matrigel™ which consists of laminin, collagen IV, fibronectin, and various growth factors and, thus, mimics basement tissue membranes [38]. We used a range of concentrations of Matrigel™ in medium (v/v) from 25% to 75%, thus subjecting spheroids to various mechanical loads [39,40] and imaged the spheroids for more than 24 h (maximum 31 h) with a frame rate of 1 every 5 min. All our data was obtained with simple wide-field microscopy, which limits sources of potential phototoxicity but complicates image segmentation. Examples of frames from the resulting time-lapse movies are shown in Fig. 1A for a spheroid in 50% ECM at time t = 0, 8, 16, 24 hours. Here it is obvious that there are two types of motility in play: (i) branch-like pattern of cells moving collectively through the ECM as well as (ii) individual cells leaving the spheroid body. These different motility modi are well-known [35,41,42] and correspond to invasion and migration, respectively.

2.1. Extracellular matrix stiffening slows down invasion

We measured the time-dependent average invaded cross sectional area, (A(t)), as given by Eq. (3) and showed in Fig. 1B for various ECM concentrations ranging from 0% to 75% Matrigel™ concentration in medium (v/v). Here a concentration of 0% corresponds to growth in medium only (gray asterisks) with an average growth rate of 23 [21–26] μm/day (N = 9). Hence, we confirm that the cross sectional area (A(t)) of the tumor spheroid grows linearly with time in fresh culture medium [43–45]. In contrast, for the invading spheroids, we found an exponential dependence between cross-sectional area and t with the characteristic growth exponent, β,

\[ A(t) \propto e^{\beta t}, \]  

(1)
as shown in Fig. 1C. To the best of our knowledge, an exponential growth of the spheroids’ cross-sectional area in ECM has not been reported before. It should be noted, however, that our findings of such fast expansion and invasion of the brain cancer spheroids for the first 24 h after ECM addition are in agreement with previous reports of various cell lines invading type I collagen [46,47]. The exponential exponent β appears to correlate with ECM concentration, such that the higher the ECM concentration, the slower the spheroids invade. However, for 67% ECM, we detect an abrupt decrease in β compared to the slightly lower and higher ECM concentrations of 50% and 75%.

However, for all cases of non-zero ECM concentration, the spheroids’ final average areas, (A(t)), are significantly larger than the 8.4 ·10^4 μm^2 spheroids in 0% ECM (gray bar) in Fig. 1D. This signifies that the resulting volume of an invading spheroid is larger than that of a non-invasive – yet still proliferating – spheroid. The most pronounced difference is found for spheroids in 0% vs. 25% ECM (p < 0.001) while further increases in ECM concentration slows down invasion. For instance, for spheroids in 75% compared to 25% ECM, the invaded (A(t)) drops by 1/3. Nevertheless, stiffening of ECM does hinder invasion but does not give rise to a full arrest. While at slower pace, spheroids still invade and remodel the surrounding matrix. This is in accordance with findings for breast cancer cells in type I collagen [46,48].

2.2. Tumor cells migrate individually

After analyzing the collective invasive motility of the investigated brain cancer tumor spheroids, we tracked the cells that succeeded in detaching from the spheroid or invading cell collections. We followed migration through the ECM for several hours over hundreds of microns and with an algorithm that tracked the center-of-mass of the cells, we distinguished translocation from membrane dynamics. With this large number of data points, even low-probability events are captured.

In Fig. 2A, still images of spheroids are overlaid with examples of cell trajectories (colored lines) for the various ECM concentrations. Cell trajectories are highly diffusive, but in contrast to the invading cell collections, the migrating cells seem indifferent to ECM density at a first glance. This indifference is confirmed by the distributions of displacements, d, between two consecutive frames (5 min) as given by Eq. (5) and shown in Fig. 2B. Because of the heavy tails at longer d, these bar plots cannot be described by a Gaussian distribution as we would expect for random walk models [23].

2.2.1. Super-diffusive migration

To further rectify this, we found the ensemble-averaged mean-squared displacement (MSD) as calculated in Eq. (6) and shown for the cell ensembles in 25% (blue), 50% (red), 67% (yellow), and 75% (purple) ECM concentrations in Fig. 2C. For completeness, mean-squared displacements of the individual cell trajectories for all ECM concentrations can be found in Supplementary figure S1.

