1 The Complex Ecosystem in Non Small Cell Lung Cancer

2 Invasion

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- 14

16 Abstract

17 Many tumors are characterized by genetic instability, producing an assortment of genetic 18 variants of tumor cells called subclones. These tumors and their surrounding 19 environments form complex multi-cellular ecosystems, where subclones compete for 20 resources and cooperate to perform multiple tasks, including cancer invasion. Our recent 21 empirical studies revealed existence of such distinct phenotypes of cancer cells, leaders 22 and followers, in lung cancer. These two cellular subclones exchange a complex array of 23 extracellular signals demonstrating a symbiotic relationship at the cellular level. Here, we 24 develop a computational model of the microenvironment of the lung cancer ecosystem to 25 explore how the interactions between subclones can advance or inhibit invasion. We 26 found that, due to the complexity of the ecosystem, invasion may have very different 27 dynamics characterized by the different levels of aggressiveness. By altering the 28 signaling environment, we could alter the ecological relationship between the cell types 29 and the overall ecosystem development. Competition between leader and follower cell 30 populations (defined by the limited amount of resources), positive feedback within the 31 leader cell population (controlled by the focal adhesion kinase and fibronectin signaling), 32 and impact of the follower cells to the leaders (represented by yet undetermined 33 proliferation signal) all had major effects on the outcome of the collective dynamics. 34 Specifically, our analysis revealed a class of tumors (defined by the strengths of 35 fibronectin signaling and competition) that are particularly sensitive to manipulations of 36 the signaling environment. This class can undergo irreversible changes to the tumor 37 ecosystem that outlast these manipulations of feedbacks and have a profound impact on 38 invasive potential. Our study predicts a complex division of labor between cancer cell 39 subclones and suggests new treatment strategies targeting signaling within the tumor 40 ecosystem.

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42 Author Summary

Cancer is an elusive disease due to the wide variety of cancer types and adaptability to
treatment. How is this adaptability accomplished? Loss of genetic stability, a hallmark of
cancer, leads to the emergence of many different types of cancer cells within a tumor.
This creates a complex ecosystem where cancer cell types can cooperate, compete,
and exploit each other. We have previously used an image-guided technology to isolate

- 48 distinct cancer subclones and identify how they interact. Here, we have employed
- 49 mathematical modeling to understand how the dynamic feedbacks between different
- 50 cancer cell types can impact the success of invasion in lung cancer. We found that
- 51 successful invasion required for feedbacks to support the less viable but more invasive
- 52 cell types. These predictions may have implications for novel clinical treatment options
- 53 and emphasize the need to visualize and probe cancer as a tumor ecosystem.

55 Introduction

Lung cancer is the second most prevalent type of cancer causing over 150,000 deaths 56 57 per year in the United States [1]. Insufficient progress has been made in achieving 58 efficacious treatments. One of the main barriers in developing new treatment strategies 59 is the vast diversity between and within cancers; heterogeneity exists between patients 60 with the same tumor type, between tumor loci within a patient (i.e. metastases and 61 primary tumor), and within the primary tumor itself [2,3]. Cancer is distinguished by loss 62 of normal control over cell processes leading to genetic instability and unregulated 63 growth. Genetic instability creates array of different clonal populations with different cell 64 fitnesses, renewal and invasion potential [4]. Competition between different cancerous 65 subclones and between cancerous and normal cell types sets the stage for classical 66 ecological dynamics in the tumor microenvironment. The outcome of this process determines success of the tumor progression and its understanding may help discover 67 68 novel treatment strategies [5,6].

Invasion of surrounding tissue, either locally or distally via metastasis, is a hallmark of cancer [7]. Extensive research has detailed that invasion is mediated by interactions between tumor and extracellular matrix [8,9] and cancer-associated fibroblasts [10], but there is a lack of focus on the cooperative interactions between distinct cancer subclones. Indeed, in mouse models of lung cancer, collective invasion of cancer cells was shown to correspond markedly more successful metastasis [3,11–13], confirming the critical role of collective invasion in driving cancer progression.

76 We recently developed a novel image-guided genomics approach termed SaGA that 77 allowed us to identify at least two distinct phenotypic cell types in lung cancer invasion 78 packs: highly migratory leader cells and highly proliferative follower cells [14]. Genomic 79 and molecular interrogation of purified leader and follower cultures revealed differential 80 gene expression prompting distinguishing phenotypes. Specifically, leader cells utilized 81 focal adhesion kinase signaling to stimulate fibronectin remodeling and invasion. Leader 82 cells also overexpressed many components of the vascular endothelial growth factor 83 (VEGF) pathway facilitating recruitment of follower cells but not the leader cell motility 84 itself [14]. However, leader cells proliferated approximately 70% slower than follower 85 cells due to a variety of mitotic and doubling rate deficiencies. These deficiencies could 86 be corrected by addition of cell media extracted from the follower only cell cultures, 87 leading to conclusion that follower cells produce an unknown extracellular factor responsible for correcting mitotic deficiencies in the leader cells. In sum, leader cells provide an escape mechanism for followers, while follower cells (and follower cell media only) help leaders with increased growth. Together, these data support a serviceresource mutualism during collective invasion, where at least two phenotypically distinct cell types cooperate to promote their escape.

93 In this new study, we developed population-level computational model to explore 94 impact of the complex interactions between leaders and followers cell types on cancer 95 progression. The model implemented effects of critical signaling factors controlling the 96 communication between cell types and the interaction between cells and environment. 97 We derived analytic boundaries dividing parameter space, representing the major 98 signaling feedbacks, by the critical changes to invasion dynamics. Our study predicts the 99 critical role of specific signaling pathways involved in the symbiotic interactions between 100 cancer subclones for the overall success of cancer progression.

102 Methods

103 Our model tracks the cell counts of leader cells, L, and follower cells, F, the 104 concentrations of extracellular factors VEGF, V, an unidentified Proliferation factor, P, 105 and Fibronectin, N, as well as the size of the domains for leader cells, Ω_L , and for 106 follower cells Ω_F . Based on the available data [14], the following processes have been 107 implemented. Leader cells can expand their domain, Ω_L , by secreting Fibronectin, which 108 in turn relaxes competitive pressure on leader cell growth. Leaders also secrete VEGF, 109 which is taken up by follower cells and causes follower cells to follow them. This was 110 modeled by increasing the domain for follower cells, which in turn relaxes competitive 111 pressure on follower cells. Follower cells secrete an unknown proliferation signal that 112 increases the reproductive capacity of leader cells (initially smaller than follower cells). 113 Leader and follower cells also must compete with each other for resources at rate c (see 114 Figure 1).

