Far from equilibrium dynamics of tracer particles embedded in a growing multicellular spheroid

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Local stresses on the cancer cells (CCs) have been measured by embedding inert tracer particles (TPs) in a growing multicellular spheroid. The utility of the experiments requires that the TPs do not alter the CC microenvironment. We show, using theory and extensive simulations, that proliferation and apoptosis of the CCs, drive the dynamics of the TPs far from equilibrium. On times less than the CC division times, the TPs exhibit sub-diffusive behavior (the mean square displacement, $\Delta_{TP}(t) \sim t^{\beta_{TP}}$ with $\beta_{TP} < 1$). Surprisingly, in the long-time limit, the motion of the TPs is hyper-diffusive ($\Delta_{TP}(t) \sim t^{\alpha_{TP}}$ with $\alpha_{TP} > 2$) due to persistent directed motion for a number of CC division times. In contrast, CC proliferation randomizes their motion resulting from jamming at short times to super-diffusive behavior, with $\alpha_{CC}$ exceeding unity, at long times. Surprisingly, the effect of the TPs on CC dynamics and radial pressure is negligible, suggesting that the TPs are reliable reporters of the CC microenvironment.

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The interplay between short-range forces and non-equilibrium processes arising from cell division and apoptosis gives rise to unexpected dynamics in the collective migration of cancer cells [1–7]. An example is the invasion of cancer cells (CCs) in a growing multicellular spheroid (MCS), which is relevant in cancer metastasis [6, 7]. Imaging experiments show that collective migration of a group of cells that maintain contact for a long period of time exhibits far from equilibrium characteristics [7–11]. Simulations and theory explain some of the experimental observations [12–14]. Dynamics in a growing MCS is reminiscent of the influence of active forces in abiotic systems [15–17]. In a growing MCS the analogue of active forces are self-generated [18], arising from biological events characterized here by cell division and apoptosis.

Experiments that probe the local stresses or pressure on the CCs [19–24] have provided insights into the mechanism by which the CCs invade the extracellular matrix. Recently, the stresses within MCSs were measured [24] by embedding micron-sized inert deformable polyacrylamide beads as probes. For a TP to be a reliable sensor, it should not alter the dynamics of the CCs and the microenvironment, as assessed by local pressure. However, the influence of TPs on the CCs is unknown – a gap that we fill here.

We used theory and simulations to probe the dynamics of the TPs in the MCS, and the impact of the TPs on the CC mobility as well as radial pressure. The central results are: (1) The TPs exhibit sub-diffusive motion in the intermediate time scale ($t \lesssim \frac{1}{k_5}$), with the mean squared displacement, $\Delta_{TP}(t) \sim t^{\beta_{TP}}$, with $\beta_{TP}$ less than unity. In the long time limit, ($t \gtrsim \frac{1}{k_5}$), the TPs undergo hyper-diffusive motion, $\Delta_{TP}(t) \sim t^{\alpha_{TP}}$ with $\alpha_{TP}$ greater than 2. In contrast, the CCs, which are jammed at $t \lesssim \frac{1}{k_3}$, exhibit super-diffusive dynamics. (2) The dynamics of the CCs in the intermediate time regime changes slightly as the size of the TPs are varied. Remarkably, the long time exponent ($\alpha_{CC}$) as well as the decrease in the pressure as a function of the distance from the center of the MCS remains unaffected in the presence of TPs.

Theory: We consider the dynamics of the TPs in a growing MCS in a dissipative environment by neglecting inertial effects. The deformable but inert TPs, do not grow, divide or undergo apoptosis. The CCs grow and divide at a given rate, and undergo apoptosis (see Figure 1). The CCs and TPs experience systematic short-ranged attractive and repulsive forces arising from other CCs and TPs [12]. To obtain the dynamics of the TPs and CCs, we introduce the density fields $\phi(r,t) = \sum_i \delta[r - r_i(t)]$ for the CCs, and $\psi(r,t) = \sum_i \delta[r - r_i(t)]$ for the TPs. A formally exact Langevin equation for $\phi(r,t)$, and $\psi(r,t)$ may be derived using the Dean’s method [25] accounting for diffusion and non-linear interactions.