The MSDs show anomalous diffusion of the migrating cells characterized by a power law dependence of the time delay, τ:

\[ \text{MSD} \propto \tau^{\alpha}, \]  

(2)

where α is the corresponding anomalous exponent. For normal diffusive motion α = 1. In contrast, anomalous diffusion is either sub-diffusive (α < 1), super-diffusive (1 < α < 2) or ballistic (α = 2). As 1 < α < 2 for all concentrations (inset of Fig. 2C), the migrating cells are super-diffusing which is a sign of active motility and in agreement with prior findings e.g. [49,50]. Thus, the investigated cells appear to overcome the steric hindrances, i.e. the added strength of the physical barrier from increased protein density and decreased pore sizes, imposed by
the increasing concentration of ECM. This is very different from what has been found for lung and bladder cells in collagen [51,52], where concentration increases in collagen caused the cell motility to drop from super- to sub-diffusive motion.

Moreover, the MSDs are almost indistinguishable up until \( t \approx 200 \) min. For the exponent \( \alpha \) of spheroids growing in 25%, 50%, and 75% ECM, the 95% CI's overlap, meaning that the anomalous coefficients are indistinguishable (inset of Fig. 2C). In contrast, the 67% \( \alpha \) is significantly larger than for the other ECM concentrations, hence, cells migrate even more super-diffusively with this mechanical load and level of growth factors etc. Therefore, \( \alpha \) evolves non-linearly with ECM concentration. For even longer lag times, \( t > 1000 \) min, the MSDs exhibit erratic and erroneous behavior due to sparse data.

### 2.2.2. Migration speeds are independent of extra-cellular matrix stiffness

We then investigated how modulating matrix stiffness affects the instantaneous speeds, \( v \), of the individual steps along the migration trajectories; as given in Eq. (8). Box plots of the distributions of \( v \) for the various ECM concentrations are given in Fig. 3A. From these results we concluded that the increased ECM stiffness has no considerable effect on \( v \) for the migrating brain cancer cells. As \( v \) is linearly proportional to the displacements, \( d \), we find the same heavy-tailed distributions as in Fig. 2B, which signifies that cells most often migrate in small steps but once in a while in large leaps. Similar bursts of motion have been observed before [53,54] and, hence, both the frequency and the amplitude contribute to the overall migration speed of the cells.

We also looked into the distributions of the migration speeds, \( \langle v \rangle \), i.e., the average speed over the contour length of the trajectory, as defined in Eq. (9) and box plots of the distributions are shown in Fig. 3B. In contrast to the instantaneous speeds, the migration speeds are normally distributed with the following mean values: \( \langle v \rangle = (1.48 \pm 0.40) \, \mu m/min \) (mean \( \pm SD \), \( N = 223 \)), \( \langle v \rangle = (1.52 \pm 0.54) \, \mu m/min \) (\( N = 252 \)), \( \langle v \rangle = (1.53 \pm 0.57) \, \mu m/min \) (\( N = 123 \)), and \( \langle v \rangle = (1.41 \pm 0.45) \, \mu m/min \) (\( N = 108 \)) in the ECM concentration range from 25% to 75%, respectively. Thus, as verified by a Student's \( t \)-test there are no significant differences between the speed distributions. Hence, the increased ECM concentrations cause no reductions of neither \( \langle v \rangle \) nor \( \langle v \rangle \), indicating a mechanism that allows for overcoming the increased tension of the surrounding matrix. As increasing ECM concentration does not only augment stiffness but also adhesion ligand density and decreases matrix porosity, the migration speed of cells in 3D ECMs is a result of the balance between stiffness, ligand density, proteolysis of the matrix and steric hindrances [39]. For instance, in response to increasing ECM concentration pancreatic cancer has been shown to upregulate the activity of matrix metalloproteinases [55], which are enzymes known to amplify cancer cell migration [56]. Our results are in favor of a similar behavior where migrating brain cancer cells boost their proteolytic activity to overcome the effects of increased matrix rigidity and cell–ECM adhesion as well as reduced pore sizes.