We modeled cell counts (*L* and *F* species) as standard Lotka-Volterra competition [15]. The carrying capacity of the leader cells was dynamic and dependent on the amount of proliferative signal, *P*, present. This capacity increased in a saturating manner with *P*, with maximum equal to the follower cell carrying capacity, K_{F0} . Intra- and interspecific competition was driven by concentration, i.e. $[L]=L/\Omega_L$, and birthrate was driven by absolute number, *L*. The extracellular species (*V*,*P*,*N*) and domain sizes all had linear dynamics for simplicity. Below primes denote the time derivative of the variable.

122
$$\frac{L'}{L} = r_L \left[1 - \frac{(L/\Omega_L) + c(F/\Omega_F)}{K_{L0} + (K_F - K_{L0}) \left(\frac{P}{\delta + P}\right)} \right]$$
(1)

123
$$\frac{F'}{F} = r_F \left[1 - \frac{(F/\Omega_F) + c(L/\Omega_L)}{K_F} \right]$$
 (2)

$$124 V' = \beta_V L - \gamma_V V (3)$$

$$125 \qquad P' = \beta_P F - \gamma_P P \tag{4}$$

$$126 N' = \beta_N L - \gamma_N N (5)$$

127
$$\Omega_L' = \beta_{OL} N - \gamma_{OL} (\Omega_L - \Omega_{L0})$$
(6)

128
$$\Omega_F' = \beta_{OF} V - \gamma_{OF} (\Omega_F - \Omega_{F0})$$
(7)

Here r_L and r_F denote the rate of expansion for leaders and followers, respectively. The parameter *c* denotes the strength of competition between the two cell types. The capacity of the environment for follower cells is given by the parameter K_F . The capacity for leaders depended on an initial capacity, K_{L0} , and on the amount of proliferation signal in a Hill-like manner with EC50, δ . Each extra-cellular species (*V*,*P*,*N*) had a production rate, β , and a degradation rate γ , the domain size variables (Ω_L and Ω_F) also had a parameter denoting initial capacity (Ω_{L0} and Ω_{F0}).

136

137 Reduction and Feedbacks

Previous 3D spheroid experiments show that invasion occurs on a much faster time scale than reproduction [14]. By assuming that factors (*V*,*P*,*N*) and domains (Ω_L, Ω_F) change much faster than cell counts, one can reduce these equations to a set of two equations (*L*,*F*), where variables in equations (3)-(7) are at their equilibria

142
$$V_{SS} = \frac{\beta_V}{\gamma_V} L;$$
 $P_{SS} = \frac{\beta_P}{\gamma_P} F;$ $N_{SS} = \frac{\beta_N}{\gamma_N} L;$ $\Omega_L^{SS} = \frac{\beta_{OL}}{\gamma_{OL}} N;$ $\Omega_F^{SS} = \frac{\beta_{OF}}{\gamma_{OF}} V;$ (8)

143 Using this reduction drastically reduced the complexity of the system. First, we defined the feedbacks based on the reduced system. The feedback that determines the leaders 144 impact on their own domain expansion was denoted by $s_L = \frac{\beta_N \beta_{OL}}{\gamma_N \gamma_{OL}}$, for the strength of the 145 leader only feedback. The feedback that determines the leaders impact on follower cell 146 growth was denoted by $s_{LF} = \frac{\beta_V \beta_{OF}}{\gamma_V \gamma_{OF}}$, for the strength of the leader to follower feedback. 147 148 The feedback that determines the followers impact on leader cell growth was denoted by $s_{FL} = \frac{\beta_P}{\gamma_R \delta}$, for the strength of the follower to leader feedback. Second, using these 149 150 assumptions, we re-wrote the leader-follower system as

151
$$\frac{L'}{L} = r_L \left[1 - \frac{(L/\Omega_L^{SS}(L)) + c(F/\Omega_F^{SS}(L))}{K_L(F)} \right]$$
(9)

152
$$\frac{F'}{F} = r_L \left[1 - \frac{(F/\Omega_F^{SS}(L)) + c(L/\Omega_L^{SS}(L))}{K_F} \right]$$
(10)

153 where

154
$$\Omega_L^{SS}(L) = s_L L + \Omega_{L0};$$
 $\Omega_F^{SS}(L) = s_{LF} L + \Omega_{F0};$ $K_L(F) = K_{L0} + (K_F - K_{L0}) \frac{s_{FL}F}{1 + s_{FL}F}$

155 Using this reduction we can derive several critical points in invasion. The reduced system (9),(10) may have five equilibrium points: extinction of leaders (O1: L=0, F>0), 156 157 followers (O_2 : L>0, F=0), both (O_3 : L=0, F=0), and two coexistence points (O_4 , O_5) (where both leaders and followers populations are non-zero: L>0, F>0; O_4 is always 158 159 stable, wheras O₅ is unstable). Changes in the feedback strengths cause fundamental 160 shifts in dynamics. In the following we used parameter values $\Omega_L = 1$, $\Omega_F = 1$. To match 161 experimental observations that leader cells grow slower and less effeciently, we set 162 $r_L = 0.3$ and $K_{L0} = 0.3$ while $r_F = 1$ and $K_F = 1$. The strengths of the various 163 feedbacks, s_L , s_{LF} , and s_{FL} are varied systematically below.

164

165 Transcritical Bifurcation at Zero

166 To determine the critical points in the leader-follower system, we calculated the Jacobian 167 of the reduced system evaluated for the leader extinction equilibrium (O₁ :*L* = 0, *F* = 168 $F_{LE} = \Omega_F \cdot K_F$).