Figure 1: Difference between the CCs (pink sphere) and the TPs (magenta sphere). Cell division creates two daughter CCs (blue and green sphere), each with $R_{d} = \frac{R_{m}}{2^{1/3}}$. Their relative position is displaced by $2R_{m}(1 - \frac{1}{2^{1/3}})$ with the orientation being random with respect to the sphere center.
To model a growing MCS, we modify the density equation for the CCs phenomenologically by adding a non-linear source term, \( \propto \phi(\phi_{\text{th}} - \phi) \), accounting for cell birth and apoptosis, and a non-equilibrium noise term that breaks the CC number conservation. The noise, \( f_{\phi} \), satisfies \( f_{\phi}(r, t) f_{\phi}(r', t') = \delta(r-r')\delta(t-t') \). The source term, \( \propto \phi(\phi_{\text{th}} - \phi) \), represents the birth and apoptosis, with \( \phi_{\text{th}} = \frac{\beta_{\text{TP}}}{k} \) [26, 27]. The coefficient \( \sqrt{k_{b}\phi + \frac{k_{b}}{2}\phi^{2}} \) of \( f_{\phi} \) is the noise strength, corresponding to number fluctuations.

The \( \phi(r, t) \) and \( \psi(r, t) \) fields obey,
\[
\begin{align*}
\frac{\partial \phi(r, t)}{\partial t} &= D_{\phi} \nabla^{2} \phi(r, t) + \nabla \cdot (\psi(r, t)J) + \eta_{\phi}, \quad (1) \\
\frac{\partial \psi(r, t)}{\partial t} &= D_{\psi} \nabla^{2} \psi(r, t) + \nabla \cdot (\phi(r, t)J) + \\
&= \frac{k_{a}}{2} \phi \left( \frac{2k_{b}}{k_{a}} - \phi \right) + \sqrt{k_{b}\phi + \frac{k_{b}}{2}\phi^{2}} f_{\phi} + \tilde{\eta}_{\psi}(2)
\end{align*}
\]
where \( J = \int \psi(r', t) \phi(r', t) \nabla U(r - r') \), \( \tilde{\eta}_{\phi}(r, t) = \nabla \cdot (\eta_{\psi}(r, t) \psi^{1/2}(r, t)) \), \( \tilde{\eta}_{\psi}(r, t) = \nabla \cdot (\eta_{\psi}(r, t) \phi^{1/2}(r, t)) \), and \( \eta_{\phi, \psi} \) satisfies \( \eta_{\phi, \psi}(r, t) \eta_{\phi, \psi}(r', t') = \delta(r-r')\delta(t-t') \). The second term in Eq.(1) accounts for the TP-TP interactions \( (\nabla \cdot (\psi(r, t) J) \nabla U(r - r')) \) and TP-CC interactions \( (\nabla \cdot (\psi(r, t) \int \phi(r', t) \nabla U(r - r')))) \). The influence of the CCs on the TP dynamics is reflected in the TP-CC coupling. The third term in Eq.(2) results from cell birth and apoptosis, and the fourth term in Eq.(2) is the non-equilibrium noise. Eq. 1 does not satisfy the fluctuation-dissipation theorem in the tumor growth phase.

To obtain the exponents describing the dependence of the mean-square displacement (MSD) on time, we consider a change in scale, \( r \rightarrow s r, \) and \( t \rightarrow s^{2} t \) where \( s \) is the dynamic exponent. We use the Parisi-Wu [28-30] technique to calculate \( s \). The main features of the method, which are not needed to understand the central results here, are relegated to section I in the SI.

Simulations: We simulated a 3D MCS with embedded TPs using an agent-based model [12, 14, 31]. The cells, treated as interacting soft deformable spheres, grow with time, and divide into two identical daughter cells upon reaching a critical radius \( (R_{m}) \). The mean cell cycle time, \( \tau = \frac{1}{\beta} \), is 15 hrs is used if not mentioned explicitly. The CCs also undergo apoptosis at the rate \( k_{a} \ll k_{b} \). The sizes and the number of TPs are held constant.

To determine the TP dynamics, we model the CC-CC, CC-TP, and TP-TP interactions using two potentials (Gaussian and Hertz) to ensure that the qualitative results are robust (simulation details are in Sec II in the SI).