### 2.2.3. Velocity auto-correlation is a double exponential

To further investigate the nature of the motility, we found the velocity auto-correlation function (VACF) from Eq. (10) as presented in Fig. 3C. In general, the mean VACF is – strictly speaking – positive for all ECM concentrations, even at longer time scales. This behavior is consistent with a super-diffusive regime of motion where cells actively project forward in a direction dependent on the previous step. Ergo, the direction of motion is conserved in most cases. Although the correlation decays with time, the cells do not forget in which direction they were heading, i.e., for lag times of \( t < 100 \) min, velocities stay correlated. This memory effect is most pronounced for spheroids in 67% ECM and the least for those in 75% ECM which indicates that the increased matrix stiffness weakens the cells’ ability to keep a consistent direction of motion.
Mammalian cells’ motility has been studied for decades and most classic models are based on the Langevin equation and paths are described as Ornstein–Uhlenbeck processes of the persistent random walk (PRW). Here the MSD depends on diffusion, dimensionality, and the so-called persistence time, \( \tau_p \), i.e. the time that the cell moves persistently in one direction [57], as given in Eq. (12). This model has been found to describe motions of e.g. fibroblasts [18,19], lung epithelial cells [58], and neutrophils [59]. The Ornstein–Uhlenbeck process from Eq. (12) predicts a fast exponential relaxation [17]. In contrast, we found a slower decrease of the VACF for long \( \tau_p \). The decays
of the experimentally observed VACFs are, therefore, characterized by two characteristic relaxation times, $\gamma$ and $\delta$, shown in the insets of Fig. 3C. As expected from the fast decay of 75% ECM (purple line), the $\delta$ is in this case significantly smaller than for 25% and 67% ECM. Such density-dependent decay suggests that augmented ECM density dampens cell motility. Similar double-exponential decay has been found previously in e.g. human fibrosarcoma cell migration in 2D and 3D [26], and in 2D migration of amoebas [49] as well as normal human dermal fibroblasts and human keratinocytes [22], while a gradual transition from exponential to power-law decay was found in mammalian kidney cells [23].

### 2.2.4. Cells migrate both with directionality and persistence

To further investigate the directionality revealed from the VACFs, we studied the angular change between successive steps, $\theta$, as defined in Eq. (11). For a randomly walking cell, we would expect a flat distribution of $\theta$ values, as the cell orient random in all directions. However, Fig. 4A shows that the $\theta$ distributions are skewed towards small angles ($\tau = 5$ min). Therefore, the migrating brain cancer cells exhibit high directional persistence in all ECM concentrations. Fig. 4B shows the distribution of $\theta$s for $\tau$ in the range from 25 to 100 min. The high occurrence of small angles does not flatten with time, even at $\tau = 100$ min. Instead, the motile cells show persistence, which goes directly against the predictions of the PRW model. Furthermore, the probability of observing large angles between two steps increases with increasing time lag. This is indicative of the tendency of the migrating cells to do 180° turns and backtrack; a behavior that is more pronounced in the higher ECM concentrations (67% and 75%), which is in accordance with other observations [26,60,61]. A possible explanation for this is that cells – due to high adhesion and small ECM pore sizes – turn around and follow the beaten micro-tracks in the ECM back towards the spheroid. This asymmetric motility is described by the anisotropic persistence random walk (APRW) model, which encompasses both the high degree of heterogeneity between cells and the anisotropic movements [61].

### 2.2.5. Identifying super-spreaders

Based on these indications, we followed the train of thoughts of Wu et al. [61] to fit the MSDs of individual migrating cells along their trajectory eigenvectors, primary (p) and non-primary (np), with the APRW model from Eq. (14). For each of the migrating cells’ trajectories, this model estimates the persistence time, $\tau_p$, along the primary axis of migration. The distributions of $\tau_p$ shown in Fig. 5A signify stochastic spreading dynamics with long ranged displacements [62]. It has previously been reported that metastatic cancer cells do not only move in a super-diffusive fashion, but also display movement patterns consistent with Lévy walks [15,50]. We, thus, considered both power law (Lévy), exponential (Brownian) as well as log-normal models, which have been used to fit the motion of T-cells within lymph nodes [63].

Fig. 4. Directionality in increased ECM stiffness. (A) Distributions of angles, $\theta$, between two consecutive ($\tau = 5$ min) steps, $r_\tau$, and $r_\tau$, as illustrated in the inset and defined by Eq. (11). The distributions for 25% ($N = 14,630$), 50% ($N = 15,739$), 67% ($N = 6,363$), and 75% ($N = 8,719$) ECM all show that the majority of $\theta$s fall within $|\theta| < 90^\circ$. (B) Angle distributions, $\theta$, for different time delays, $\tau$, from 25 to 100 min, see legend. As the counts vary for different $\tau$’s, occurrences were normalized with respect to the counts for $\theta = 0$.
stay within the field of view more than 10 h after detachment from the spheroid body. As expected, this higher persistence of the super-spreaders is also reflected in the angle distributions. This is apparent from the sub-population medians which become significantly different for all tested ECM concentrations between 25% to 67% when evaluated with a Wilcoxon rank-sum test ($p < 0.001$). For 25% ECM the median $\theta$ is 23° and 33° for high $\tau_p$ and low $\tau_p$, respectively. Likewise for 50%, we found 18° vs. 30° and for 67% 15° vs. 26°. In contrast, for 75% ECM we found a less significant difference between the median $\theta$s, 27° vs. 32° ($p = 0.0002$), suggesting that the super-spreader motions are damped at very high ECM concentrations.