169
$$J|_{L=0;F=F_{LE}} = \begin{pmatrix} r_L(1 - \frac{cK_F}{K_L^{SS}}) & 0\\ r_F\left(\frac{K_{F_0SFL}}{\Omega_{F_0}^2} - \frac{c}{\Omega_{L_0}}\right) & -r_F \end{pmatrix}$$
 (11)

170

171 Here $K_L^{SS} = K_{L0} + (K_F - K_{L0}) \frac{s_{FL} \cdot F_{LE}}{1 + s_{FL} \cdot F_{LE}}$, the value of K_L when $F = F_{LE}$. The Jacobian has 172 eigenvalues

173
$$\lambda = \left[r_L \left(1 - c \frac{K_F}{K_L^{SS}} \right), -r_F \right]$$
(12)

For $c < \frac{K_L^{SS}}{K_F}$, O₁ is unstable and O₄ (steady state where both L>0 and F>0) is stable. At $c = \frac{K_L^{SS}}{K_F}$ these two equilibria coincide, and for $c > \frac{K_L^{SS}}{K_F}$ equilibrium O₄ moves to the left of the L=0 axis and becomes unstable while O₁ gains stability. Thus, extinction of leaders (O₁) is stable as long as $c > \frac{K_L^{SS}}{K_F}$, which determines an upper bound on competition where leader and followers can coexist and a bifurcation we call the transcriticalbifurcation at zero.

180

181 Saddle Node Bifurcation

The system undergoes a saddle node bifurcation when two coexistence equilibria (O_4 and O_5), representing non-zero populations of both leaders and followers, coincide and disapper. Beyond this bifurcation point the leader/follower populations undergo unbounded growth. This bifurcation was determined numerically using MatCont [16]. We found that this bifurcation point depends critically on both the leader feedback strength, s_L , and on the competition strength, *c*. One of these coexistence points is effected by the transcritical bifurcation, below.

189

190 Transcritical Bifurcation at Infinity

191 When the leader feedback strength is sufficiently high relative to competition, leaders 192 and followers may undergo unbounded growth from the initial conditions belonging to the 193 certain regions of the phase space. We describe this scenario as an attractor basin in 194 the phase space for the stable infinity attractor. However, if s_L is reduced (or *c* is 195 increased) beyond a certain threshold, infinity becomes unstable. This corresponds 196 precisely with the loss of an unstable coexistence equilibrium with non-zero values of 197 both leaders and followers (O_5). Leaders and followers that are coexisting must satisfy

$$198 \qquad \frac{L}{\Omega_L} + c \frac{F}{\Omega_F} = K_L(F) \tag{13}$$

199 and $\frac{F}{\Omega_F} + c \frac{L}{\Omega_L} = K_F$ or equivalently,

$$200 \qquad \frac{F}{\Omega_F} = K_F - c \frac{L}{\Omega_L}.$$
(14)

201 In the case that follower populations are large relative to δ , $K_L(F) \rightarrow K_F$, we substituted 202 (13) into (14) to find

203
$$L = \frac{K_F \Omega_{L0}}{(1+c)\left(1 - \frac{K_F S_L}{1+c}\right)}$$
(15)

204 which has a discontinuity at

$$205 c = K_F s_L - 1 (16)$$

206 defining the loss of one of the coexistence equilibrium points (O_5) when it moves to

207 infinity. We describe this as the transcritical bifurcation at infinity as the stability of infinity

changes at this point.

209 Results

210 Leader and Follower Ecosystem

211 Leader and follower cell types in non-small cell lung cancer spheroids were previously 212 isolated using a fluorescence technique termed SaGA [14] (Figure 1A,B). We found that 213 leaders and followers are genotypically and phenotypically distinct populations of cancer 214 cells that exchange a variety of signaling molecules to coordinate complex behavior 215 during invasion. In this new work, we focus on four main channels of communication 216 (see Figure 1C). Leader cells secrete fibronectin in an autocrine manner. This leads to 217 ECM restructuring and expansion of leader cell domain, Ω_L , (see Methods) which 218 ultimately increases the leader cell count. The strength of this positive feedback is 219 characterized in our model by s_L (strength of Leader only feedback). Leader cells also 220 secrete VEGF. In the leader-follower ecosystem this promotes follower cells to track 221 expanding leader cells, increases follower domain size (Ω_F) , and ultimately, follower cell 222 count. In our model, the strength of this feedback is given by s_{LF} (strength of Leader to 223 Follower feedback). Follower cells secrete an undetermined proliferation signal, as 224 evidenced by the observation that follower-only cell media increases leader cell growth 225 rate [14]. The strength of this feedback is given by s_{Fl} (strength of Follower to Leader 226 feedback) in the model. Finally, both cell types compete for resources, which is modeled 227 here by the feedback c.

228 These feedback mechanisms were incorporated into a modified Lotka-Volterra 229 type competition-cooperation model. We chose a Lotka-Volterra model to focus on the 230 ecological aspects of competition in the cancer ecosystem. Here, the leader cells could 231 grow to a total capacity K_L , which is an increasing function of the proliferation signal 232 secreted by the follower cells. This capacity was reached when a combination of leader 233 and follower cell densities (cell counts divided by domains) exceeds K_L (see Methods). 234 Increases in the domain size of each type (by Fibronectin secretion in the leader case 235 and VEGF in the follower case) limited the overall density of that cell type and mitigated 236 its impact on the overall capacity of the system. Increasing competition, for example by 237 limiting resources, increased the impact of either cell type on the conjugate capacity type 238 (e.g. how leader density, L/Ω_L , impacts follower capacity K_F).

This system of the feedbacks between the leader and follower cells describes a complex dynamical ecosystem. The impact these feedbacks may have on cancer growth or invasion is unclear. Leader and follower cells are engaged in competition for resources but can also be engaged in cooperation and play supportive roles. For example, invasive leader cells provide new territory for the follower cell population and are supported by proliferative follower cells. In the following, we analyzed the model to find critical turning points for the ecosystem dynamics.

246

247 Multiple Types of Invasion Dynamics

248 We found that multiple feedbacks between the leader and follower cell populations could 249 produce a wide variety of complex dynamics. When competition strength, c, was high 250 and the strength of the leader only feedback, s_{L} , was moderate, population dynamic was 251 bounded and resulted in a stable cell count for both leader and follower cell populations 252 as well as a stable domain size (Fig. 2A). In contrast, when feedback was large and 253 competition was moderate, population dynamics revealed an unbounded growth (Fig. 254 2B). Intermediate values of both c and s_i led to dynamic regimes that depended on the 255 initial cell count: ecosystems with large initial cell count underwent unbounded growth, 256 while small ecosystems attained a stable size (Figure 2C). These types of dynamics are 257 in a qualitative agreement with experimental studies which revealed (a) rapid expansion 258 of intact leader-follower ecosystem and (b) that blocking specific feedback mechanisms 259 in vitro can reduce or block cell population growth. Specifically, blockade of fibronectin 260 signaling or blockade of VEGF signaling led to significantly reduced invasion [14].