The equation of motion for the dynamics of TP and CCs is taken to be [12], \( \hat{r}_{i} = \frac{\nabla\psi}{m} + \gamma_{i} \hat{v}_{i} + F_{i} \), where \( \hat{r}_{i} \) is the velocity of \( i^{th} \) CC or TP, \( \hat{v}_{i} \) is the force on \( i^{th} \) CC/TP, and \( \gamma_{i} \) is the damping term. We used a pressure inhibition mechanism to model the observed growth dynamics in solid tumors [12]. Dormancy or the growth phase of the CCs depends on the local microenvironment, determined by the pressure on the \( i^{th} \) cell (Figure 1). Cell division and the placement of the daughter cells add stochasticity in the dynamics of the CCs. We initiated the simulations with 100 TPs and 100 CCs. The spatial coordinates of the CCs and TPs were sampled using a normal distribution with mean zero and standard deviation, 50 \( \mu m \).

Results: Theory and simulations predict that, in the limit \( t < \frac{1}{\beta \tau} \), the TPs exhibit sub-diffusive behavior (Table I). The non-linear terms for the TP-TP and TP-CC interactions determine the scaling laws. A power counting analysis shows that the dynamical exponent \( z = 2 + \frac{d}{2} \). As a consequence, the MSD of a TP behaves as, \( \Delta(t) = \sum [r(t) - r(0)]}^{2} \sim t^{2z/2} = t^{2/2} \). Because \( \beta_{TP} \) is less than unity we surmise that the TPs are jammed. We also calculated \( \Delta(t) = \langle [r(t) - r(0)]^{2} \rangle \), by averaging over \( \approx 2,000 \) trajectories, for the Gaussian (Figure 2) and Hertz (Figure S2a in the SI) potentials. Similar behavior is found in both cases. The duration of the plateau (Figure 2) increases as the cell cycle time increases. Although jamming behavior predicted theoretically is consistent with the simulations the \( \beta_{TP} \) ex-

| Theory | Hertz | Gaussian |
|--------|-------|----------|
| \( \beta_{TP} \) | 0.57 | 0.12 | 0.11 |
| \( \alpha_{TP} \) | 2.28 | 2.30 | 2.30 |
| \( \alpha_{CC} \) | 1.45 | 1.47 | 1.50 |

Figure 2: MSD of the TPs (\( \Delta_{TP} \) using the Gaussian potential. The curves are for 3 cell cycle times (blue (0.5\( \tau \)), red (\( \tau \)), and green (2\( \tau \))). Time to reach the hyper-diffusive behavior, which is preceded by a jamming regime (\( \Delta_{TP} \sim t^{2/2}, \beta_{TP}=0.11, \) shown in black dashed line), increases with \( \tau \). The inset focusses on the hyper-diffusive regime (\( \frac{d}{2} > 1 \)). \( \tilde{\tau} \) is the cell cycle time for the respective curves. Time is scaled by \( \frac{1}{\tau} \). \( \Delta_{TP} \sim t^{2/2}, \alpha_{TP} = 2.3 \) (dashed black line).
ponents differ (Table I).

In the $t > \frac{1}{\tau}$ limit, theory and simulations predict hyper-diffusive dynamics, $\Delta_{TP} \sim t^{\alpha_{TP}}$ with $\alpha_{TP} > 2$. Non-linearities, due to the TP-CC interactions together with cell division and apoptosis, determine the long-time behavior of the TPs. Using the scaling analysis as before, the theory predicts hyper-diffusive dynamics for the TPs (Table I). Variations in the theory predicts hyper-diffusive dynamics for the TPs (Table I). Variations in $\alpha_{CC} > 1$, $r_{TP} = 2r_{CC}$, and $r_{TP} = r_{CC}$, and red - $r_{TP} = 0.5r_{CC}$, and blue - $r_{TP} = 0.5r_{CC}$, where $r_{CC} = 4.5\mu m$ is the average cell radius. Inset shows $\Delta_{CC}$ for the four curves, focusing on the super-diffusive regime. The black dashed line is drawn with $\alpha_{CC} = 1.47$.

![Figure 3](image-url): CC dynamics in the presence of the TPs. $\Delta_{CC}$ using the Hertz potential. The curves are for different TP radius ($r_{TP}$) and $r_{CC}$, green - $r_{TP} = r_{CC}$, and red - $r_{TP} = 0.5r_{CC}$, and blue - $r_{TP} = 0.5r_{CC}$, where $r_{CC} = 4.5\mu m$ is the average cell radius. Inset shows $\Delta_{CC}$ for the four curves, focusing on the super-diffusive regime. The black dashed line is drawn with $\alpha_{CC} = 1.47$.