Following the surprising revelation of super-spreaders, we investigated whether this minority population was only characterized by more persistence overcoming the diffusive motion, or whether cells also moved faster along their path. Hence, we re-distributed the migration speeds, $\mathcal{V}$, as shown in Fig. 5C. However, the slight differences we found in average migration speeds, $\mathcal{V}$, were non-significant ($p > 0.05$). This is also apparent from $k$-means clustering of the $\mathcal{V}$ versus $\tau_p$ scatter plots in Supplementary figure S3. So in conclusion, super-spreaders do not move faster — but more persistently.

3. Discussion

Spheroids are known to be very heterogeneous with an outer proliferating zone, an intermediate quiescent region with limited oxygen, nutrients, and metabolites [64,65] as well as a necrotic core [35]; similar to solid in vivo tumors. Furthermore, when suspended in growth medium spheroids grow from the outer proliferating zone. However, the growth cannot be described by a logistic model, rather rates depend on population size [66–69]. When moved from serum-containing growth medium to ECM, our brain cancer spheroids invaded the surrounding matrix by long finger-like protrusions. This is a sign of collective migratory behavior where multicellular groups infiltrate the ECM through gained motility and retained cell–cell adhesion [70]. These findings are paralleling earlier findings for similar spheroids (U87-MG) [35,47] as well as for other glioblastoma spheroids (KNS42) [35], glioma spheroids (U-251-MG) [71], and breast carcinoma spheroids [35,72]. This infiltrative nature makes it impossible to eliminate all cells in a host by surgery, thus leading to tumor recurrence.

From the acquired time-lapse sequences of invading spheroids, we found that the cross sectional area of the invading spheroid grows exponentially with time with no slowing of the growth rate within 24 h. Nevertheless, a gradual augmentation of the ECM (Matrigel™) concentration slows down invasion. Thus, the collective motility of cells in the finger-like protrusions responds to the polymer concentration in the matrix. However, increased ECM concentration will necessarily also increase the concentration of individual ECM components such as growth factors (IGF, EGF, PDGF etc.) which activate and sustain cancer proliferation and growth [73,74]. Naively, one could therefore think that increased Matrigel™ concentration would augment the invasive capabilities of the spheroid. However, this augmentation is only observed when comparing ECM-embedded spheroids to spheroids in growth medium alone (0% ECM). In contrast, increasing the ECM concentration from 25% to 75% decreased spheroid invasion. One possible explanation is mechanosensing as several studies have shown that increased matrix stiffness triggers mechanosensitive transcriptional activators such as YAP and TAZ, which in turn activate e.g. proliferation, differentiation and metastasis in the tumor [75–78]. However, increased matrix stiffness is accompanied by a decrease in matrix porosity which limits the ability of cell collections to move through the matrix [11]. Like so many other functions in the tumor [79],
these mechanical cues might also be conveyed through YAP/TAZ and favor the full transformation into individual migrating cells rather than invasive cell collections. Consequently, the mechanical stiffness of the tumor microenvironment not only triggers spheroid invasion, but also regulates it effectively through mechanical feedback [80].

We further detected how individual cells detached from the spheroid – or the neighboring cells in the protrusion – to start individual migration through the ECM. This is indicative of a complete epithelial to mesenchymal transition, EMT, of the cells associated with weakening of the intercellular adhesion, loss of polarity, and increased motility [81,82]. The cells were observed to first extend their pseudopodia, then displace the cell body followed by parallel retraction of pseudopodia. This behavior has been observed before in migrating glioblastoma cells [54] and in the highly invasive human breast carcinoma cells (MDA-MB-231) [53]. Particularly, bursts of speed were pronounced when cells escaped from areas crowded by non-transformed breast epithelium cells (MCF10 A). Moreover it was suggested that such fast movements contribute to the overall migration speed of each cell and they are the reason for their effective migration [54].