This array of behaviors can be explained by the critical shifts in the cell population dynamics due to the changes in the feedbacks strength. We found that depending on the level of competition, *c*, and the strength of invasiveness of leaders, s_L , the leader-follower ecosystem can operate in one of five different regimes, as described below (Figure 3).

Leader Extinction: When competition was high and invasive feedback was minimal, the leader cells (the weaker competitor) were forced to extinction while the follower cells persisted and its population reached a stable size (Figure 3A,B). There was a critical level of competition between leaders and followers, given by $c > \frac{K_L^{SS}}{K_F}$ (see Methods, *Transcritical bifurcation at zero* for derivation), required for this type of dynamics. This critical level of competition, the ratio of the capacity of leader cells, K_L^{SS} , to that of the follower cells, K_F , essentially depends on the fitness differences between leader only and follower only cell populations. Leader and follower populations with similar fitness would tolerate a much higher competition threshold without driving one species to extinction. From the dynamical systems perspective, when $c > \frac{K_L^{SS}}{K_F}$ and s_L is sufficiently low (see below), the only stable equilibrium in the phase space is O₁ and all system trajectories converge to this equilibrium point representing the leader cells extinction state (Figure 3B).

279 Leader Extinction with Escape: If competition was above the leader extinction limit, $c > \frac{K_L^{SS}}{K_{F0}}$, but not high enough to balance the impact of the leader only feedback, 280 281 $c < K_F s_L - 1$, there were two possible outcomes depending on the initial population size 282 (see Methods, Transcritical Bifurcation at Infinity for derivation) (Figure 3A,C). The 283 second condition, $c < K_F s_L - 1$, can be interpreted as a balance between positive feedback, s_L , and negative feedback, c. In this regime, leaders could go extinct if the 284 285 initial population of leader cells was sufficiently small. Alternatively, if initial populations 286 of leaders and followers both were large enough, the ecosystem could grow 287 unboundedly. Thus, our model predicts, that the ability to undergo successful collective 288 invasion depends on whether the initial bulk size is larger than a critical amount. These 289 types of dynamics with divergent outcomes occur when competition is large enough to 290 be able to drive leaders extinct, but small enough so it can be outbalanced by the strong 291 invasive effects of the leader cells.

In the phase space of the model, the basins of attraction of the two distinct dynamical regimes are separated by a critical boundary (separatrix of a saddle equilibrium O_5) where the cell bulk size determines its ultimate fate (Figure 3C). Both infinity and the leader extinction equilibrium (O_1) are stable attractors representing two possible end solutions of the system dynamics.

Non-invasive Dynamics: When competition was smaller than the extinction limit, $c < \frac{K_L^{SS}}{K_{F0}}$, but large enough to balance leader feedback strength, $c > K_FS_L - 1$, the ecosystem size remained bounded and both leaders and followers attained a stable and non-zero population size. In the phase space, this type of dynamics corresponds to conversion to the stable equilibrium O₄ (Figure 3A,D). We refer to this as non-invasive dynamics, as the cells cannot grow beyond a defined size. In this case, while competition was present, it was too weak to lead to extinction, while leader population
was not invasive enough to promote unlimited growth. This scenario represents stable,
non-invasive dynamics.

306 **Multimodal Dynamics:** If competition was (a) small enough to allow leader existence, $c < \frac{K_L^{SS}}{K_{EQ}}$ (b) small enough relative to the leader feedback strength, so that escape was 307 308 possible, $c < K_F s_L - 1$, but (c) high enough, so that for small initial population of leader it 309 could balance the positive leader feedback, leader and follower cell dynamics depended 310 on the initial population size (Figure 3A,E). Ecosystems with a large initial cell count 311 would grow without bound but those with a small initial cell count would reach a stable 312 population size, due to the competition as in the non-invasive dynamics case. On the 313 phase plane the last outcome was represented by a contraction to a stable equilibrium 314 O_4 . This critical boundary was defined by a separatrix of a saddle fixed point O_5 (Fig. 3E). 315 (This separatrix was determined numerically by reversing time [17].)

Aggressive Dynamics: When leader invasive strength was sufficiently high and competition was sufficiently low, the only possible outcome was unbounded growth of both cell populations (Figure 3A, F). In this case, the only stable attractor in the phase space is infinity where all system trajectories are converged to.

320 In summary, our analysis revealed that the complex balance of the feedbacks in 321 the leader-follower ecosystem can lead to the multiple types of population dynamics. 322 When the leaders' invasiveness was low, the outcome depended on the competition 323 between two populations – strong enough competition promoted leader extinction, while 324 weak competition allowed stable coexistence states with bounded size of both leader 325 and follower cell populations. As leader invasiveness rate increased, the system 326 revealed a new state with unbounded growth. This aggressive dynamic state coexisted 327 with a stable attractor representing a bounded size of both populations if competition 328 between leaders and followers was strong enough. Otherwise, unlimited population 329 growth was the only outcome. Based on the system dynamics derived above, next we 330 will show how critical boundaries between parameter regimes could be exploited to lead 331 to profound changes in the ecosystem dynamics.

332

333 Limiting leader feedback Leads to Irreversible Changes in Invasion

334 In the multimodal dynamics (e.g., Fig. 3C or 3E), leader and follower cell populations can 335 undergo explosive growth or achieve a stable count depending on the initial size of the 336 ecosystem. We examined the impact of limiting the invasive leader feedback in 337 scenarios of this type (Fig. 4). Even when the ecosystem was initially sufficiently large to 338 support unbounded growth, after reducing invasive leader feedback s_i (Fig. 4A), the 339 ecosystem was forced into the non-invasive dynamics type and the total bulk of the cell 340 population reduced reaching a steady-state (Fig. 4E). Importantly, the leader and 341 follower cell populations remained stable and bounded after restoring invasive leader 342 feedback to its original strength (Fig 4E, right side).

From the point of view of the dynamical systems analysis, reducing leader feedback changed the phase space, so the only stable attractor was non-zero equilibrium (O_4) (Fig. 4C). In this regime, unlimited growth was abandoned and the system converged to the equilibrium state (O_4) corresponding to the bounded size of both cell populations. This equilibrium remained stable even after the feedback was restored to its original level (Fig. 4D).