Figure 4: (a) CC correlation function, $(g(r))$, as a function of inter-cellular distance. Red (blue) curve shows $g(r)$ in presence (absence) of TPs. The two dashed lines (black and orange) are power law fits to $g(r)$ in the large $r$ limit. (b) Force force autocorrelation (FFA), as a function of delay time ($t_d$). The red (blue) curve shows the FFA for TPs (CCs). Inset shows FFA on log-log scale. The black (yellow) dashed line is a power law fit with exponent of $-0.7$ (-1.2).

$g(r) = \frac{\nu}{4\pi r^2 N^2} \sum_{i=1}^{N} \sum_{j \neq i}^{N} \delta(r - |r_i - r_j|) \sim r^{-0.5}$, in the presence and absence of TPs at $t \approx 8\tau$ (Figure 4a). The dynamically-induced CC correlations is independent of the TPs, thus explaining the insignificant effect of the TPs on the CC dynamics.

To explain the finding, $\alpha_{TP} < \alpha_{CC}$, we calculated the force-force autocorrelation function, FFA=\langle F(t+t_d) \cdot F(t) \rangle. Here, $\langle ... \rangle$ is the time average and $t_d$ is a delay time. Since, the TPs (CCs) exhibit hyper-diffusion (super-diffusion), we expect that the FFA of TPs should decay slower relative to the CCs, which is confirmed in Figure 4b, which shows that the FFA for the TPs (CCs) decays as $t_d^{-0.7}$. Thus, the TP motion is significantly more persistent than the CCs, which explains the
We define \( \cos(\theta) \) as the angle between the dynamics of the TPs and CCs, we calculated by measuring the extent of skewness for both the CCs and TPs. The data analysis revealed that the distribution of \( \cos(\theta) \) is skewed towards \( \cos(\theta) > 0 \), indicative of persistent motion. The data for the TPs is skewed to \( \cos(\theta) > 0 \), whereas for the CCs the distribution is skewed to \( \cos(\theta) < 0 \). We conclude with a few comments. (i) The excellent agreement for \( \alpha_{TP} \) between simulations and theory suggests that the CC dynamics is determined by the overall tumor growth, and not the details of the TP-CC interactions. Although \( \alpha_{TP} > \alpha_{CC} \), the CCs move a larger distance than the TPs (compare Figure 2 and Figure 3), which implies that the effective diffusion constant of the CCs is greater than the TPs. (ii) The radial dependence of the pressure (Figure 5b), with the core experiencing higher values than the periphery cells, is another manifestation of the pressure profile found in the experiments. Pressure decreases roughly by factor of four, as the distance \( r \) from the center of the tumor increases. The pressure is almost constant in the core, with a decrease that can be fit using the logistic function, as the boundary of the tumor is reached (Figure (5b). The high core pressure is due to small number of cell divisions. As a consequence, the CCs are jammed, leading to high internal pressure. As \( r \) increases, the CCs proliferate, resulting in a decrease in self generated stress, and consequently a decreases in the pressure (Figure (5b).(ii) Most importantly, the CC pressure profiles are unaltered even in the presence of the TPs (compare blue circles and greenish yellow squares in Figure (5b), which shows that the latter does faithfully report the microenvironment of the CCs.

We also calculated pressure experienced just by the TPs. As \( \alpha_{TP} > \alpha_{CC} \), the CCs move a larger distance than the TPs (compare Figure 2 and Figure 3), which implies that the effective diffusion constant of the CCs is greater than the TPs. (ii) The radial dependence of the pressure (Figure 5b), with the core experiencing higher values than the periphery cells, is another manifestation of the pressure profile found in the experiments. Pressure decreases roughly by factor of four, as the distance \( r \) from the center of the tumor increases. The pressure is almost constant in the core, with a decrease that can be fit using the logistic function, as the boundary of the tumor is reached (Figure (5b)). The high core pressure is due to small number of cell divisions. As a consequence, the CCs are jammed, leading to high internal pressure. As \( r \) increases, the CCs proliferate, resulting in a decrease in self generated stress, and consequently a decreases in the pressure (Figure (5b).(ii) Most importantly, the CC pressure profiles are unaltered even in the presence of the TPs (compare blue circles and greenish yellow squares in Figure (5b), which shows that the latter does faithfully report the microenvironment of the CCs.

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