Surprisingly, we found that glioblastoma cells exhibit similar migratory behaviors in all tested ECM concentrations. In contrast, if mechanosensing causes a shift from invasion towards migration as hypothesized above, we would expect an increase in the overall migration speeds or diffusion constants with ECM concentration. The notion that increased stiffness increases cancer cell migration due to YAP/TAZ activation supports this argument. However, Matrigel™ has been found to play a dual role: While the presence of Matrigel™ in collagen gels promoted migration of lung cancer cells (H1299) at low concentration, it hindered migration at higher concentrations, probably due to increased adhesion-ligand concentration [11] and increased elastic modulus [40]. Thus, our hypothesis – that increased ECM concentration favors increased single cell migration – might be counteracted by increased adhesion-ligand concentration and result in no apparent ECM concentration dependence [60]. Moreover, earlier studies suggest that proteolysis serves as an important additional mechanism for migration of cancer cells, allowing them to move through the matrix — no matter how dense it is [39,56,83]. Further, it has been established that during collective cell invasion, the leading edge cells carve paths for the remaining cells by proteolytic remodeling of the ECM [84,85]. For example for fibrosarcoma cells (HT-1080) invading 3D collagen, leader cells utilize proteolysis by a membrane-bound collagenease (MT1-MMP) to generate tracks [86]. Our detection of two different migrating populations is indicative of a similar leader/follower relationship for migrating cells: leader cells that proteolytically generate paths for subsequent cells to follow. However, we could not detect such a relationship between super spreaders and other migrating cells.

Additionally, tumor cells can switch between mesenchymal and amoeboid migration modes and thereby increase invasive capabilities [87]. Amoeboid migration is characterized by e.g. fast speeds and high levels of actomyosin contractility [88] and have also been reported for U87-MG in 3D hydrogels [89]. To decide whether the super spreading population found in this study is amoeboid will demand further investigations, e.g., of membrane flows [90].

Further, it has been shown that collagenase activity in pancreatic cancer cells (Panc-1) increases with matrix stiffness [55]. This indicates that the cells have mechanosensing abilities that cause them to increase proteolytic activity in order to overcome increased matrix stiffness, thus apparently rendering migration of U87-MG glioblastoma cells ECM-concentration independent. Nevertheless, we did observe slight concentration dependencies, such as decreased VACF decay times and increased probability of backtracking at long time scales in high ECM concentrations.

Any Brownian random walk model of cell migration would predict a uniform distribution of turning angles between successive steps, while the PRW model predicts a peaked distribution that flattens for time lags longer than the persistence time. In contrast to both these random walk models, we found that the probability of complete 180° turnarounds in the 3D matrix increased with time. This result indicates that the probability of observing cells backtracking through the tunnels – formed by the cells during their initial exploration – increases at longer time scales. Given such anisotropic persistence, we fitted our results to the APRW model and extracted the according persistence time distributions for the primary directions of migration. We found long-range displacement dynamics interspersed between long periods of short-range steps, which is characteristic of Lévy walks. Furthermore, a recent study proved that the motion of metastatic cancer cell lines performed Lévy walks, while their non-metastatic counterparts did not [50]. As Lévy walks also have uniformly distributed turning angles, the arguments for describing the motion of migrating glioblastoma cells as a Lévy walk were, hence, very compelling. However, our persistence time distributions did not follow a power-law. Instead we – unexpectedly – found the persistence time distributions to be log-normal and bimodal. This is suggestive of the existence of two subpopulations of which one is a population of super-spreaders with boosted persistence. Such contradictory observations have been done previously, e.g., for immune T-cell motion, which has been identified as either log-normal [63] or a Lévy walk [91] depending on the tissue they migrated in. Moreover, evidence that individual persistent random walkers exhibit Lévy walk movement patterns at the ensemble level has been published in the case of single bacteria vs. swarms [92,93]. Evidently, the distinction between variants of PRWs and Lévy walks warrants further investigation [94,95].

Glioblastoma is a particular fast-spreading form of brain cancer [96], due to its high plasticity, heterogeneity and therapy resistance [97]. Understanding metastasis and migration in these malignant cells is a critical step towards better prognoses. As the focus of this study is to compare uniform spheroids under different conditions, we used an immortalized human tumor-derived cell line. However, often long-term cultivation of tumor cell lines results in loss of genetic heterogeneity of the original tumor. In particular, the cell line used in this study (U87-MG) was recently found to be genetically different from its tumor of origin [98]. Therefore to extend the validity, a follow-up study should include the use of spheroids that are either derived from human glioma xenografts [99] or built as multi-cell type glioma spheroids [100].

In summary, we analyzed cell migration from glioblastoma spheroids embedded in varying concentrations of ECM-like Matrigel™. While collective migration was partially inhibited by high-density gels, single cell migration was not. However, migration of single cells did become more anisotropic with increasing ECM density. Modeling the migration as an APRW, we identified a subpopulation of super-spreading cells with extraordinary directional persistence. Development of a model that captures all the behaviors we have observed – namely double-exponential decay of the VACF, highly directional and persistent migration, anisotropic behavior at high ECM density and the existence of super-spreading cells – would allow prediction of the motility mode of the migrating cells further from the spheroid. This would be an important step en route to a unifying model of cancer metastasis. Moreover, although the concept of intra-tumoral heterogeneity has been established years ago, it is still an area of research that receives much attention. Our work presents yet another aspect of between-cell tumor heterogeneity with high clinical relevance since targeting the population of super-spreading cells would be the most efficient methodology for prevention of metastasis and relapse after surgery. Future studies will have to uncover whether our observations represent actual differences between cells within the tumor spheroids or whether they arise in response to different local environments.