Our model predictions (Fig. 5A) are consistent with in vitro data (Fig. 5B). Using siRNA blocking we previously showed that expression of fibronectin (which is characterized by the strength of leader only feedback, s_L , in the model) led to the low invasion potential and a stable cell population size [14].

353

354 Increasing Competitive Signals Leads to Leader Extinction

355 We next tested effect of increasing competition between leader and follower cell 356 populations on the ecosystem dynamics (Fig. 6). Leader cells excrete extracellular 357 factors that induce the death of the followers and leaders alike [14], which supports 358 competition. Here, we again started from aggressive unbounded type of dynamics and 359 then increased competition strength (Fig. 6A). This caused change of the ecosystem 360 dynamics. Both cell populations reduced the size, with leader cell population going to 361 extinction state (Fig. 6E). However, upon restoring competition to the original level, 362 leader and follower cells reemerge and grow unboundedly again. The last can be 363 avoided if no leader cells remain (complete extinction).

364 Again, this dynamic can be easily understood using bifurcation analysis. Increasing 365 competition strength made leader extinction equilibrium state O_1 stable (Fig. 6C). 366 However, when competition was restored to its original level, O₁ became unstable again 367 and leader and follower cells returned to escape dynamics (Fig. 6D). Importantly, in the 368 extreme case of very small cell populations, cells undergo discrete and stochastic 369 dynamics and complete extinction of a small population of leaders is possible in a finite 370 time, leading to irreversible changes due to competition increase (similar dynamics was 371 described in our previous study [18]).

372

373 Support For Leaders has Large Impact on Aggressiveness

374 Changing the strength of the feedbacks that determine the interaction between 375 leaders and followers (s_{LF} and s_{FL}) could also impact the dynamics. Leader cells secrete 376 VEGF (denoted here by s_{LF}) that helps follower cells to expand their territory and follower 377 cells secrete a proliferation signal (denoted here by s_{Fl}) that allows leaders to increase 378 their proliferative capacity. These two feedbacks have distinct impacts on the overall 379 ecosystem dynamics. Perturbations to s_{LF} (changing the impact that leaders have on 380 followers) changed the system dynamics (assuming that the cell count was small 381 enough at the time of the intervention) from unlimited growth to the bounded type. The 382 size of both leader and follower cell populations decreased reaching non-zero steady-383 state (Fig. 7E). This regime persisted as long as the feedback from the leaders to 384 followers remained low. However, increasing sLF to its original level restored the system 385 dynamics with unlimited cell population growth (Fig. 7E, right size).

386 Using bifurcation analysis, we found that reducing impact that leaders have on 387 followers shifted the location of the saddle node bifurcation boundary that separated 388 state with unlimited growth only dynamics and a state with coexistence of the unlimited 389 growth and a stable equilibrium attractor (O_4) regimes (Fig. 7A). Effectively, decreasing 390 s_{LF} increased the threshold level of the invasive leader feedback (s_L) needed to cause 391 unbounded growth. Thus, reducing s_{1F} made the system to converge to the stable 392 equilibrium state O_4 corresponding to the bounded size of both cell populations (Fig. 7C). 393 However, increasing s_{LF} to its original level changed the phase space again, so infinity 394 became the only stable attractor (Fig. 7D) and unlimited growth dynamics resumed.

395 Our model predictions (Fig. 5C) are consistent with in vitro data (Fig. 5D). Using 396 siRNA to block the VEGF receptor VEGFR2 (siKDR in Fig. 5D), we previously showed 397 that blocking the leader to follower feedback led to the limited invasion potential and 398 stable cell population size (Fig. 5D) [14].

399 Finally, we tested the role of the follower to leader feedback (s_{FL}) and found that 400 perturbations to s_{FL} have a significant impact on the system dynamics. In contrast to s_{LF} , 401 changes to the s_{FI} changed both the location of the saddle node bifurcation boundary 402 and the transcritical bifurcation boundary of the leader extinction (Fig. 8A). Therefore, 403 decreasing s_{FL} both increased the threshold on the leader invasion strength (s_L) needed 404 to cause unbounded population growth and decreased the threshold of the competition 405 strength (c) needed to induce leader population extinction. We have exploited this to 406 show that decreasing s_{FL} can cause irreversible change in the cell population bulk. Again, 407 starting with unlimited growth dynamics (Fig. 8B), decreasing follower to leader feedback, 408 s_{LF} , reversed the dynamics and both leader and follower cell population reduced in size 409 converging to the steady-state (Fig. 8E). This regime with bounded ecosystem size 410 persisted after the feedback was restored (Fig. 8E, right side).

411 Using dynamical systems analysis, we found that reducing follower to leader 412 feedback (s_{FL}) triggered the system convergence to the stable attractor (O_1) representing the leader extinction state (Fig. 8C). When the feedback was restored, O1 becomes 413 414 unstable but the ecosystem fell to the attraction basin of the stable equilibrium O₄ and 415 avoided regime of unlimited growth (Fig. 8D). In more general case, the outcome 416 depended on the balance between the leader to follower, s_{FI} , and follower to leader, s_{IF} , 417 feedbacks, with higher s_{LF} requiring more significant s_{FL} decrease to avoid unbounded 418 growth (Fig. 8F).

419

420 Summary of Perturbations to Cancer Ecosystem

Complex balance of the feedbacks within the cancer cell ecosystem allows for some alterations of the feedback parameters to have significant impacts on the ecosystem dynamics. We summarized these different possibilities in Table 1 from the perspective of achieving the goal to reduce cell population bulk. Hence, manipulating s_L , s_{LF} , s_{FL} should be interpreted as decreasing these feedbacks, whereas manipulating c should be interpreted as increasing c. We also examined the possibility of non-targeted cell death, 427 such as might occur during non-specific chemotherapy. Manipulations were either 428 irreversible, so the system dynamics remained altered upon cessation of the 429 perturbation (e.g. irreversible leader extinction or irreversible stabilization of the cell 430 count), or caused only temporal and reversible reduction of the cell bulk. In some cases, 431 such as leader extinction with escape and multimodal dynamics (see Fig. 3), the size of 432 the initial cell bulk dictated possible outcomes of the feedback perturbations. The 433 outcomes described in the Table 1 represent the best-case scenario. Thus, 434 perturbations were started from an appropriate initial state and maintained long enough 435 to achieve the desired effect.