4. Materials and methods

4.1. Cell culture

The glioblastoma multiforme cell line U87-MG was cultured under standard conditions (37 °C, 5% CO₂, 95% humidity) in Dulbecco’s
Modified Eagle Medium (DMEM) supplemented with 10% Fetal Bovine Serum (FBS) and 1% penicillin–streptomycin; all provided from Gibco (Gibco, Life Technologies Ltd. Paisley, UK).

4.1. Gravitational-assisted tumor spheroid formation

For the formation of spheroids, 650 cells (3250 cells/mL) were seeded in ultra-low attachment round-bottomed 96-well plates (Corning B.V. Life Sciences, Amsterdam, The Netherlands) with 200 μL DMEM and incubated for 4 days [37].

4.2. Spheroid invasion/migration

Our invasion assay was based on classic protocols described by Vinci et al. [35] and Richards et al. [101]. Matrigel™ (Corning B.V. Life Sciences, Amsterdam, Netherlands) was thawed on ice overnight at 4 °C. On the day of experimentation, the round-bottomed well plates with fully-formed spheroids were placed on ice to avoid untimely Matrigel™ solidification. For experimentation, we varied the concentration of Matrigel in medium (v/v) between 0%, 25%, 50%, 67%, and 75%. For this aim, medium was removed from the wells and replaced with ice-cold Matrigel™ (using pipette tips kept at −20°C) to the desired final concentration. When interchanging medium with Matrigel™ we took care to keep the spheroids centered in the bottom of the well and to remove all bubbles. Afterwards, plates were incubated for 1 h for solidification of the Matrigel™ and lastly 100 μL DMEM was added on top of each well to a final volume of 300 μL.

4.2. Time-lapse imaging

96-well plates were mounted on an inverted Juki Stage Real-Time cell history recorder (NanoEntek, Guro-gu, Seoul, Korea) with bright-field (10× objective) placed inside a cell incubator (37 °C, 5% CO₂, 95% humidity). The focus of the microscope was set manually for each well and we used the built-in software to acquire time-lapses with a frame rate of 1 every 5 min, exposure time of 170 ms, and field of view (105×640) μm² with a resolution of 440 nm/pixel. For 24 or 31 h of imaging, 50 μL of DMEM was added to the wells during experimentation to prevent dehydration of wells. Please, note that our imaging modality results in a 2D approximation of the 3D migration as discussed in detail by Wu et al. [26].

4.3. Image analysis

4.3.1. Spheroid invasion

To quantify spheroid invasion we used the Fiji - ImageJ [102] macro tool Analyze Spheroid Cell Invasion In 3D Matrix to obtain the projected area of the main spheroid body [103]. From the resulting cross sectional area, A, we found the average area of all spheroids to be:

$$\langle A(t) \rangle = \frac{1}{N} \sum_{n=1}^{N} A_{{n}(t)} \tag{3}$$

where $A_{{n}(t)}$ is the area of the nth tumor spheroid at time $t$ and $N$ is the total number of spheroids.

4.3.2. Image pre-processing

We used the open-source software ilastik, which enables automated – albeit supervised by the user – segmentation of spheroids and individual cells [36]. For training of the algorithm, we used a representative stack of 50 consecutive images from 0 to 250 min to minimize computation time. Please note that some manual track-editing was needed for our data because of over-segmentation, i.e., one cell is detected as two, and under-segmentation (vice versa). We also had to remove data points from objects (of size similar to cells) that mistakenly had been detected as cells. All these errors were filtered manually to ensure meaningful results.

4.3.3. Single cell migration

For automated single particle tracking we used the TrackMate plugin for Fiji - ImageJ, which consists of a spot detection followed by a particle tracking procedure. Since our cells expand over a rather large area in the segmented image (≥30 pixels ~ 9 μm), we used the Downsample LoG detector [104] with the down sample factor set to 4, the estimated blob diameter to 50 μm, and a cell/background threshold value of 1 appropriate for our binary images. With these settings, the spot detection algorithm detected all centers of mass of migrating cells as well as cells on the spheroid surface. Once spot detection is completed, cells were tracked in every frame of the image stack using the The Simple Linear Assignment Problem algorithm since – upon visual inspection of the images – mitosis events were rare within our time frame. For post-processing, we used the TrackScheme module to edit the cell trajectories and we ended a track in the following cases: (i) if a cell moved out of the field of view (even if it returned later), (ii) if part of the cell left the field of view for more than 15 min, (iii) if cells never really detached from the spheroid before returning, (iv) after cell division [105].

4.4. Data analysis

4.4.1. Displacement analysis

An ensemble of particles undergoing Brownian motion is distributed as [106]:

$$P(r, t) = \frac{1}{\sqrt{4\pi Dt}} e^{-\frac{r^2}{4Dt}} \tag{4}$$

where $D$ is the diffusion coefficient and $t$ is the time step. In the $n$th cell trajectory we define a position vector in the projected plane as: $r^{(n)}(t)$. The length of the vector is the displacement, $d$:

$$d = |r^{(n)}(t) - r^{(n)}(t - 1)| \tag{5}$$

4.4.2. Mean squared displacement analysis

The mean squared displacement (MSD) describes the displacement of cells by determining the ensemble-averaged distance that the ensemble of cells move in a given time interval, $r$, which is designated as the lag or delay time [107]:

$$\text{MSD} \equiv \langle |r(r) - r(0)|^2 \rangle = \frac{1}{N} \sum_{n=1}^{N} |r^{(n)}(r) - r^{(n)}(0)|^2 \tag{6}$$

where $N$ is the total number of trajectories for all measured spheroids, vector $r^{(n)}(0)$ is the starting position of the $n$th cell trajectory, and vector $r^{(n)}(r)$ is the position of the $n$th cell after time lag $r$.

The MSD of a random walk, i.e., normal diffusion, of cells in 3D scales linearly with time:

$$\text{MSD} = 6D r \tag{7}$$

4.4.3. Instantaneous speed distributions and migration speed

We defined the instantaneous speed, $v$, at the $i$th step of a cell as the displacement in the projected $x$, $y$-plane per time step:

$$v = |\dot{r}(i)| = \frac{|r(i) - r(i-1)|}{t_i - t_{i-1}} \tag{8}$$

and the migration speed of a cell, $\bar{v}$, as the mean over the entire contour length of the trajectory from $i = 1$ to the total number of steps, $i = L$:

$$\bar{v} = \frac{1}{L} \sum_{i=1}^{L} v_i \tag{9}$$
4.4.4. Motion persistence and directionality

We also investigated the velocity auto-correlation function (VACF) of the ensemble average of cell trajectories which is:

\[ \text{VACF} = \langle v(0) \cdot v(\tau) \rangle = \frac{1}{N} \sum_{i=1}^{N} v^{(i)}(0) \cdot v^{(i)}(\tau), \]  

(10)

where the velocity at time \( \tau \) is \( v^{(i)}(\tau) \) of the \( i \)th cell trajectory and the initial velocity is \( v^{(i)}(0) \). In general, the VACF diminishes with increased \( \tau \), as the cells’ velocities become uncorrelated due to interactions with the surrounding environment. Furthermore, the angle between the position vectors between two successive steps \( r_{(i)} \) and \( r_{(i-1)} \) fulfill the relation:

\[ \tan(\theta) = \frac{\| r_{(i)} \times r_{(i-1)} \|}{r_{(i)} \cdot r_{(i-1)}} \]  

(11)

4.4.5. The Ornstein–Uhlenbeck process

The Ornstein–Uhlenbeck process describes a random walk model but with an added persistence (PRW) [22]. This model of cell motility is derived from a stochastic differential equation describing the motion of a self-propelled cell. By introducing persistence, the MSD in 3D as given in Eq. (7) for a simple random walk is described by Fürth’s formula [24]:

\[ \text{MSD} = 6D \left( t - \tau_p \left( 1 - e^{-\tau_p/\tau_f} \right) \right). \]  

(12)

where \( \tau_p \) is the persistence time of the motion, i.e., the time for which a certain velocity is retained by the system [17]. Therefore, \( \tau_p \) is the time before a cell changes direction. Eq. (12) can be rewritten as:

\[ \text{MSD} = 2r^2\beta^2 \left( e^{-\tau_p/\tau_f} + \frac{\tau_f}{\tau_p} - 1 \right) \]  

(13)

if we assume an exponential distribution of persistence times [108]. Furthermore, the Ornstein–Uhlenbeck velocity auto-correlation function, VACF, which can be derived by differentiating equation (12), is predicted to be a single-exponential decay.