436 This analysis revealed that certain parameter regimes are more sensitive to the 437 perturbations than others. Specifically, in the leader extinction with escape regime (area 438 (2) in Figure 3A) and the multimodal dynamics regime (area (4) in Figure 3A) 439 perturbations could have irreversible impacts on the ecosystem. In these cases, any 440 perturbation (death, reduction in s_L , s_{LF} , s_{FL} , or increase in c) can potentially force the 441 system to cross the critical boundary (separatrix) and transition from explosive growth to 442 a steady-state dynamic. These regimes give a unique opportunity to impact the 443 invasiveness of the ecosystem.

Also, certain perturbations could force the ecosystem into a state where leader extinction (O_1) is stable. This occurs when applying sufficient increases in the competition pressure, *c*, or decreases in the support from followers to leaders, s_{LF} . In these cases, it is possible for the discrete and stochastic nature of the cell population dynamics to define the ecosystem fate. Thus, a sufficiently long perturbation could irreversibly eradicate a sufficiently small discrete number of leader cells [18].

451

Type of dynamics	Initial State	Manipulation Type	Outcome
1. Leader	N/A	death, <i>c, s_L, s_{LF}, s_{FL}</i>	Reversible cell bulk reduction.
Extinction			
2. Leader	Infinity attractor	death, <i>s_L, c, s_{LF}, s_{FL}</i>	Irreversible leader extinction.
Extinction w/	basin		
Escape	Stable attractor	death, <i>c, s_L, s_{LF}, s_{FL}</i>	Reversible cell bulk reduction.
	basin		
3. Non-invasive	N/A	<i>c, s_{FL} s_L, s_{LF}</i> , death	Reversible cell bulk reduction.
Dynamics			
4. Multimodal	Infinity attractor	<i>c, s_{FL}, s_L, s_{LF},</i> death	Irreversible stabilization of cell bulk.
	basin		
	Stable attractor	<i>c, s_{FL,} s_L, s_{LF}</i> , death	Reversible cell bulk reduction.
	basin		
5. Aggressive	N/A	<i>c, s_{FL} s_L, s_{LF}</i> , death	Reversible cell bulk reduction.
Dynamics			

Table 1: Effect of the different feedback alterations on the ecosystem dynamics. 452 453 First column - the different types of the ecosystem dynamics in the (s_L, c) parameter 454 space are as shown in Figure 3. In some cases, initial population size dictated the 455 outcome. These outcomes are distinguished in the initial state given in the 2d column. 456 An initial state in the infinity attractor basin denotes that the cell bulk exceeded the 457 critical amount and could grow unboundedly; initial state in the stable attractor basin 458 denotes the case when the cell counts were less than the critical value and the system 459 converged to the stable attractor. Third column - each transient manipulation to the 460 model parameters had a goal to reduce cell population size (decrease of s_{L} , s_{FL} , s_{FL} or 461 increase of c). The "death" indicates a non-targeted "enforced" reduction of the cell 462 population. The last column indicates the system dynamics after the original values of 463 the model parameters were restored. The changes were either irreversible upon 464 cessation of the perturbation (e.g. irreversible leader extinction and irreversible 465 stabilization of cell bulk) or caused reversible reduction in the cell population bulk.

467

468 Discussion

469 Heterogeneity of tumors, at the genetic, epigenetic, and phenotypic levels, is one 470 of the main obstacles to developing new effective treatment strategies. Tumor cells 471 rapidly evolve forming highly efficient symbiotic systems with well-defined labor division 472 targeted to augment tumor survival and expansion. In lung cancer collective invasion 473 packs observed in vitro, two distinct populations of cancer cells - highly migratory leader 474 cells and highly proliferative follower cells - have been recently identified [14]. In this 475 new study, we used computational models to explore collective dynamics of the leader-476 follower ecosystem and to exploit approaches that can effectively disrupt it.

477 We found that competition between two populations (defined by the limited 478 amount of resources), the positive feedback within the leader cell population (controlled 479 by the focal adhesion kinase and fibronectin signaling) and impact of the follower cells to 480 the leaders (represented by yet undetermined proliferation signal) all had major effects 481 on the outcome of the collective dynamics. While increase of the positive feedback 482 within the leader cell population would ultimately lead to the system state with 483 unbounded growth, manipulating follower to leader feedback or increasing competition 484 between leader and follower cell populations was able to reverse this dynamic and to 485 form a stable configuration of the leader and follower cell populations.

486 Our model highlights the importance of focal adhesion kinase (FAK) and 487 fibronectin signaling. Our previous empirical work showed that FAK signaling was a key 488 distinguishing feature between leader and follower cells and critical for invasive leader 489 behavior [14]. Our model predicts that FAK is the main driver of invasion by leader cells 490 and disruptions in the FAK driven feedback loop cause critical changes in the leader-491 follower population dynamics. Indeed, FAK is a well-known regulator of the tumor micro-492 environment: promoting cell motility and invasion [19]. FAK expression is upregulated in 493 ovarian [20] and breast cancer [21] tumors with expression levels correlating with survival [22,23]. Many FAK inhibitors, such as defactinib, are currently in clinical trials 494 495 with promising results [19,24,24–28]. A key advantage of FAK inhibitors is that they 496 impact both the tumor itself and the surrounding stroma where tumor associated 497 fibroblasts also utilize FAK signaling to promote tumor invasiveness [29,30].

498 While commonly associated with angiogenesis in healthy and cancerous tissue, 499 our previous work showed that VEGF mediates communication between leader and 500 follower cells [14]. There is a long history of targeting VEGF to limit tumor invasiveness 501 [31,32]. While great success has been seen in preclinical models [33,34], only moderate 502 success was seen in clinical trials with anti-VEGF drugs such as bevacizumab [35,36]. 503 This is largely due to cancers developing resistance to specific VEGF-therapeutics. In 504 our model, VEGF stimulated followers to shadow leaders and expand their domain. 505 However, we found that inhibition of VEGF had little impact on the ecosystem dynamics 506 relative to the perturbations of the other axes (such as FAK or competition for resources).

507 Competition for resources is one of the principal forces that structures any 508 ecosystem, including tumor ecosystems [6,37]. Our modeling work predicts that 509 competition was a critical component in the leader-follower ecosystem. We found that 510 when the strength of competition exceeded a critical threshold, leaders (the weaker 511 competitor) were driven to extinction. Further, enhancements of the competition in the 512 model changed the fundamental cell population dynamics. In some cases this meant 513 stopping unbounded growth and promoting the extinction of the leader cells. Our 514 previous in vitro work demonstrated that leaders may inhibit the growth of followers 515 through an unknown secreted factor in cell media [14]. While still in the early stages, 516 exploiting this inhibition may also provide similar benefits to those shown here as 517 increases in competition.