4.4.6. Anisotropic persistent random walk model

We have adopted the anisotropic persistent random walk model (APRW) for the migration of cells in a 3D matrix [26,61]. In the APRW, the velocity of the migrating cells is assumed to be spatially anisotropic and display different persistence times on two orthogonal axes, namely the primary \( p \) migration axis and the non-primary \( np \) migration axis; these are found through singular vector decomposition of cell velocities.

The migration of the cells in the APRW model is described by the same parameters as in the PRW model, the persistent time \( \tau_p \) and cell speed \( V_p \), however they are different for each individual axis of motion. Therefore, for the primary axis they are denoted as \( (V_p, \tau_p) \) and for the non-primary they are \( (V_{np}, \tau_{np}) \). The MSDs in the primary direction, \( p \), are given by:

\[ \text{MSD}_p = \frac{1}{N_p} \sum_{i=1}^{N_p} v^2_{(i)} \left( e^{-\tau_p/\tau_f} + \frac{\tau_f}{\tau_p} - 1 \right) + 2\sigma^2 \]  

(14)

and in the non-primary orthogonal direction \( np \):

\[ \text{MSD}_{np} = \frac{1}{N_{np}} \sum_{i=1}^{N_{np}} v^2_{(i)} \left( e^{-\tau_{np}/\tau_f} + \frac{\tau_f}{\tau_{np}} - 1 \right) + 2\sigma^2, \]  

(15)

where \( 2\sigma^2 \) is the error in the position of the cell and the total MSD for each cell is given by:

\[ \text{MSD} = \text{MSD}_p + \text{MSD}_{np}. \]  

(16)

4.5. Statistical analysis

When examining migration characteristics for different Matrigel™ concentrations, more than 100 cell trajectories originating from at least 4 different spheroids were evaluated for each condition (corresponding to >6,000 steps).

4.5.1. MSD curve fitting

For all MSD analysis of trajectories, the small time delays, \( \tau \), are weighted more in the average MSD than longer delays. For example, if a trajectory has \( N \) points, the delay, \( \tau \), that is equal to 1 time step will have \( N - 1 \) points in the average, whereas the delay that corresponds to \( N \) time units will have only 1 point in the average [109]. Therefore, the accuracy of the MSD becomes increasingly worse with \( \tau \) as the variance depends on the number of trajectories available. Moreover, the curvature of the MSD is important when investigating whether there is active transport in the system under study. In the ensemble-averaged MSD, there are contributions from different kinds of motion which makes the curvature change over time thus increasing statistical uncertainty. Therefore, to limit the relative errors of the MSD, the plots should not be analyzed and fitted to any model further than 25% of the longest time lag. Furthermore, when fitting an MSD to a model, one should use proper weights, such as the inverse of the variance or standard deviation for each time lag, in order to give more weight to the parts of the MSD with the highest certainty [16,110]. Both suggestions were adopted in this study.

From curve fitting of the various models to the MSD or VACF, the coefficients \( \alpha, \beta, \gamma, \) and \( \delta \) are obtained and given (in Figs. 1, 2, and 3) with error bars, which corresponds to the 95% confidence intervals (CI).

Furthermore, when analyzing MSD data, there are two possible approaches: (i) to truncate all trajectories to the same length (number of time steps) or (ii) to truncate all trajectories to the same distance from the spheroid (cut of by field of view). The advantage of the first approach is that there is no oversampling of the slow trajectories, which stay longer in the field of view. The strength of the second approach, is better statistics. For this analysis we used (ii) but we found that when using (i), the second populations of Fig. 5A grow substantially [111]. Specifically for this analysis, we only registered the \( \tau_p \)'s for fits with the coefficient of determination \( R^2 > 95\% \).

4.5.2. Confidence and accuracy

Throughout the present study we state data as mean ± standard deviation (SD). However, for the investigation of spheroid invasion (Fig. 1) we presented the mean ± standard error of the mean (SEM) and for the persistence analysis (Fig. 5) we used the multiplicative standard deviation (SD); as detailed in the text.

For the box plots in Figs. 3 and 5 the red line signifies the ensemble average, the pink box represents the 95% confidence interval (CI), and the edges of the colored boxes are one standard deviation (SD).

CRediT authorship contribution statement

Aimilia Nousi: Spheroid imaging and single-particle tracking.

Maria Tangen Søgaard: Supportive experiments. Mélanie Audoin: Supportive data analysis. Liselotte Jauffred: Supervised the project.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Data availability statements

All cell trajectories – as spatial coordinates and time – presented in this study are openly available in FigShare at doi:10.6084/m9.figshare.14229515.
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