518 Our previous study also revealed a currently unknown extracellular factor 519 secreted by followers that corrects mitotic deficiencies and enhances leader proliferation 520 [14]. Our modeling highlights this factor as having critical impact on the ecosystem 521 dynamics. We found that blockade of this proliferation factor, modeled here by the 522 strength of the followed to leader feedback, can cause critical shifts in the population 523 dynamics. More work needs to be done to identify and understand the mechanism of this 524 action, but preliminary results suggests that this may be a potential novel treatment axis 525 that specifically targets the mutualistic interaction between leaders and followers.

526 Ecological forces shape the exchange of biomaterial between different biotic and 527 abiotic environmental agents. These forces determine capacity of the ecosystem for 528 different species (subclones) and the environment ultimately sets the fitness of each of 529 the competitors. Classic ecological theory dictates that an abundance of many similar 530 species (such as similar subclonal populations) will lead to a high competition for 531 resources [38,39]. This competition can force the exclusion of inferior competitors and 532 ultimately may reduce heterogeneity of the system. However, when symbiotic and 533 mutualistic interactions occur, otherwise competitive species support each other and 534 increase the capacity of the ecosystem [40,41]. Symbiosis between different subclonal 535 populations may be particularly important during critical times when the tumor survival is 536 in peril (such as hypoxia, metastasis or therapy). One critical moment in tumor 537 progression occurs when highly proliferative tumor cells saturate the resource potential 538 of their current environment. In order to obtain more resources, tumors need to invade 539 new territory.

540 Previous results to model complex tumor cell population dynamics range from 541 detailed cellular level models (e.g. [9,42-44]) to continuous models with a different 542 degree of complexity (e.g. [18,45–48]) similar to that proposed in our new study. While 543 cellular level model can directly incorporate heterogeneous cell types and intrinsic tumor 544 properties, including proliferation, metabolism, migration, protease and basement 545 membrane protein expression, and cell-cell adhesion, they typically have high-546 dimensional variables and parameter space that is difficult to explore. Advantages of the 547 reduced type of models include the low dimensional parameter space, where 548 parameters have clear biophysical meanings, and which allows for systematic analysis 549 to rapidly explore and determine the sensitive parameter space. We previously applied 550 this approach to study cell interactions in chronic cancers and predicted conditions for 551 explosive tumor growth [18]. Similar approach was applied to model cancer cell 552 population dynamics in many other types of cancer [45,48,49].

553 The vast diversity between different cancers and even between different cell 554 types within a single tumor remains one of the biggest hurdles to overcome to achieve 555 personalized cancer treatment. This diversity leads to a complex array of interactions 556 between different tumor cell types and the healthy surrounding tissue: the tumor 557 ecosystem. Our work has isolated phenotypically unique lung cancer cells and taken a 558 dynamical approach to understanding the interactions within the tumor ecosystem. We 559 identified the critical features and interactions composing the leader-follower ecosystem, 560 to explore vulnerabilities of the lung cancer invasive cell populations.

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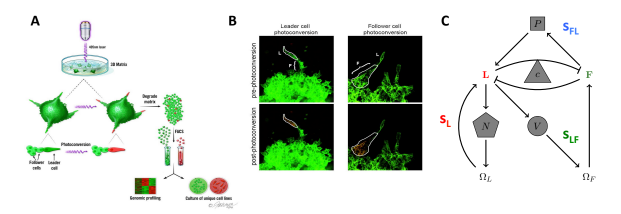
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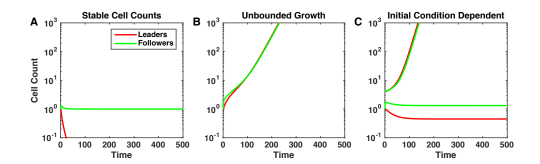
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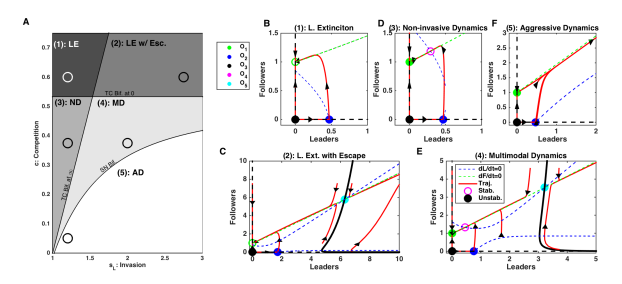
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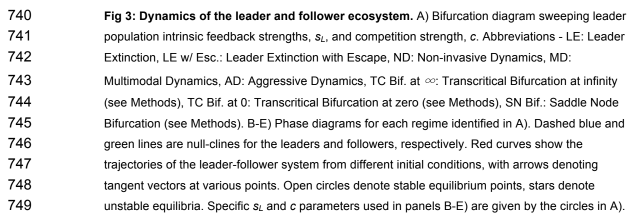


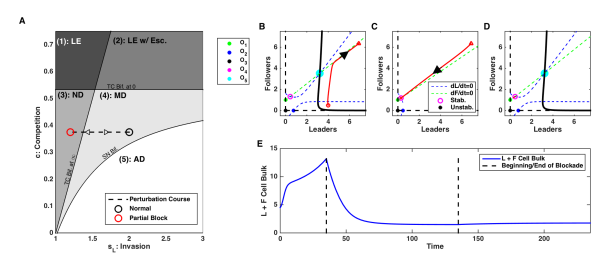
717 718	Fig 1: Leader and Follower system. A) Pictorial representation of the <u>spa</u> tio-temporal <u>g</u> enomic and cellular analysis (SaGA). Laser excitation of Dendra2 drives a change in fluorescence of user-
719	specified cells. After degradation of cell matrix, fluorescence-based cell sorting is used to separate
720	cells into leader and follower groups allowing for genetic analysis on specified groups. Printed with
721	permission from Fairman Studios, LLC. This image is not included in the Creative Commons
722	licence for the article. Adapted from Figure 1 in [14]. B) Photo-conversion examples using 3-D
723	spheroids of H1299-Dendra2 cells. L= leader cell, F = follower cell. Adapted from Figure 1 in [14].
724	C) Stick representation of mathematical model of leader and follower cell interactions and invasion.
725	Positive feedbacks are given by arrows, while negative feedbacks are given by flat-ended curves.
726	The strength of leader only feedback (s_L) is mediated by fibroNectin (N). The strength of leader to
727	follower feedback (s_{LF}) is mediated by VEGF (V). The strength of follower to leader feedback (s_{FL})
728	is mediated by a proliferation signal secreted by followers (P). The strength of competition is given
729	by c.

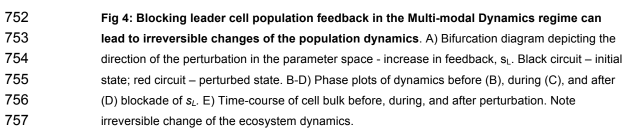


733	Fig 2: Ecosystem dynamics depend strongly on the feedbacks strength. Characteristic
734	examples of the cancer cell population dynamics in the model for different strength of the
735	competition between leader and followed cell populations, c, and leader population intrinsic
736	feedback, s_L . Cancer cell populations may attain a stable size (A: s_L = 1.2, c = 0.6), grow
737	unboundedly (B: s_L = 1.2, c = 0.05), or be dependent on the initial tumor size (C: s_L = 2, c = 0.375).









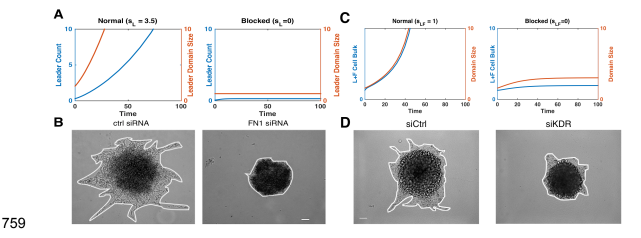
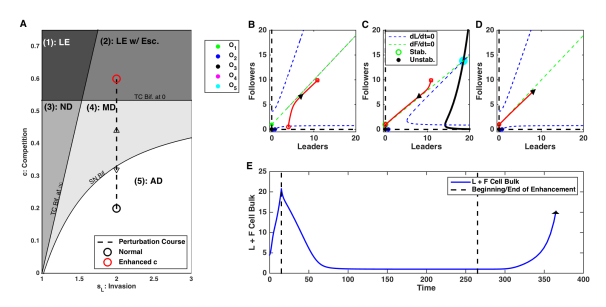
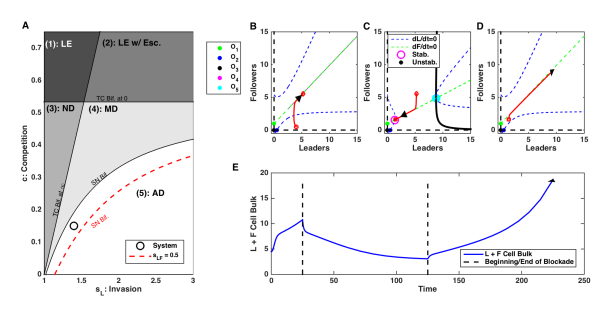


Fig 5: Model reproduces in vitro experimental results. A) In the model, leader cell count (blue) and domain size (red) are given in both normal (left) and leader population feedback, s_L , blocked (right) conditions. B) In cell culture, invasion of leader cells was significantly reduced during siRNA block of focal adhesion kinase (right), compare to control (left). Scale bar 100 μ m. Reproduced from [14]. C) In the model, blocking leader to follower feedback, s_{LF} , limited invasive area and cell count. D) Impact on invasion of leader cell cultures during siRNA block of VEGFR2 (siKDR). Scale bar 100 μ m. Invasive area was significantly reduced after VEGFR2 block (p<0.0001) (right), compare to control (left). Reproduced from [14].



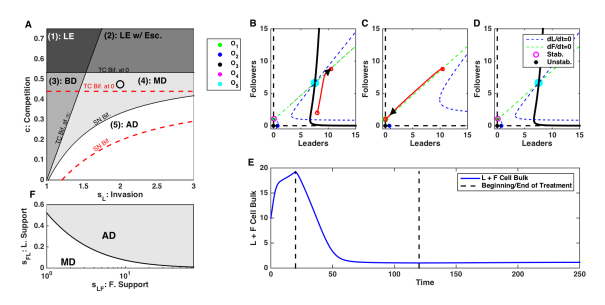
770	Fig 6: Enhancing competition between leader and follower populations can drive transient
771	extinction of leaders. A) Bifurcation diagram depicting the direction of the perturbation - increase
772	in competition, c. Black circuit – initial state; red circuit – perturbed state. B-D) Phase plots of
773	dynamics before (B), during (C), and after (D) enhancement of competition. E) Time-course of cell
774	count. Here, we assume total extinction of leaders occurs during treatment, i.e. at some point
775	during treatment L=0. Note that ecosystem dynamics is reversed after perturbation is removed.





778Fig 7: Disrupting Leader to Follower feedback, s_{LF} , can trigger transient changes in the779population dynamics. A) Bifurcation diagram depicting the direction of the perturbation in780parameter space. Perturbations in s_{LF} change the location of the saddle node bifurcation boundary.781Black line – initial location of the bifurcation boundary; red line – perturbed location. B-D) Phase782plots of dynamics before (B), during (C), and after (D) blockade of s_{LF} . E) Time-course of cell count.783Note that ecosystem dynamics is reversed after perturbation is removed

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788	Fig 8: Disrupting Follower to Leader feedback, s_{FL} , can have irreversible changes in
789	dynamics leading to stabilization of cell count. A) Bifurcation diagram depicting the direction of
790	the perturbation in parameter space. Perturbations in s_{FL} change the location of the saddle node
791	bifurcation and transcritical bifurcation at zero boundaries. Black lines - initial location of the
792	bifurcation boundaries; red lines – perturbed location. B-D) Phase plots of dynamics before (B),
793	during (C), and after (D) blockade of s_{FL} . E) Time-course of cell count. Note irreversible change of
794	the ecosystem dynamics. F) Bifurcation diagram depicting the position of the saddle node
795	bifurcation point as a function of s_{LF} and s_{FL} . AD: Aggressive Dynamics; MD: Multi-modal Dynamics.
796	Here, $s_L = 1.4$ and $c = 0.2$.